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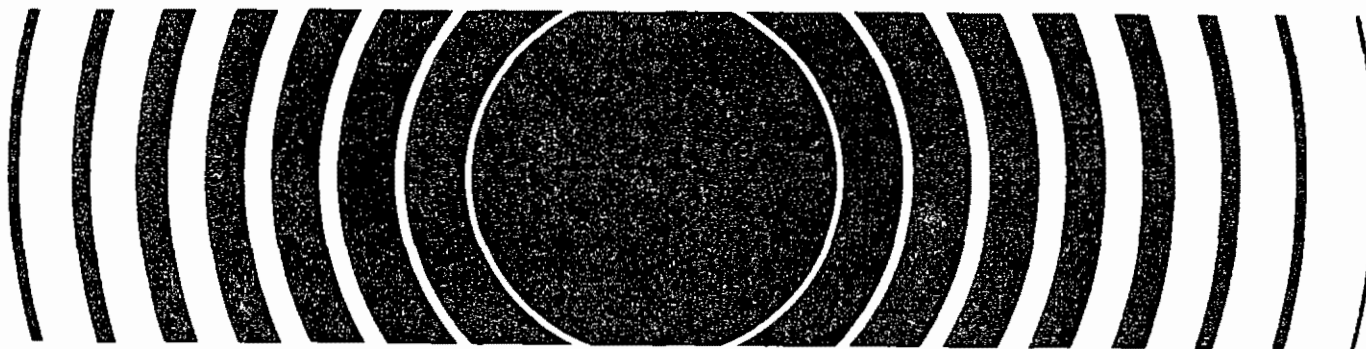
Radiation



Potential Health and Environmental Hazards of Uranium Mine Wastes

Executive Summary

Report To The Congress Of The United States



Volume 1 of 3 Volumes

POTENTIAL HEALTH AND ENVIRONMENTAL
HAZARDS OF URANIUM MINE WASTES

Executive Summary

A Report to the Congress of the United States
in Response to Public Law 95-604

June 10, 1983

U.S. Environmental Protection Agency
Office of Radiation Programs
Washington, D.C. 20460

CONTENTS

<u>SECTION</u>		<u>Page</u>
I	INTRODUCTION	1
	Purpose of the Report	1
	Contents of the Report	1
	Scope of the Report	3
	Brief Description of Uranium Mining Operations	6
II	ACTIVE MINES	8
	Number of Mines	8
	Health Impact of Air Emissions	8
	Health Impact of Water Emissions	10
	Health Impact of Solid Wastes	12
III	INACTIVE MINES	14
	Number of Mines	14
	Health Impact of Air Emissions	14
	Health Impact of Water Emissions	16
	Health Impact of Solid Wastes	16
IV	CONCLUSIONS AND RECOMMENDATIONS	19
	Conclusions	19
	Recommendations	22
V	OTHER FINDINGS	23

SECTION I

INTRODUCTION

PURPOSE OF THE REPORT

Uranium mining operations release some radioactive materials into both air and water and generate large quantities of solid wastes containing low levels of radioactive materials. Solid wastes produced by past mining operations remain on the surface at many inactive mining sites, and represent a potential health and environmental hazard similar in concept to uranium mill tailings. Contamination of surface and subsurface water supplies also represents a potential problem. To evaluate these potential problems, the Congress, in Section 114(c) of the Uranium Mill Tailings Radiation Control Act of 1978 (UMTRCA), instructed the Administrator of the Environmental Protection Agency (EPA) to prepare a report "which identifies the location and potential health, safety, and environmental hazards of uranium mine wastes together with recommendations, if any, for a program to eliminate these hazards."

This report analyzes the potential health and environmental impacts of both active and inactive uranium mines, lists the locations of these mines, identifies additional information needs, and recommends needed actions.

CONTENTS OF THE REPORT

This Executive Summary contains a brief description of the material presented in the main text, including the principal findings, conclusions, and recommendations. The full report consists of this Executive Summary, a main text, and appendices. The full report has been reviewed by the uranium mining industry, States and the Nuclear Regulatory Commission. Comments have been incorporated where possible.

Main text

The main text consists of seven chapters covering the following subject matter:

- a general description of uranium mining
- an inventory of both active and inactive uranium mines
- sources and amounts of pollutants released to the environment
- amounts of solid waste generated
- pathways of human exposure to pollutants
- health risks and environmental impacts
- recommendations and conclusions

Appendices

The appendices cover the following subjects:

- a detailed listing of the active and inactive uranium mines in the United States and their locations
- observations of existing conditions at selected inactive mines
- a description of the methodology used in the health risk and environmental impact assessments

SCOPE OF THE REPORT

This report addresses potential health impacts caused by air and water emissions and solid wastes at active and inactive underground and surface mines. We emphasize radiological impacts because we believe these to represent the most significant health hazards although nonradiological aspects of ground water and air contamination were also studied. Impacts from other mining activities, such as exploration, site preparation, and in situ leaching, were evaluated in proportion to their potential significance and the amount of available information about them.

Pathways of Exposure

Underground and surface mining release radioactivity and chemicals into air and water and generate solid wastes that may spread through wind and water erosion and release radon-222 into air. We have examined the extent to which people may be exposed to these released materials or residual solid wastes and thereby incur an increased chance of cancer or other health effects from:

- breathing air containing radon daughters,
- drinking water containing uranium and its daughters,
- eating food contaminated by either air or water, and
- living in homes on land covered by mine wastes.

Estimates of the health risks from each of these pathways are presented in this report.

Method of Analysis

Our preliminary evaluations indicated little actual environmental data is available to evaluate the impacts of releases from uranium mines. Therefore, we developed models of active and inactive mines using the available data and evaluated these impacts on a broad generic basis. To the extent possible, operating parameters and pollutant release rates characteristic of the various classes of mines were used in our models. Finally we extrapolated the health risks from the model mines to obtain an estimate of the total health effects from all active and inactive mines on regional populations within 50 miles from each mine. We estimate the risk to the total U.S. population is no greater than a factor of 3 or 4 higher than our estimates for regional populations.

The availability of information to assess the health and environmental impact from uranium mines varied greatly depending

upon the type of release and pathway of exposure. In some cases, we had to assume the most appropriate values to use in the analyses. For some release-pathway combinations, we were able to make a quantitative risk assessment. For other release-pathway combinations, the information was so limited that we could identify only the potential for impact.

We have expressed the health and environmental impacts in this report in a number of different ways:

- Estimates of the risk of cancer to individuals and to population groups
- Estimates of the risk of genetic effects to the descendants of exposed individuals and population groups
- Estimates of radioactivity and chemical concentrations in the environment and a comparison of these concentrations with air or water standards or with existing background levels
- Estimates of land areas disturbed, amounts of solid wastes generated, quantities of water discharged, and quantities of contaminants released to air and water
- Qualitative observations of a potential health impact

It must be recognized that the primary effect of radiation exposure is cancer although genetic effects are also evaluated.

Uncertainty of Health Risk Estimate

To assess the increased chance of cancer and of genetic effects occurring after exposure to radiation, Federal agencies base risk estimates on studies of persons exposed at high doses and assume that the effects at lower doses will be proportionately less. Such assessments are based on a statistical risk to all persons in a large population exposed to a known radiation dose. Because of uncertainties in the health risk analyses presented in this report, these estimates should be used carefully.

BRIEF DESCRIPTION OF URANIUM MINING OPERATIONS

The two major mining methods used in the United States are underground and surface (open pit) mining. During 1978, underground mines produced 5.5 million metric tons of ore containing 8300 metric tons of uranium oxide (U_3O_8) while surface mines produced 7.5 million metric tons of ore containing 8700 metric tons of U_3O_8 . In situ leaching, heap leaching, and mine water extraction methods accounted for the remaining 1300 metric tons of U_3O_8 production.

Underground Mining

Underground mining uses shafts and tunnels to gain access to the ore. A mine may extend underground for a mile or more at several depths. The ore is moved to the surface and stored for transport to a uranium mill. Waste rock and sub-ore* generated during mining are also stored at the surface as a waste pile. At most underground mines, these wastes remain on the surface when mining ceases.

Large capacity ventilation systems are used at underground mines to keep the radon-222 decay product concentrations in the working areas below occupational exposure limits. Air is usually forced down through the main shaft along the tunnels to the working areas and then exhausted through ventilation shafts. Large underground mines may have as many as a dozen ventilation shafts. However, while ventilation removes radon-222 decay products from the working areas, it discharges radon-222 to the atmosphere.

*Sub-ore contains uranium at a concentration uneconomical to mill. This concentration varies with the "cutoff level" of the mill receiving the ore. The cutoff level is usually determined by the cost of milling vs. the value of the recovered uranium.

Surface Mining

Surface mining is done by excavating one or more pits. The top soil and overburden above the ore are removed and stockpiled. The uranium ore is then removed and stockpiled for shipment to a uranium mill. Sub-ore is also removed from the pit during these operations and stockpiled for possible future use.

The present practice at most surface uranium mines is to backfill the mined out pits with overburden as part of a reclamation program. However, even though backfilling is performed, some waste remains on the surface after mining is completed, and the final pit may not be backfilled. Most older inactive mines were not backfilled and little or no reclamation was done.

Mine Dewatering

Since most uranium ore deposits are below the water table, groundwater must be controlled to prevent mines from flooding. Underground mines and most surface mines are dewatered to allow for excavation or shaft sinking and ore removal. Both underground and surface mines discharge this water to natural surface drainage systems. The discharged water, if necessary, is treated with barium chloride and allowed to settle to reduce radium and suspended solids before it is released. In addition to local effects, the long-term impacts on regional water availability and quality are also important considerations.

Exploratory and Development Drilling

The uranium industry has drilled approximately 1,300,000 exploratory and development drill holes through 1977. It appears from mine site surveys and aerial photography that very few drill sites have been reclaimed. Some States do require backfilling of drill holes.

The average drilling depth has increased with time and will probably continue to do so in the future. Deeper drilling will tend to increase the possibility that aquifers with good quality water may be degraded by being connected, via the drill holes, with aquifers of poorer quality water.

SECTION II

ACTIVE MINES

NUMBER OF MINES

In 1978 there were about 340 active uranium mines in the United States. A list of these mines is presented in Appendix E and includes the type of mine, location, and owner. Table 1 summarizes the locations, numbers, and types of active mines:

Table 1

Location of Active Mines in United States in 1978

<u>State</u>	<u>Surface</u>	<u>Underground</u>	<u>In situ</u>	<u>Other</u>
Colorado	5	106	0	4
New Mexico	4	35	0	3
Texas	16	0	8	1
Utah	13	108	0	3
Wyoming	19	6	3	2
Other	3	1	0	0
Total	60	256	11	13

HEALTH IMPACT OF AIR EMISSIONS-ACTIVE MINES

Radiological Impacts

Exposure to radionuclide emissions into air from active uranium mines increases the chance of cancer. These risks of cancer are the

primary public health impact from air emissions due to active uranium mines. Individuals who might be living near uranium mines are exposed to higher radiation risks than those farther away. Our estimates of potential impacts are based on model mines in the absence of adequate field data. For our model of a large underground mine we estimate that individuals living for a lifetime 1 mile from the mine would have an increased chance of fatal lung cancer of 2 in a thousand resulting primarily from breathing radon-222 decay products. The increased risk caused by the mine to an individual living 25 miles away is several hundred times lower. Risks from other types of uranium mines are somewhat lower.

We estimated the health impact from all active uranium mines operating in 1978 by multiplying the risks from the model mines by the number of active mines of each type. This procedure provides only a very rough estimate of the total population risks from all mines and is accurate only to the extent the model mine represents an average for all operations. Based on this rough extrapolation of the total risks from all mines, we estimate that the radionuclide emissions into air from all active uranium mines operating in 1978 would cause less than one fatal cancer in the regional population living around these sites.

The risk of genetic defects in future generations due to airborne radiation exposure from uranium mines is very small compared to the natural occurrence of hereditary disease. The largest potential increase in genetic defects would occur near large surface mines. Exposure of the population near a large surface mine for one year is estimated to result in a very small chance of additional genetic effects to their descendants (less than 0.0001 such effects in the population).

Nonradiological Impacts

We estimated the air concentrations of nonradioactive pollutants produced by our model mines at an assumed location of the nearest individual--1 mile from center of mine site--and determined the following emissions presented minimal potential risk to the population:

- airborne stable trace metals
- airborne combustion products from heavy equipment operation
- nonradioactive gas emissions at in situ leach mines

However, the estimated concentrations of particulates in the form of dust in ambient air near large surface mines exceeded the national ambient air quality standard. Most dust near active surface mines is caused by vehicle traffic.

HEALTH IMPACT OF WATER EMISSIONS-ACTIVE MINES

Radiological Impacts

The health risks due to radionuclide emissions to water from active uranium mines are lower than those caused by radionuclide emissions to air. Although we were able to estimate cancer risks caused by radionuclides in discharged water from our model underground and surface mines, we could not do so for in situ leach mines because of insufficient data. However, radionuclide releases in water appear to be low from in situ mines. As with our estimates of air emission impacts, models utilizing some actual data were used to develop this information.

For our model of an average underground mine, we estimated that individuals living for a lifetime 1 mile from an underground mine would have an increased chance of cancer of about one in a hundred thousand due to releases to surface water. We estimated that about one additional cancer in several hundred years might occur from the normal controlled releases from these mines.

However, mine water discharged to nearby streams can recharge shallow aquifers, many of which are presently used for drinking water or may be in the future. We do not have enough information at this time to evaluate the potential health risks from using these aquifers, but using these aquifers for drinking water could result in increased radiation exposure.

Where such a problem may exist, the state radiological program should investigate existing records to determine the contaminant levels in these aquifers due to mining, and evaluate the significance of the health risks from using these shallow aquifers. If a state determines that sufficient data do not exist to perform an evaluation, additional sampling and analyses should be performed by the state to acquire the necessary data.

Nonradiological Impacts

We estimated the concentrations of nonradioactive pollutants in the streams used by the general population of the region from our mine models. These concentrations were from dewatering the model mines and were calculated after the discharge was diluted by the receiving stream. Under these conditions, none of the pollutant concentrations alone or in combination exceeded the EPA Water Quality Criteria concentrations for use in irrigation and livestock water. However, the recharge of shallow aquifers and the use of these aquifers for drinking water present a potential problem similar to that discussed for radionuclide emissions. Thus States may want to evaluate pollution concentrations to ensure drinking water standards are met.

HEALTH IMPACT OF SOLID WASTES-ACTIVE MINESRadiological Impacts

Uranium mining operations generate large quantities of solid wastes containing low levels of radioactive materials. An average surface mine generates about 6 million metric tons of solid waste per year, while an underground mine generates considerably less--about 20 thousand metric tons per year. These wastes consist of sub-ore, waste rock, and overburden. At surface mines the sub-ore comprises only a few percent of the waste while at underground mines, because much less waste is produced, the sub-ore may comprise up to 90 percent of the waste.

Through wind, water erosion, and release of radon-222, these wastes can potentially contribute to air and water pollution. These wastes pose this hazard because they contain elevated concentrations of radium-226. Sub-ore (depending upon the cutoff grade for milling) may contain up to 50 picocuries per gram (pCi/g) of radium-226, and, even though the overburden and waste rock contain lower concentrations of radium-226 than the sub-ore, large quantities of these wastes can contain concentrations of radium-226 in excess of 5 pCi/g.* EPA has proposed that uranium mine wastes containing radium-226 in quantities greater than 5 pCi/g be listed as "hazardous wastes" under the Resource Conservation and Recovery Act (RCRA) and has also proposed regulations for the treatment, storage, and disposal of these wastes (43 FR 58946, December 18, 1978). The EPA is currently conducting an extensive study of solid wastes from mines, including uranium mines at the request of Congress. If warranted, further regulations on mining would be promulgated.

*The radium-226 concentration of most soil and rock is about 1 pCi/g.

Use of Wastes in Building Construction

Using wastes containing elevated levels of radium-226 as land fill for residential construction or building homes on land contaminated by these wastes can greatly increase the chance of lung cancer to individuals living in these structures. Radon-222 formed from the decay of radium-226 is an inert gas that readily seeps through foundations, floors, and walls and accumulates in the inside air of a house. The radon-222 then decays to daughter products which, when breathed, will lodge in the lungs and cause radiation exposure to the lung tissues. For example, the use of uranium mill tailings in the construction of homes in Grand Junction, Colorado, resulted in radon-222 decay product concentrations inside the homes that required a Federal-State remedial action program for the affected structures (Public Law 92-314). These mill wastes, however, contain much higher concentrations of radium-226 than mine wastes. A survey of homes in Florida on reclaimed land containing wastes from phosphate mining showed about 20 percent of these homes have radon-222 decay product concentrations in excess of 0.03 working level (WL).^{*} Lifetime residency in a home with this level could increase the chance of lung cancer by as much as 4 in 100--thus doubling the normal risk of lung cancer.

The mechanisms by which uranium mine wastes may cause health risks are similar to those which have occurred from uranium mill tailings and phosphate wastes. Although uranium mine wastes usually have a lower radionuclide content and are less suitable as a construction material than the sand-like tailings, these wastes are still a potential health hazard to individuals if effective waste disposal methods are not used. EPA has provided to the States survey reports of radiation anomalies that may be due to use of mine wastes in construction and will continue to support State use of this data.

^{*}A working level (WL) is any combination of short-lived radon decay products in one liter of air that will result in the ultimate emission of alpha rays with a total energy of 130,000 MeV. The working level expresses a concentration of radioactivity in the air, not how much radiation a person receives. EPA estimates that the average working level in U.S. homes is about 0.004 WL.

SECTION IIIINACTIVE MINESNUMBER OF MINES

There are about 3400 inactive uranium mines in the United States. A list of these mines developed from computer listings maintained by the U. S. Department of Energy is presented in Appendix F including the type of mine, location, and owner. The following table summarizes the numbers and types of inactive mines by State:

Table 2

Location of Inactive Mines in United States

<u>State</u>	<u>Surface</u>	<u>Underground</u>	<u>Other</u>
Arizona	135	189	2
Colorado	263	902	52
New Mexico	34	142	12
South Dakota	111	30	0
Utah	378	698	17
Wyoming	223	32	10
Other	108	43	8
Total	1252	2036	101

HEALTH IMPACT OF AIR EMISSIONS- INACTIVE MINESRadiological Impact

Radionuclide emissions into air at inactive mine sites are small compared to the emissions from active mines according to our estimates of model mines. The principal radionuclide emitted,

radon-222, emanates from unsealed mine vents, portals and residual waste piles. This causes only small increases in the risk of lung cancer to individuals living near these mine sites. Utilizing the same models as for the active mines, we estimated risks of cancer from radon-222 emissions to air from our model inactive mines.

By multiplying the risks from our model mines by the number of inactive mines of each type, we extrapolated the total number of potential cancers from all inactive mines. This procedure provides only a very rough approximation of the total risk from all inactive mines.

By these estimates, radon-222 emissions from inactive uranium mines would produce the following cancer risks:

- Individuals living for a lifetime 1 mile from an inactive mine would have an increased chance of lung cancer of about 2-3 in 100,000.
- The amount of radon-222 released each year from all inactive uranium mine sites would cause about 0.1 lung cancers in the regional population around these sites.

Nonradiological Impacts

We did not identify any significant health impact associated with nonradiological air emissions at inactive uranium mines. Our estimates of dust emissions from wind erosion of waste piles showed that insignificant concentrations of nonradiological pollutants would exist in air at these inactive sites.

HEALTH IMPACT OF WATER EMISSIONS-INACTIVE MINES

The extent to which inactive surface and underground mines harm water quality is poorly understood. Ground water in contact with ore bodies and consequently in mines typically contains radionuclides and trace elements, and the flow of the water away from the site carries dissolved and suspended radionuclides and trace elements.

Site specific studies are needed to determine the present and potential impacts of inactive uranium mines on both surface and groundwater quality. As with active mines, the potential exists for contamination of drinking water supplies. States may desire to conduct sampling of drinking water at a few sites in the vicinity of inactive mining districts to provide data to evaluate whether such a potential is valid.

HEALTH IMPACT OF SOLID WASTES-INACTIVE MINES

Surface Mines

We estimate that over 1 billion tons of solid wastes were generated at surface uranium mines through 1978. These wastes consist of sub-ore and overburden. The sub-ore, which may comprise about 3 percent of the total wastes, contains significantly elevated concentrations of radium-226 (up to 100 pCi/g).^{*} Although the overburden contains much lower concentrations of radium-226 than the sub-ore, large quantities of these wastes can contain radium-226 in concentrations in excess of 5 pCi/g--the level EPA has proposed be used to judge whether wastes should be considered as a candidate for designation as hazardous waste under RCRA. Such a determination would require that specified disposal methods be developed for these mine wastes.

^{*}The radium-226 concentration of normal soil and rock is about 1 pCi/g.

In many surface mines opened since 1970, the general practice is to backfill the mined-out pits with wastes as part of a reclamation program. However, at most older inactive surface mines, little or no reclamation was done.

Underground Mines

We estimate that about 30 million tons of solid waste consisting mostly of sub-ore were generated at underground uranium mines through 1978. As in surface mining, the sub-ore contains significantly elevated concentrations of radium-226 (up to 100 pCi/g). There has been very little reclamation at inactive underground mine sites, so most of these wastes remain on the surface at these sites.

Use of Wastes in Building Construction

As discussed in the section on active mines, uranium mine wastes would present a significant hazard to individuals if homes are built on land contaminated by these wastes or if these wastes are used in construction materials for homes. Individuals living in these homes could have an increased chance of lung cancer from breathing radon-222 decay products. The extent to which uranium mine wastes have previously been used for these purposes is not well known.

However, some information is available which shows that uranium mine wastes may have been widely used as landfill in the construction of various types of buildings. In 1972 EPA and the former Atomic Energy Commission (AEC) tried to identify locations of higher-than-normal levels of gamma radiation in an attempt to locate uranium mill tailings. During this study, over 500 locations were identified where uranium ore was believed to be the source of elevated gamma radiation. Since it is unlikely that ore-grade

material would be used as landfill, we suspect that uranium mine wastes (perhaps sub-ore) may be the source of the abnormal gamma radiation at these sites.

In order to better define the off-site use of uranium mine wastes, EPA is studying the extent to which these wastes have been used away from the mine sites for landfill or in construction materials for use in homes. If mine wastes were involved in construction of homes, a health risk from radon-222 emissions would exist. A preliminary survey has already been completed and the information has been shared with the interested agencies in appropriate States.

SECTION IVCONCLUSIONS AND RECOMMENDATIONS

The evaluation of the potential impacts of uranium mining was performed largely by means of analytical studies of model facilities. We believe that the results give an adequate representation of the industry. In order to determine the extent of possible problems, our studies were specifically designed to give conservative results. It should be recognized that actual mines may operate under conditions producing substantially smaller impacts than the results presented.

Compared to uranium milling, health and environmental effects of uranium mining are not as well understood, despite the existence of over 3000 active and inactive mines. We have noted throughout this report instances of the absence or inadequacy of pertinent information.

CONCLUSIONSSolid Wastes

Solid uranium mining wastes are potentially hazardous to health when used as building materials or when buildings are constructed on land containing such wastes. The hazard arises principally from increased risk of lung cancer due to radon-222. In a 1972 survey of communities in uranium milling and mining regions, EPA and the former Atomic Energy Commission found more than 500 locations where such wastes had been used.

Airborne Effluents

a) Individuals living very near active underground mine exhaust vents would have an increased risk of lung cancer caused by exposure to radon-222 emissions. Surface mines and in situ mines are less hazardous, and inactive mines do not have significant radon-222 emissions. Other airborne radioactive emissions from all types of mines are judged to be smaller.

b) The number of additional cancers committed per year in regional populations due to radionuclide air emissions from the approximately 340 active mines and 3300 inactive mines was estimated to be about 0.6 cancers in 1978. This number of estimated additional cancers is small, about *one-third of the estimated* additional cancers in regional populations due to radon emissions from the 24 inactive uranium mill tailings piles addressed by Title I of the Uranium Mill Tailings Radiation Control Act. (These mill tailings piles represent about 13 percent of all tailings currently existing due to U.S. uranium milling and mining). These potential effects are not of sufficient magnitude to warrant corrective measures, especially considering the large number of sites involved.

c) The following emissions were judged to cause an insignificant health risk at all types of mines:

1. airborne nonradioactive trace metals
2. airborne combustion products from heavy-duty equipment operations
3. nonradioactive emissions from in situ leach sites

d) Airborne dust near large surface mines (primarily caused by vehicular traffic) may exceed the National Ambient Air Quality Standard for particulate matter.

Waterborne Effluents

a) We estimate that an insignificant health risk accrues currently to populations from waterborne radioactivity from an average existing mine.

b) Uranium mine dewatering and water discharges, which are increasing as more and deeper mines are created, may in the future have significant effects on water quality. Current treatment practices are controlling the release of radioactivity into surface waters.

c) Water in inactive surface and underground mines usually contains radionuclides and trace elements in concentrations comparable to ground water in contact with ore bodies. Some abandoned underground mines in certain areas of Colorado and Utah probably discharge such waters to nearby streams and shallow aquifers. Available data is not sufficient to conclude whether or not there is a problem.

d) We could not determine, using models, that there is no health hazard to individuals who drink water drawn from surface or underground sources. Water discharges from active mines to nearby streams and stream channels may extensively recharge shallow aquifers, many of which are either now used or could be used for drinking water. Such determinations must be made on a site-specific basis, and take account of the additive effects of multiple mines. These studies can be made easily a part of State or utility surveillance programs.

Exploratory and Development Drilling

Harm from effluents due to exploratory and developmental drilling is probably small compared to effects of operating mines. Under current regulations and practices, however, aquifers penetrated at different levels can mix, creating the potential for degrading good quality groundwater.

RECOMMENDATIONS TO CONGRESS

1) Based on this study, we do not believe at this time that Congress needs to enact a remedial action program like that for uranium mill tailings. This is principally because uranium mine wastes are lower in radioactivity and not as desirable for construction purposes as uranium mill tailings. Nonetheless, some mining waste materials appear to have been moved from the mining sites, but not to the extent that mill tailings were.

2) Some potential problems were found that might require regulatory action, but none of these appear to require new Congressional action at this time.

SECTION VOTHER FINDINGS

1) Regulations may be needed to control wastes at active uranium mines to preclude off-site use and to minimize the health risks from these materials. These regulations would need to address the use of the materials for construction purposes as well as ultimate disposal of the materials.

EPA proposed such regulations in 1978 under the Resource Conservation and Recovery Act (RCRA). In 1980, Congress amended RCRA to require further EPA studies before promulgating general regulations for mining wastes. An EPA study by the Office of Solid Wastes on all types of mines, including uranium mines, is currently being conducted. The amendment does not affect EPA's authority to regulate use of uranium mine wastes in construction or reclamation of lands containing such wastes.

2) Standards are needed to control human exposure from radioactive air emissions from uranium mines. This is principally because of potential exposure to individuals living near large underground uranium mines rather than concerns regarding the exposure of regional populations. We have proposed such standards under Section 112 of the Clean Air Act.

3) EPA has conducted two field studies in 1972 and 1978 which define possible sites at which mine wastes may have been used in construction or placed around buildings. The information developed in these studies has been sent to State health departments. The States should conduct follow-up studies, as appropriate, to determine whether there are problems at these sites.

4) The adequacy with which NPDES permits protect individuals who may obtain drinking water near the discharge points for uranium mine dewatering should be evaluated by States. Under the Public Water Systems provision of the Safe Drinking Water Act, radionuclide standards now exist for drinking water.

5) Some site specific studies should be considered by States to determine the extent to which inactive uranium mines may be significant water pollution sources.

6) States with uranium mines should determine the feasibility of controlling fugitive dust from large surface mines and incorporate the recommendations in State Implementation Plans.

7) States should require borehole plugs in drilling operations that will prevent interaquifer mixing (exchange) and also seal drilling holes at the surface.

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HAZARDS OF URANIUM MINE WASTES

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in Response to Public Law 95-604

Volume 2 of 3 Volumes

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	Page
1.4.2 State Regulations	1-42
1.4.2.1 Colorado	1-45
1.4.2.2 New Mexico	1-47
1.4.2.3 Texas	1-49
1.4.2.4 Utah	1-49
1.4.2.5 Washington	1-50
1.4.2.6 Wyoming	1-51
1.5 References	1-53
2.0 Inventory of Uranium Mines	2-1
2.1 References	2-16
3.0 Potential Sources of Contaminants to the Environment and Man	3-1
3.1 Background Concentrations of Radionuclides and Trace Metals	3-1
3.1.1 Naturally Occurring Radionuclides	3-1
3.1.2 Stable Elements	3-6
3.2 Water-Related Aspects of Uranium Mining	3-11
3.2.1 Previous and Ongoing Hydrologic and Water Quality Studies Related to Uranium Mining	3-11
3.2.2 Mine Water Management	3-13
3.2.3 Water Quality Effects of Mine Water Discharge	3-23
3.2.3.1 Behavior of Contaminants in the Aqueous Environment	3-23
3.2.3.1.1 Dilution and Suspended Sediment Transport	3-25
3.2.3.1.2 Sorption and Desorption	3-25
3.2.3.1.3 Precipitation	3-28
3.2.3.1.4 Biological Assimilation and Degradation	3-30
3.2.3.1.5 Complexation	3-31
3.2.3.2 Results of Field Studies in Uranium Mining Areas	3-32

	Page
3.2.3.2.1 Colorado	3-32
3.2.3.2.2 Wyoming	3-34
3.2.3.2.3 Texas	3-38
3.2.3.2.4 New Mexico	3-39
3.2.3.3 Summary	3-43
3.3 Surface Mining	3-44
3.3.1 Solid Wastes	3-44
3.3.1.1 Overburden Piles	3-45
3.3.1.2 Ore Stockpiles	3-54
3.3.1.3 Sub-Ore Piles	3-60
3.3.1.4 Reclamation of Overburden Piles	3-62
3.3.2 Mine Water Discharge	3-63
3.3.2.1 Data Sources	3-63
3.3.2.2 Quantity and Quality of Discharge	3-64
3.3.3 Hydraulic and Water Quality Effects of Surface Mine Discharge	3-68
3.3.3.1 Runoff and Flooding in the Model Surface Mine Area	3-68
3.3.3.1.1 Study Approach	3-68
3.3.3.1.2 Description of Area	3-70
3.3.3.1.3 Method of Study	3-72
3.3.3.1.4 Discussion of Results	3-77
3.3.3.2 Impacts of Seepage on Groundwater	3-89
3.3.4 Gases and Dusts from Mining Activities	3-93
3.3.4.1 Dusts and Fumes	3-93
3.3.4.2 Radon-222 from the Pit, Storage Piles, and Ore Handling	3-99
3.4 Underground Mining	3-107
3.4.1 Solid Wastes	3-107
3.4.1.1 Waste Rock Piles	3-109
3.4.1.2 Ore Stockpiles	3-110
3.4.1.3 Sub-Ore Piles	3-112

	Page
3.4.2 Mine Water Discharge	3-113
3.4.2.1 Data Sources	3-113
3.4.2.2 Quality and Quantity of Discharge	3-114
3.4.3 Hydraulic and Water Quality Effects of Underground Mine Discharge	3-120
3.4.3.1 Runoff and Flooding in the Model Underground Mine Area	3-120
3.4.3.1.1 Study Approach	3-120
3.4.3.1.2 Description of Area	3-122
3.4.3.1.3 Estimate of Sub-basin Flood Flow	3-124
3.4.3.1.4 Prediction of Sub-basin Water Quality	3-132
3.4.3.2 Impacts of Seepage on Groundwater	3-149
3.4.4 Gases and Dusts from Mining Activities	3-155
3.4.4.1 Radon-222 in Mine Exhaust Air	3-155
3.4.4.2 Aboveground Radon-222 Sources	3-157
3.4.4.3 Dusts and Fumes	3-159
3.5 In Situ Leach Mining	3-168
3.5.1 Solid Wastes	3-169
3.5.2 Associated Wastewater	3-172
3.5.3 Airborne Emissions	3-174
3.5.4 Excursion of Lixiviant	3-178
3.5.5 Restoration and Reclamation	3-179
3.6 Other Sources	3-185
3.6.1 Mineral Exploration	3-185
3.6.1.1 Environmental Considerations	3-187
3.6.1.2 Radon Losses from Drill Holes	3-192
3.6.1.3 Groundwater	3-193
3.6.1.4 Fumes	3-193
3.6.1.5 Model Drilling	3-194
3.6.2 Precipitation Runoff from Uranium Mines	3-194
3.7 Inactive Mines	3-204
3.7.1 Inactive Surface Mines	3-204
3.7.1.1 Waste Rock Piles	3-213

	Page
3.7.1.2 Radon-222 from the Mine Area	3-216
3.7.1.3 Land Surface Gamma Radiation	3-222
3.7.2 Inactive Underground Mines	3-227
3.7.2.1 Waste Rock Piles	3-229
3.7.2.2 Radon-222 from the Mine Area	3-232
3.7.2.3 Land Surface Gamma Radiation	3-237
3.8 References	3-242
4.0 Description of Model Mines	4-1
4.1 Surface Mine	4-1
4.2 Underground Mine	4-4
4.3 In Situ Leach Mine	4-7
4.4 Inactive Surface Mine	4-8
4.5 Inactive Underground Mine	4-9
5.0 Potential Pathways	5-1
5.1 General	5-1
5.1.1 Vegetation	5-1
5.1.2 Wildlife	5-1
5.1.3 Land Use	5-2
5.1.4 Population Near Mining Areas	5-2
5.1.5 Population Statistics of Humans and Beef Cattle	5-12
5.2 Prominent Environmental Pathways and Parameters for Aqueous Releases	5-12
5.2.1 Individual Committed Dose Equivalent Assessment	5-13
5.2.2 Collective (Population) Dose Equivalent Assessment	5-15
5.3 Prominent Environmental Pathways and Parameters for Atmospheric Releases	5-16
5.3.1 Individual Committed Dose Equivalent Assessment	5-16
5.3.2 Collective (Population) Dose Equivalent Assessment	5-18
5.4 Mine Wastes Used in the Construction of Habitable Structures	5-19
5.5 References	5-20
6.0 Health and Environmental Effects	6-1
6.1 Health Effects and Radiation Dosimetry	6-1

	Page
6.1.1 Radioactive Airborne Emissions	6-1
6.1.2 Nonradioactive Airborne Emissions	6-26
6.1.2.1 Combustion Products	6-26
6.1.2.2 Nonradioactive Gases	6-28
6.1.2.3 Trace Metals and Particulates in the Form of Dust	6-28
6.1.3 Radioactive Aquatic Emissions	6-35
6.1.4 Nonradioactive Aquatic Emissions	6-42
6.1.5 Solid Wastes	6-44
6.1.5.1 Radium-226 Content	6-44
6.1.5.2 Estimates of Potential Risk	6-45
6.1.5.3 Using Radium Bearing Wastes in the Construction of Habitable Structures	6-46
6.1.5.3.1 Use of Uranium Mine Wastes	6-48
6.2 Environmental Effects	6-48
6.2.1 General Considerations	6-48
6.2.2 Effects of Mine Dewatering	6-52
6.2.3 Erosion of Mined Lands and Associated Wastes	6-54
6.2.4 Land Disturbance from Exploratory and Development Drilling	6-56
6.2.5 Land Disturbance from Mining	6-59
6.2.5.1 Underground Mines	6-59
6.2.5.2 Surface Mines	6-59
6.2.6 Retirement Phase	6-59
6.3 References	6-73
7.0 Summary and Recommendations	7-1
7.1 Overview	7-1
7.2 Sources and Concentrations of Contaminants	7-1
7.2.1 Surface and Underground Mines	7-1
7.2.2 In Situ Leach Mines	7-7
7.2.3 Uranium Exploration	7-8
7.3 Exposure Pathways	7-8
7.4 Potential Health Effects	7-9

	Page
7.4.1 Radioactive Airborne Emissions	7-9
7.4.2 Nonradioactive Airborne Emissions	7-13
7.4.3 Radioactive Aqueous Emissions	7-13
7.4.4 Nonradioactive Aqueous Emissions	7-15
7.4.5 Solid Wastes	7-16
7.5 Environmental Impacts	7-16
7.5.1 Land and Water Contamination	7-17
7.5.2 Effects of Mine Dewatering	7-22
7.5.3 Erosion of Mined Lands and Associated Wastes	7-22
7.5.4 Exploratory and Development Drilling	7-23
7.5.5 Underground Mining	7-23
7.5.6 Surface Mining	7-24
7.6 Regulatory Perspective	7-25
7.7 Conclusions and Recommendations	7-25
7.7.1 Conclusions	7-26
7.7.1.1 Solid Wastes	7-26
7.7.1.2 Airborne Effluents	7-26
7.7.1.3 Waterborne Effluents	7-27
7.7.1.4 Exploratory and Development Drilling	7-27
7.7.2 Recommendations to Congress	7-28
7.8 Other Findings	7-28
7.9 References	7-29

Appendixes (See Volume 3)

- A. Summary of Federal Laws Potentially Affecting Uranium Mining
- B. Federal Water Programs and Rights Activities
- C. Congressionally Approved Compacts that Apportion Water
- D. State Laws, Regulations, and Guides for Uranium Mining
- E. Active Uranium Mines in the United States
- F. Inactive Uranium Mines in the United States
- G. General Observations of Uranium Mine Sites in Colorado, New Mexico, Texas, and Wyoming.

- H. Influence of Mine Drainage on Seepage to Groundwater and Surface Water Outflow
- I. Computation of Mass Emission Factors for Wind Erosion
- J. Aquatic Dosimetry and Health Effects Models and Parameter Values
- K. Airborne Pathway Modeling
- L. Health Risk Assessment Methodology

FIGURES

	Page
1.1 Uranium mining regions in the western United States	1-4
1.2 The percent of \$50 U_3O_8 reserves located in the principal mining states	1-8
1.3 The percent of \$50 U_3O_8 reserves located in the various mining regions	1-9
1.4 Artist's conception of open pit mining operation and support facilities.	1-13
1.5 Generalized underground mine showing modified room and pillar method of mining	1-17
1.6 Diagrams of some common injection-recovery well patterns used in uranium in situ leach mining	1-26
2.1. Location of active and inactive uranium mines and principal uranium mining districts in Colorado	2-6
2.2 Location of active and inactive uranium mines and principal uranium mining districts in the Uravan Mineral Belt of western Colorado	2-7
2.3 Location of active and inactive uranium mines in the Grants Mineral Belt and other areas of New Mexico	2-8
2.4 Location of active, inactive, and proposed surface and in situ uranium mines in Texas	2-9
2.5 Location of uranium mines and mining districts in Utah	2-10
2.6 Location of uranium mines and principal uranium mining districts in southeastern Utah	2-11
2.7 Location of active and inactive uranium mines and principal uranium mining areas in Wyoming	2-12
2.8 Location of active and inactive uranium mines in the Gas Hills and Crooks Gap-Green Mountain areas of central Wyoming	2-13
2.9 Location of active and inactive uranium mines in the Shirley Basin, South Powder River Basin, and Pumpkin Buttes areas of Wyoming.	2-14

3.1	The uranium decay series showing the half lives and mode of decay	3-2
3.2	The thorium decay series showing the half lives and mode of decay	3-3
3.3	Disposition of drainage water from active surface and underground uranium mines	3-14
3.4	Location of mines, ore and waste storage areas and monitoring stations at the Morton Ranch mine, South Powder River Basin, Wyoming	3-37
3.5	Location of study areas, sampling stations and uranium mines, Poison Canyon area, McKinley County, New Mexico	3-41
3.6	Sample locations for radionuclides and select trace metals in sediments, San Mateo mine, New Mexico	3-42
3.7	Potential sources of environmental contamination from active open pit uranium mines	3-46
3.8	Storage pile configurations assumed at surface and underground mines	3-48
3.9	Sketch of sub-basin, basin, and regional basin showing orientation of principal drainage courses, areas of drainage, and location of mines	3-69
3.10	Average monthly flows for the Cheyenne River and Lance Creek near Spencer, Wyoming, for the period 1948-1970	3-71
3.11	Suspended sediment concentration to discharge, Salt Wells Creek and Tributaries, Wyoming	3-75
3.12	Relation of discharge and specific conductance to time at Salt Wells Creek, Green River Basin, Wyoming	3-76
3.13	Periods of no flow in Lance Creek and the Cheyenne River near Riverton, Wyoming, for the period 1948-1978.	3-82
3.14	Configuration of open pit model mines	3-105
3.15	Potential sources of environmental contamination from active underground uranium mines	3-108

3.16 Sketch of sub-basin, basin, and regional basin showing orientation of principal drainage courses, areas of drainage, and location of mines in the New Mexico model area	3-121
3.17 Average monthly flows for the period of record for the Rio San Jose and the Rio Puerco in New Mexico	3-129
3.18 Periods of no flow in the Rio San Jose and Rio Puerco	3-131
3.19 Total flow volumes in one-day periods for floods of various recurrence intervals in the sub-basin and basins in New Mexico	3-133
3.20 Total flow volumes in seven-day periods for floods of various recurrence intervals in the sub-basin and basins in New Mexico	3-134
3.21 Principal streams and surface water sampling stations in the Churchrock and Gallup areas	3-145
3.22 Average depth of exploratory drilling in the U.S. uranium industry from 1948 to present	3-188
3.23 Annual waste to ore ratios for surface mining of uranium (1948 to 1979)	3-209
3.24 Cross section of model inactive surface mine	3-214
3.25 Results of gamma exposure rate survey at the 1601 pit and environs, Morton Ranch uranium mine, Converse County, Wyoming	3-226
3.26 Waste to ore ratios for inactive underground uranium mines from 1932 to 1977	3-228
3.27 Radon-222 concentrations in mine air discharged by natural ventilation	3-236
3.28 Gamma radiation survey around an inactive underground uranium mine in New Mexico	3-240
5.1 Potential airborne pathways in the vicinity of uranium mines	5-17

6.1	Average indoor radon-222 decay product measurements (in working levels) as a function of average radium-226 concentration in soil	6-47
6.2	Example of natural reclamation of drill sites	6-57
6.3	Inactive underground mine site	6-60
6.4	Example of active and inactive surface mining activities	6-63
6.5	Mine wastes eroded by ephemeral streams in the Mesa Montanosa area, New Mexico	6-64
6.6	Basal erosion of a uranium mine waste pile by an ephemeral stream in the Mesa Montanosa area, New Mexico	6-65
6.7	Scattered piles of mine waste at the Mesa Top mine, Mesa Montanosa, New Mexico	6-66
6.8	Close up view of easily eroded sandy and silty mine waste from the Mesa Top mine, Mesa Montanosa, New Mexico	6-66
6.9	Gullying and sheet erosion of piled and spread mine wastes at the Dog Incline uranium mine, Mesa Montanosa, New Mexico	6-67
6.10	Recent erosion of unstabilized overburden piles at the inactive Galen mine, Karnes County, Texas	6-68
6.11	Unstabilized overburden piles and surface water erosion at the Galen mine, Karnes County, Texas	6-68
6.12	Aerial view of the Manka mine, Karnes County, Texas	6-70
6.13	Overburden pile showing the weak vegetative cover and gullying associated with improper stabilization at the Manka mine, Karnes County, Texas,	6-70
6.14	Inactive Hackney mine, Karnes County, Texas	6-72

Appendix G

G.1	Plan view of inactive underground uranium mine No. 1, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-3
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	Page
G.2 Sectional view of inactive underground uranium mine No. 2, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-4
G.3 Plan view of inactive underground uranium mine No. 3, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-6
G.4 Sectional view of inactive underground uranium mine No. 4, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-7
G.5 Plan view of inactive underground uranium mine No. 5, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-8
G.6 Plan view of inactive underground uranium mine No. 6, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-9
G.7 Plan view of inactive underground uranium mine No. 7, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado	G-11
G.8 Plan view of inactive underground uranium mine No. 8, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado	G-12
G.9 Plan view of inactive underground fluorspar uranium mine No. 9, related waste rock piles, and surface gamma exposure rates, near Jamestown, Colorado	G-13
G.10 Plan view of inactive underground uranium mine No. 10, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado	G-15
G.11 Typical mine waste pile associated with a small- to medium-sized inactive underground uranium mine, Uravan Mineral Belt, Colorado	G-16
G.12 Side view of a typical underground uranium mine located on the rim of a sandstone mesa, Uravan Mineral Belt, Colorado	G-16

	Page
G.13 Mine waste accumulations near the portal of a typical under-ground rim-type uranium mine, western Colorado	G-17
G.14 Mine waste dump associated with a typical rim-type under-ground uranium mine, western Colorado	G-17
G.15 Movement of fluorspar-uranium mine wastes from a tailings pile into a stream, Jamestown area, Colorado	G-19
G.16 1972 aerial photograph of the Galen and Pawelek open pit mines, Karnes County, Texas	G-27
G.17 1978 aerial photograph of the Galen and Pawelek open pit mines, Karnes County, Texas	G-27
G.18 Results of gamma exposure rate survey at the 1601 pit and environs, Morton Ranch uranium mine, Converse County, Wyoming	G-31
G.19 Location of sampling stations at the Morton Ranch mine, South Powder River Basin, Wyoming	G-33
G.20 Sample locations for radionuclides and select trace metals in sediments, San Mateo mine, New Mexico	G-34

Appendix H

H.1 Wyoming model area sub-basin drainage system	H-3
H.2 Model area stream cross section	H-3
H.3 New Mexico model area sub-basin drainage system	H-9

Appendix J

J.1 Surface stream flow pattern within drainage area	J-3
J.2 Conservation of mass relationship for resuspension model	J-13

TABLES

		Page
1.1	Domestic uranium production by state from 1948 to January 1, 1979	1-3
1.2	Projected annual nuclear capacity (GWe) in the U.S. .	1-6
1.3	Domestic uranium reserves by state as of January 1, 1979	1-7
1.4	The quantities of U_3O_8 produced in 1978 by the various mining methods	1-10
1.5	The predicted methods of mining ore reserves . . .	1-10
1.6	Summary of current in situ leaching operations as of January 1, 1978	1-21
1.7	Trace metal concentrations of recirculated acid and alkaline lixiviants	1-25
1.8	Federal laws, regulations, and guides for uranium mining	1-32
1.9	Requirements to obtain rights to prospect or explore by federal, state and private lands	1-37
1.10	Requirements to obtain rights to mine ore by federal, state and private lands	1-39
1.11	Requirements for mining and environmental plans by federal, state and private lands	1-41
1.12	State laws, regulations, and guides for uranium mining .	1-43
2.1	Type of U.S. uranium properties	2-3
2.2	The location and type of active uranium properties .	2-4
2.3	The location and type of inactive uranium properties .	2-5
2.4	Cumulative ore production through January 1, 1979 . .	2-15
3.1	Gamma-ray energy released by one gram of rock . .	3-4
3.2	Radionuclide content and dose equivalent rates from common rocks and soil	3-4
3.3	Average dose equivalent rates due to terrestrial radiation in western mining states	3-5
3.4	Radionuclide concentrations in surface and groundwater in the vicinity of a proposed uranium project . .	3-7
3.5	Concentrations of selected elements in igneous and sedimentary rocks	3-8

3.6	Concentrations of selected elements in surface water at five locations in the vicinity of a proposed uranium project	3-9
3.7	Concentrations of selected elements in groundwater at six locations in the vicinity of a proposed uranium project.	3-10
3.8	Estimated average concentrations of three metals in U.S. streams	3-6
3.9	Summary of feed water sources for active U.S. uranium mills	3-17
3.10	Current and projected uranium mine discharges in the Grants Mineral Belt, New Mexico	3-21
3.11	Estimated surface areas associated with overburden piles	3-50
3.12	Particle size distributions of mill tailings and mine overburden	3-52
3.13	Natural radionuclide concentrations in various common rock types	3-52
3.14	Annual average airborne radionuclide concentrations in the vicinity of an open pit uranium mine	3-53
3.15	Uranium and stable element concentrations measured in rock and soil samples from two uranium mines	3-55
3.16	Concentration of radionuclides and stable elements in overburden rock from the model surface mines	3-56
3.17	Estimated average areas of ore pile surface and pad	3-57
3.18	Distribution of ore reserves by the type of host	3-59
3.19	Average stable element concentrations in sandstone ores of New Mexico	3-59
3.20	Estimated average surface areas of sub-ore piles during the 17-year active mining period	3-61
3.21	Summary of average discharge and water quality data for uranium mines in Wyoming and a comparison with NPDES limits	3-65
3.22	Water quality associated with surface and underground mines in various stages of construction and operation	3-67

	Page
3.23 Peak discharge and total volume for floods of 2, 5, 10, 25, 50 and 100 year recurrence intervals	3-78
3.24 Summary of calculated total flow in the Wyoming model area sub-basin using the USGS and SCS methods	3-79
3.25 Annual contaminant loading from one uranium mine and resulting concentrations in floods within the sub-basin for return periods of 2 to 100 years	3-84
3.26 Concentrations in basin and regional basin streams as a result of surface mine discharge	3-86
3.27 Comparison of potable and irrigation water standards and surface water quality affected by surface mine drainage	3-87
3.28 Northeastern Wyoming groundwater sources	3-91
3.29 Groundwater quality of wells sampled by the three major uranium producers in the South Powder River Basin, Wyoming	3-92
3.30 Estimated air pollutant emissions from heavy-duty equipment at surface mines	3-94
3.31 Average annual dust emissions from mining activities	3-97
3.32 Average annual emissions of radionuclides (μ Ci) and stable elements (kg) from vehicular dust at the model surface mines	3-100
3.33 Average annual emissions of radionuclides (μ Ci) and stable elements (kg) from mining activities at the model surface mines	3-101
3.34 Average annual emissions of radionuclides (μ Ci) and stable elements (kg) in wind suspended dust at the model surface mines	3-102
3.35 Radon-222 releases during surface mining	3-107
3.36 Estimated average surface areas of waste rock piles at underground mines	3-111
3.37 Estimated surface areas of ore stockpiles at underground mines	3-111
3.38 Estimated average surface areas of sub-ore piles at underground mines	3-113

3.39	Summary of average discharge and water quality data for underground uranium mines in the Colorado Plateau Region (Colorado, New Mexico, Utah) and a comparison with NPDES limits	3-116
3.40	Water quality associated with underground mines in various stages of construction and operation	3-119
3.41	Total flow volume for sub-basin floods of 1- and 7-day durations and return periods of 2, 5, 10, and 25 years	3-126
3.42	Summary of area, discharge, and irrigated acreage for the sub-basin, basin, and regional basin hydrographic units in New Mexico	3-127
3.43	Dilution factors for the Rio San Jose, Rio Puerco, and Rio Grande for 1-day flood flows with a 2-year recurrence interval	3-135
3.44	Annual contaminant loading from 14 uranium mines and resulting concentrations in sub-basin floods and in the average annual flow of the Rio San Jose, Rio Puerco, and Rio Grande	3-137
3.45	Comparison of potable and irrigation water standards and surface water quality affected by underground mine drainage.	3-140
3.46	Radiochemical and stable element/compound water quality for selected acid and alkaline leach uranium mill tailings ponds in the United States	3-142
3.47	Summary of flood runoff water quality and uranium millpond quality	3-143
3.48	Flow and water quality in the Puerco River near Churchrock and Gallup, New Mexico	3-146
3.49	Groundwater quality in principal aquifers in the Grants Mineral Belt, New Mexico	3-151
3.50	Groundwater quality associated with the San Mateo Creek and Rio Puerco (west) drainages in the Grants Mineral Belt, New Mexico	3-154

3.51	Estimated annual radon-222 emissions from underground uranium mining sources	3-158
3.52	Estimated air pollutant emissions from heavy-duty equipment at underground uranium mines	3-161
3.53	Estimated average annual dust emissions from underground mining activities	3-163
3.54	Average annual emissions of radionuclides (μ Ci) and stable elements (kg) from mining activities at the model underground mines.	3-165
3.55	Average annual emissions of radionuclides (μ Ci) and stable elements (kg) in wind suspended dust at the model underground mines	3-166
3.56	Average annual emissions of radionuclides (μ Ci) and stable elements (kg) from vehicular dust at the model underground mines	3-167
3.57	Estimated quantities of wastewater produced by an in situ leaching operation	3-173
3.58	Estimated average concentrations and annual accumulation of some contaminants in waste water	3-175
3.59	Estimated average annual airborne emissions from the hypothetical in situ leaching facility	3-176
3.60	Estimated average concentrations and annual and total accumulations of some contaminants in restoration wastewater	3-181
3.61	A comparison of contaminant concentrations in pre-mining groundwater and pre-restoration mine water	3-182
3.62	Estimates of exploratory and development drill holes (1948-1979)	3-189
3.63	Estimated source terms per bore hole for contemporary surface drilling for uranium	3-195
3.64	Airborne dusts produced at an average mine site from exploratory and development drilling	3-196

	Page
3.65	Estimates of emissions from drill rig diesel power source 3-196
3.66	Sediment yields in overland flow from uranium mining areas 3-202
3.67	Consolidated list of inactive uranium producers by State and type of mining 3-206
3.68	Uranium mine waste and ore production 3-207
3.69	Cumulative uranium mine waste and ore production 3-211
3.70	Average annual emissions of radionuclides (μ Ci) and stable elements (kg) in wind suspended dust at the model inactive surface mine 3-217
3.71	Average radon flux of inactive uranium mill tailings piles 3-218
3.72	Average radon flux measured at inactive uranium mine sites 3-220
3.73	Background radon flux estimates 3-221
3.74	Summary of estimated radon-222 releases from inactive surface mines 3-223
3.75	Summary of land surface gamma radiation surveys in New Mexico, Texas, and Wyoming 3-225
3.76	Average annual emissions of radionuclides (μ Ci) and stable elements (kg) in wind suspended dust at the model inactive underground mine 3-231
3.77	Summary of radon-222 releases from inactive underground mines 3-238
3.78	Summary of land surface gamma radiation surveys in Colorado and New Mexico 3-239
5.1	Number of uranium mines and population statistics for counties containing uranium mines. 5-3
5.2	Population statistics for humans and beef cattle 5-12
5.3	Aquatic environmental transport pathways initially considered 5-14

6.1	Annual release rates (Ci) used in the dose equivalent and health effects computations for active uranium mines	6-2
6.2	Annual release rates (Ci) used in the dose equivalent and health effects computations for inactive uranium mines	6-4
6.3	Annual working level exposure from radon-222 emissions from model uranium mines	6-6
6.4	Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model average surface uranium mine	6-7
6.5	Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model average large surface uranium mine	6-8
6.6	Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model average underground uranium mine	6-9
6.7	Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model average large underground uranium mine	6-10
6.8	Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model inactive surface uranium mine	6-11
6.9	Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model inactive underground uranium mine	6-12
6.10	Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a hypothetical in situ uranium solution mine	6-13
6.11	Individual lifetime fatal cancer risk for one year of exposure and estimated additional fatal cancers to the regional population due to annual radioactive airborne emissions from model uranium mines	6-15

6.12	Individual lifetime fatal cancer risk due to lifetime exposure to radioactive airborne emissions from model uranium mines	6-16
6.13	Genetic effect risk to descendants for one year of parental exposure to atmospheric radioactive airborne emissions from model uranium mines	6-17
6.14	Genetic effect risk to descendants for a 30-year parental exposure to atmospheric radioactive airborne emissions from model uranium mines	6-18
6.15	Percent of the fatal cancer risk for the maximum individual due to the sources of radioactive emissions at model uranium mines	6-20
6.16	Percent of the fatal cancer risk for the average individual in the regional population due to the sources of radioactive emissions at model uranium mines	6-21
6.17	Percent of fatal cancer risks due to radon-222 daughter concentrations at model uranium mine sites	6-22
6.18	Percent of the fatal cancer risk for principal nuclides and pathways due to radioactive particulate and Rn-222 emissions at model uranium mines.	6-23
6.19	Natural background concentrations and average urban concentrations of selected airborne pollutants in the United States	6-27
6.20	Combustion product concentrations at the site of the maximum individual with comparisons	6-29
6.21	A comparison of the airborne concentrations of nonradioactive gases at the hypothetical in situ leach site with threshold limit values	6-30
6.22	Stable trace metal airborne concentrations at the site of the maximum individual	6-32
6.23	Comparison of stable trace metal airborne concentrations at the location of the maximum individual with natural background concentrations and average urban concentrations of these airborne pollutants	6-33

6.24	Comparison of trace metal airborne concentrations at the site of the maximum individual with threshold limit values (TLV's) in the workroom environment adjusted for continuous exposure to the general public	6-34
6.25	Annual radiation dose equivalent rates due to aquatic releases from the New Mexico model underground mine	6-36
6.26	Annual radiation dose equivalent rates due to aquatic releases from the Wyoming model surface mine	6-37
6.27	Individual lifetime fatal cancer risk and committed fatal cancers to the population residing within the assessment areas	6-38
6.28	Genetic risks to succeeding generations of an individual and committed genetic effects to descendants of the present population residing within the assessment area	6-41
6.29	Comparison of nonradiological waterborne emissions from uranium mines with recommended agricultural water quality limits	6-43
6.30	Estimated lifetime risk of fatal lung cancer to individuals living in homes built on land contaminated by uranium mine wastes	6-45
6.31	Gamma radiation anomalies and causes	6-49
7.1	Distribution of United States uranium mines by type of mine and state	7-2
7.2	Sources of contaminants at uranium mines	7-3
7.3	Concentration of contaminants in waste rock (overburden), ore, and sub-ore	7-6
7.4	Summary of harm from radioactive airborne emissions of model uranium mines :	7-11
7.5	Percent additional lifetime fatal cancer risk for a lifetime exposure to the individual and the percent additional cancer deaths in the regional population per year of exposure estimated to occur as a result of uranium mining	7-12

7.6	Summary of the fatal cancer risks caused by radioactive aqueous emissions from model uranium mines	7-14
7.7	Estimated lifetime risk of fatal lung cancer to the average person living in a home built on land contaminated by uranium mine wastes	7-16
7.8	Summary of contaminant loading and stream water quality from a model surface uranium mine	7-20
7.9	Summary of contaminant loading and stream water quality from a model underground uranium mine	7-21

Appendix A

A.1	Federal laws, regulations, and guides for uranium mining	A-1
-----	--	-----

Appendix D

D.1	State laws, regulations, and guides for uranium mining .	D-1
-----	--	-----

Appendix E

E.1	Active uranium mines in the United States	E-1
-----	---	-----

Appendix F

F.1	Inactive uranium mines in the United States	F-1
-----	---	-----

Appendix G

G.1	Uravan and Jamestown areas	G-14
G.2	Inactive uranium mine sites surveyed in New Mexico .	G-21
G.3	Status and location of uranium mines in Texas. . .	G-25
G.4	Trace elements and radionuclides in water in the south fork of Box Creek drainage at UNC Morton Ranch lease .	G-35
G.5	Radionuclides and trace metals in sediments in the south fork of Box Creek at UNC Morton Ranch lease . . .	G-36
G.6	Radionuclides and trace metals in soils near the 1601 open pit mine, UNC Morton Ranch lease, Wyoming . .	G-37

		Page
G.7	Radionuclides and trace metals in soil profiles at the open pit mines, UNC Morton Ranch lease, Wyoming	G-38
G.8	Radionuclides and trace metals in sediments from the drainage of the San Mateo mine and from San Mateo Creek, New Mexico	G-40
G.9	Radium-226 and trace elements in water from San Mateo Creek near San Mateo mine discharge point	G-40

Appendix H

H.1	Characteristics of the sub-basin containing the model mines	H-2
H.2	Seepage and outflow calculations for the Wyoming model mine drainage system	H-6
H.3	Characteristics of the sub-basin hydrographic unit in the model underground uranium mine area	H-8
H.4	Seepage and outflow calculations for the New Mexico model mine area drainage system	H-11

Appendix J

J.1	Aquatic environmental transport pathways examined	J-6
J.2	Characteristics of the generic sites	J-20
J.3	Stream data for Valencia County	J-23
J.4	Estimation of meat production in Valencia County for 1977	J-25
J.5	Estimates of meat production in Converse County, Wyoming for 1976	J-26
J.6	Annual radionuclide release rates to streams for active uranium mines	J-29
J.7	Freshwater fish concentration factors	J-29
J.8	Normalized human intake rate factors for radionuclide uptake via plant root systems	J-31
J.9	Irrigated land usage	J-31

J.10	Soil removal rate constants and radioactive decay constants	J-33
J.11	Milk and beef concentration factors	J-33
J.12	Dose equivalent conversion factors	J-36
J.13	Health effects conversion factors for internal pathways.	J-37
J.14	Health effects conversion factors for external pathways.	J-37

Appendix K

K.1	Characteristics of the generic sites	K-1
K.2	Animal and vegetable crop distribution for use with AIRDOS-EPA	K-3
K.3	Sources of food for the maximum individual (percent)	K-4
K.4	Selected input parameters to AIRDOS-EPA	K-5
K.5	Selected terrestrial pathway parameters by radionuclide.	K-7
K.6	Effective radioactive decay constants	K-8

Appendix L

L.1	Radionuclide dose rate and health effect risk conversion factors used in uranium mine assessments	L-4
L.2	Additional input data used by DARTAB in the health impact assessment of airborne emissions	L-21
L.3	Example input data file for DARTAB	L-22
L.4	Maximum individual fatal cancer risk for one year of exposure to atmospheric radioactive emissions from model uranium mines	L-23
L.5	Fatal cancer risk to an average individual in the regional population for one year of exposure to atmospheric radioactive emissions from model uranium mines	L-24
L.6	Fatal cancer risk to the population for one year of exposure to atmospheric radioactive emissions from model uranium mines	L-25

L.7	Genetic effect risk to descendants of maximum exposed individual for one year of parental exposure to atmospheric radioactive particulate and Rn-222 emissions from model uranium mines	L-26
L.8	Genetic effect risk to descendants of average individual of the population for one year of parental exposure to atmospheric radioactive particulate and Rn-222 emissions from model uranium mines	L-27
L.9	Genetic effect risk to descendants of the regional population for one year of parental exposure to atmospheric radioactive particulates and Rn-222 emissions from model uranium mines	L-28

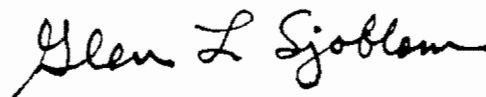
FOREWORD

The Uranium Mill Tailings Radiation Control Act of 1978 stipulates that "...the Administrator, in consultation with the Commission, shall provide to the Congress a report which identifies the location and potential health, safety, and environmental hazards of uranium mine wastes together with recommendations, if any, for a program to eliminate these hazards." It is our understanding that the intent of Congress was to determine if remedial actions similar to those for uranium mill tailings are required for mine wastes.

The report was prepared by the Office of Radiation Programs and addresses potential health effects caused by air emissions, water effluents, and solid wastes at active and inactive uranium mines. It is probably the single most comprehensive report on the subject. The effects from other mining activities such as exploration, site preparation, and in situ leaching were evaluated in proportion to their potential significance and the amount of available information about them. Comments on this report from the uranium mining industry, States, and the Nuclear Regulatory Commission have also been considered.

The conclusions and recommendations are in the Executive Summary and Chapter 7 of the report. The principal findings of this report are as follows:

1. No problems were identified that require Congressional action.
2. Standards are probably needed to control human exposure from radioactive emissions from underground uranium mines. We have proposed a standard for underground uranium mines under the Clean Air Act program to develop radionuclide standards.
3. Regulations should be considered for maintaining the control of solid wastes at active uranium mines to prevent off-site use and to minimize the health risks from these materials. This is part of the overall agency consideration of mining wastes and is being carried out under the auspices of the Solid Waste Disposal Act.
4. The report also identifies additional studies that are needed to completely elucidate the potential for local adverse effects as a result of possible misuse of the mine waste materials in construction of buildings. Preliminary reports of field studies by EPA identifying possible sites at which mine wastes may have been utilized in construction or around buildings have been sent to the States for follow-up studies.



Glen L. Sjoblom, Director
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SECTION 1
INTRODUCTION

1.0 Introduction

1.10 Purpose

This report was prepared in response to Section 114(c) of Public Law 95-604 dated November 8, 1978 (USC78). This Section of the Law stipulates that, "Not later than January 1, 1980, the Administrator, in consultation with the Commission, shall provide to the Congress a report which identifies the location and potential health, safety, and environmental hazards of uranium mine wastes together with recommendations, if any, for a program to eliminate these hazards." The purpose of this report is to comply fully with this request, as accurately and completely as available information will permit.

1.1.1 Contents

This volume has seven major sections. The content of each section is described generally below:

Section 1: Brief reviews of predicted future uranium production requirements; descriptions of methods of extracting uranium from the earth; and presently enforced standards and regulations governing uranium mining.

Section 2: A description of the active and inactive uranium mine inventory with a discussion of its limitations. The actual mine listings are presented in Appendixes E and F.

Section 3: A comprehensive discussion of potential sources of radioactive and stable contaminants to the environment and man from uranium mining operations. Annual release rates of contaminants from the identified sources computed on a generic basis.

Section 4: A description of model underground, surface, and in situ leach mines with operational parameters and source terms. Both active and inactive model underground and surface mines are described.

Section 5: A brief and general discussion of the environment that exists about uranium mines, including vegetation, wildlife, domestic animals, and human populations. The potential atmospheric and aquatic pathways of contaminants from the mines to man are also defined.

Section 6: Computation of individual and population dose equivalents and potential health effects from mine wastes and effluents based on the source terms developed in Section 3, the model mines defined in Section 4, and the pathways described in Section 5. A qualitative description of the environmental effects based on site visits is also presented.

Section 7: A brief summary of the report followed by the conclusions and recommendations.

1.2 Uranium Ore Production and Future Uranium Needs

1.2.1 Past Production

Table 1.1 lists the quantities of ore mined and uranium (U_3O_8) produced in the various uranium mining states between 1948 and January 1, 1979. Two states, New Mexico and Wyoming, have been the source of about 64 percent of the uranium mined in the United States. The Colorado Plateau, which includes parts of New Mexico, Arizona, Colorado, and Utah (see Figure 1.1), has been the largest source area of mined uranium, accounting for about 70 percent of the U_3O_8 production through 1976 (ST78). During this same period, the Gas Hills and the Shirley and Powder River Basins of Wyoming produced about 22 percent of the total U_3O_8 (ST78).

To produce 302,370 MT of U_3O_8 required the mining of 145,811,000 MT of uranium ore during the 31-year period from 1948 to 1979 (DOE79). The average grade of ore, reported as percent of U_3O_8 , was 0.208 percent during this period.

1.2.2 Projected Needs for Uranium

The expected growth in the use of nuclear energy for the production of electric power in the United States during the remainder of this century will require an expansion of the uranium mining industry. However, the magnitude of this expansion is difficult to estimate, because the forecasts that predict the growth of the nuclear power industry differ considerably (AEC74, ERDA75, EPA76, NRC76, NUS76, Cu77, He77, NER77, EW78, Ni78, and St78). However, all forecasts predict a continued growth of the industry. Expansion of the uranium mining industry will be influenced also by decisions regarding fuel reprocessing and commercial utilization of breeder reactors.

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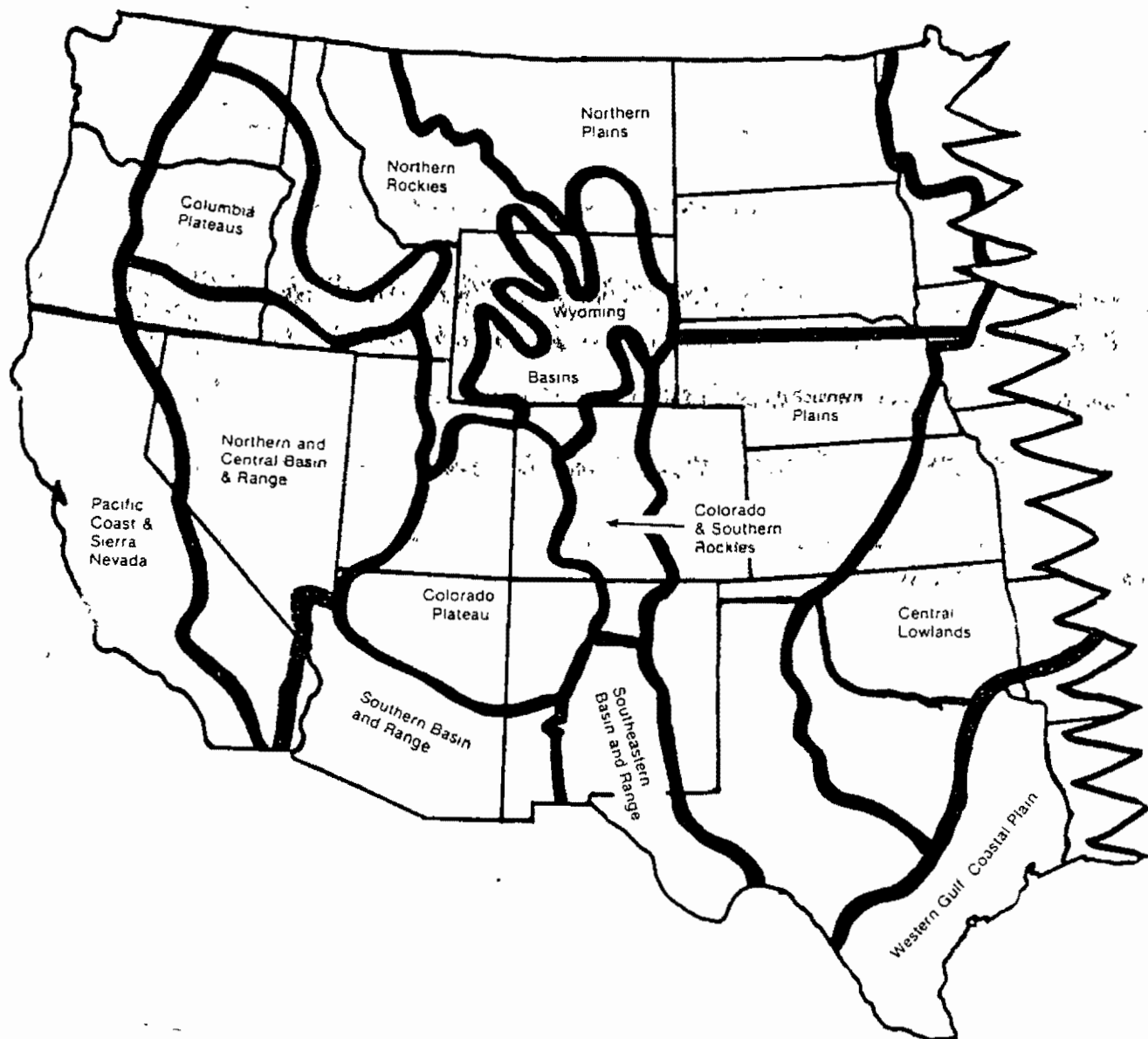


Figure 1 1 Uranium mining regions in the western United States

Table 1.2 gives examples of four typical forecasts. The Nuclear Regulatory Commission's (NRC) projected annual nuclear capacity is far below former predictions. This lower projection, which is in line with the administration's National Energy Plan (NEP77), is believed to be more realistic in view of recent drops in demand for electricity, labor problems, equipment delays, litigations initiated by environmental groups, the absence of a publicly accepted waste disposal program, and concern over nuclear proliferation. The Department of Energy predicts that 293,120 MT of U_3O_8 will be required to provide nuclear generating capacity through 1990 (DOE79). This assumes no uranium or plutonium recycling.

Table 1.3, which gives domestic uranium reserves by state, shows that the reserves are near the areas already mined. Figure 1.2 shows the distribution of \$50 ore reserves by state, and Fig. 1.3 shows reserves by resource region (see Fig. 1.1 for region locations). Future major mining activities probably will be in the same general areas that have already been mined. To obtain the 834,600 MT of \$50 U_3O_8 reserves will require mining about 1.14×10^9 MT of ore with an average grade of 0.073 percent U_3O_8 .

1.3 Overview of Uranium Mining Operations

1.3.1 General

The two major uranium mining methods used in the United States are underground mining and surface (open pit) mining. These two methods accounted for more than 98 percent of the uranium mined in the United States in 1971 (AEC74). This has decreased only slightly to about 93 percent in 1978 (DOE79). However, various types of solution mining are currently being tested and probably will be employed commercially more frequently.

Table 1.4 shows the current production capacities of U_3O_8 for the various mining methods. Although underground mines are far more numerous than surface mines, production by the two methods is nearly equal. This is because surface mines have a much larger capacity. During 1978, 305 underground mines accounted for about 46 percent (8,350 MT) of the U_3O_8 production while 63 surface mines produced about 47 percent (8,710 MT) of the U_3O_8 . In situ leaching, heap leaching, mine water extraction, and other alternative methods accounted for the remaining 7 percent (1,270 MT).

Table 1.2 Projected annual nuclear capacity (GWe) in the U.S.

Source	1980	1985	1990	1995	2000
ERDA (ERDA75)	71-92	160-245	285-470	445-790	625-1250
USEPA (EPA76)	80	188	350	578	820
Electric World (EW78)	92	160	194	237	---
NRC (NRC79)(a)	61	127	195	280	380

(a) Schedule assumed for this document.

Note. -- The actual nuclear capacity realized in 1977 was 49 GWe.

Table 1.3 Domestic uranium reserves by state
as of January 1, 1979

State	Ore		U ₃ O ₈ , MT	% Total U ₃ O ₈
	Ore, MT	Grade, % U ₃ O ₈		
New Mexico	482,200,000	0.09	434,000	52
Wyoming	431,300,000	0.06	258,800	31
Texas	83,400,000	0.05	41,700	5
Arizona, Colorado, & Utah	107,300,000	0.07	75,100	9
Others (a)	35,700,000	0.07	25,000	3
Total	1,139,900,000	0.073 (b)	834,600	100

(a) Includes Alaska, California, Idaho, Montana, Nevada, North Dakota, Oregon, South Dakota and Washington.

(b) Weighted average.

Note.--The uranium reserves in this table include ore from which U₃O₈ can be obtained at a forward cost of \$50 per pound or less. Costs do not include profits or cost of money.

Source: DOE79.

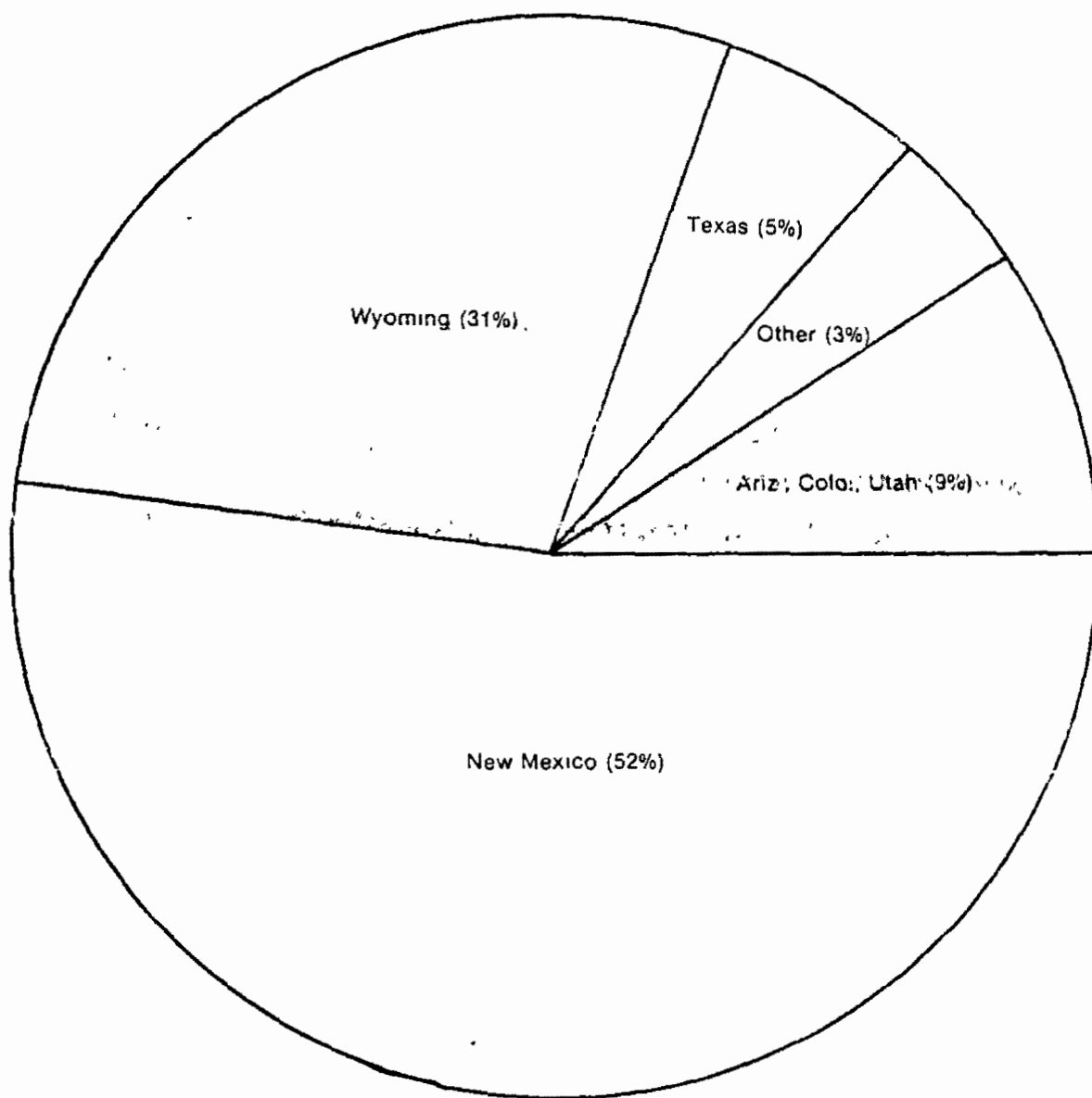


Figure 1 2. The percent of \$5Q U₃O₈ reserves located in the principal mining states. (DOE-79)

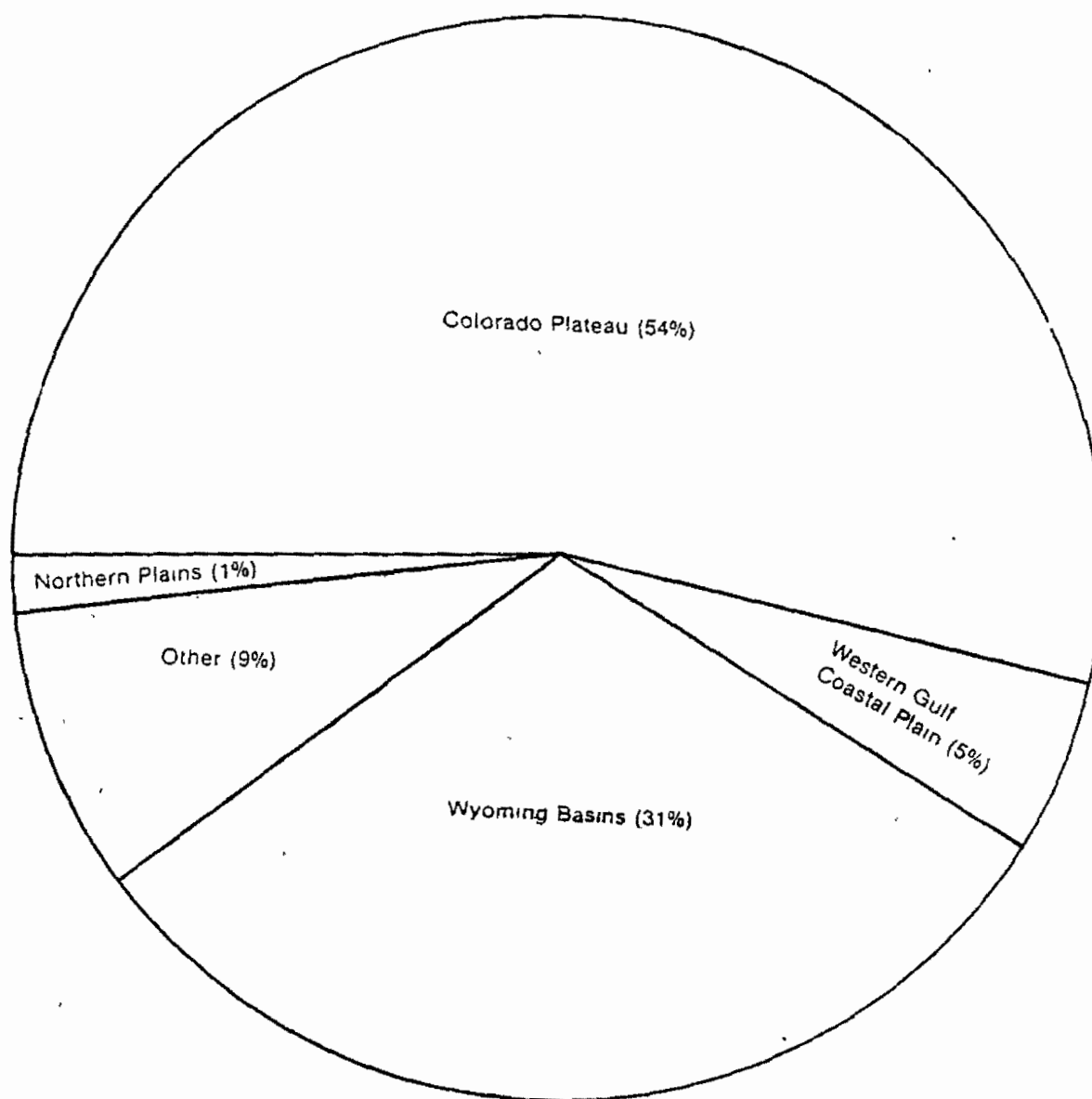


Figure 1.3 The percent of \$50 U_3O_8 reserves located in the various mining regions (DOE 79)

Table 1.4 Quantities of U_3O_8 produced in 1978
by the various mining methods

Mining Method	MT of U_3O_8	Percent ^(b)
Underground Mines (305) ^(a)	8350	46
Surface Mines (63)	8710	47
Other (23): In situ leaching, heap leaching, & mine water	<u>1270</u>	<u>7</u>
Total	18,330	100

(a) The number of mines or sites are given in parentheses.

(b) Rounded to total 100 percent.

Source: DOE79.

Table 1.5 Predicted methods of mining ore reserves

Mining Method	MT of U_3O_8	Percent
Underground Mining	547,000	66
Surface Mining	260,400	31
Other: In situ leaching, heap leaching, & mine water	<u>27,200</u>	<u>3</u>
Total	834,600	

Note.--These are reserves of the \$50 per pound U_3O_8 or less cost category.

Source: DOE79.

Although some mines produced much more than others, one can compute a rough estimate of the average capacity of underground and surface mines as follows. The average grades of ore removed from underground and surface mines in 1978 were reported to be 0.155 percent and 0.120 percent U_3O_8 , respectively (DOE79). Dividing the annual U_3O_8 productions (Table 1.4) of the two mining methods by their respective grades indicates that the 305 underground mines accounted for 5,387,100 MT of ore while 7,258,300 MT were removed from the 63 surface mines. Hence, the average ore production capacities of underground and surface mines are about 1.8×10^4 MT and 1.2×10^5 MT, respectively. From this assessment, the average ore capacity of a surface mine is about seven times that of an underground mine.

The trend during the past few years of an increasing percentage of U_3O_8 being mined underground will continue, because shallow deposits of high grade ore have tended to be surface mined first. Table 1.5, which displays the distribution of \$50 reserves by mining method, shows the continued increase in the proportion of U_3O_8 mined by the underground method. By the Department of Energy predictions, future production from underground mines will more than double that from surface mines. The NRC predicts U_3O_8 production by in situ leaching to peak in 1990 at about 4000 MT/yr and total 76,000 MT by the year 2000 (NRC79). If this prediction is realized, this resource will undoubtedly draw from those assigned to underground and surface mining in Table 1.5. The production of U_3O_8 by heap leaching and mine water extraction is predicted to be relatively small.

It is very difficult to predict how much U_3O_8 will be produced as a by-product from other mineral mines, but by-product production should increase total U_3O_8 production during the next 20 years. Approximately 180 MT of U_3O_8 are currently recovered each year from the phosphoric acid production at wet phosphate plants. The NRC predicts that this will increase to 1800 MT/yr by 1985 and possibly to 7000 MT/yr by the year 2000 (NRC79). If planned leaching facilities are actually built at copper waste dumps at Yerington, Nevada; Butte, Montana; and Twin Buttes, Arizona, to supplement the operating facility at Bingham Canyon, Utah, recovery of U_3O_8 from these operations could reach 900 MT/yr (NRC78). Hence, by-product U_3O_8 production could conceivably account for about eight percent of the required annual U_3O_8 production by the year 2000.

Although the depth of the ore deposit is the fundamental consideration in selecting the mining method to be applied to a particular ore body, the size and grade of the deposit are also important factors. The shape of the deposit, the overburden rock strength, environmental considerations, and other factors may also influence the selection. Surface mining is generally used for relatively shallow deposits; rarely for those below 400 feet (St78). However, under some conditions, it may be cheaper to mine a small, shallow, high-grade ore deposit by underground methods; whereas, a larger low-grade deposit at a greater depth may be cheaper to mine by surface mining. Because productivity is greater by surface mining, it is generally preferred when conditions are favorable. Other factors must be examined when considering the use of in situ leaching (see Section 1.3.4).

1.3.2. Surface Mining.

The use of surface (open pit) mining methods is most prevalent in the Gas Hills Region and the Shirley and Powder River Basins in Wyoming, the Laguna District of New Mexico, the coastal plains in south Texas, and some areas of Colorado and Utah (St78). Fig. 1.4 illustrates a typical open pit mine.

In surface mining, an open pit is dug to expose the uranium deposits. After the topsoil is removed and stockpiled nearby, the overburden is removed by the method best suited to the nature of the rock. If the rock is easily crumbled, it is removed by tractor-mounted ripper bars, bulldozers, shovels, or pushload scrapers; if it is not, blasting and drilling are required. The broken rock is then trucked to a nearby waste dump. Occasionally, dikes and ditches are constructed around these waste piles to collect runoff and divert it to sedimentation ponds. Overall, an area of a hundred or more acres may be covered by stored overburden wastes (AEC74).

As mining progresses, the overburden is used as it is removed to backfill mined out areas of the pit. When an area is completely backfilled, it is graded to conform to the surrounding topography and to restore the natural drainage patterns. The area is then covered with topsoil and seeded to blend with the natural terrain. Most of the older surface mines were not backfilled (see Section 3.7.1), and neither are many of the currently active surface mines.

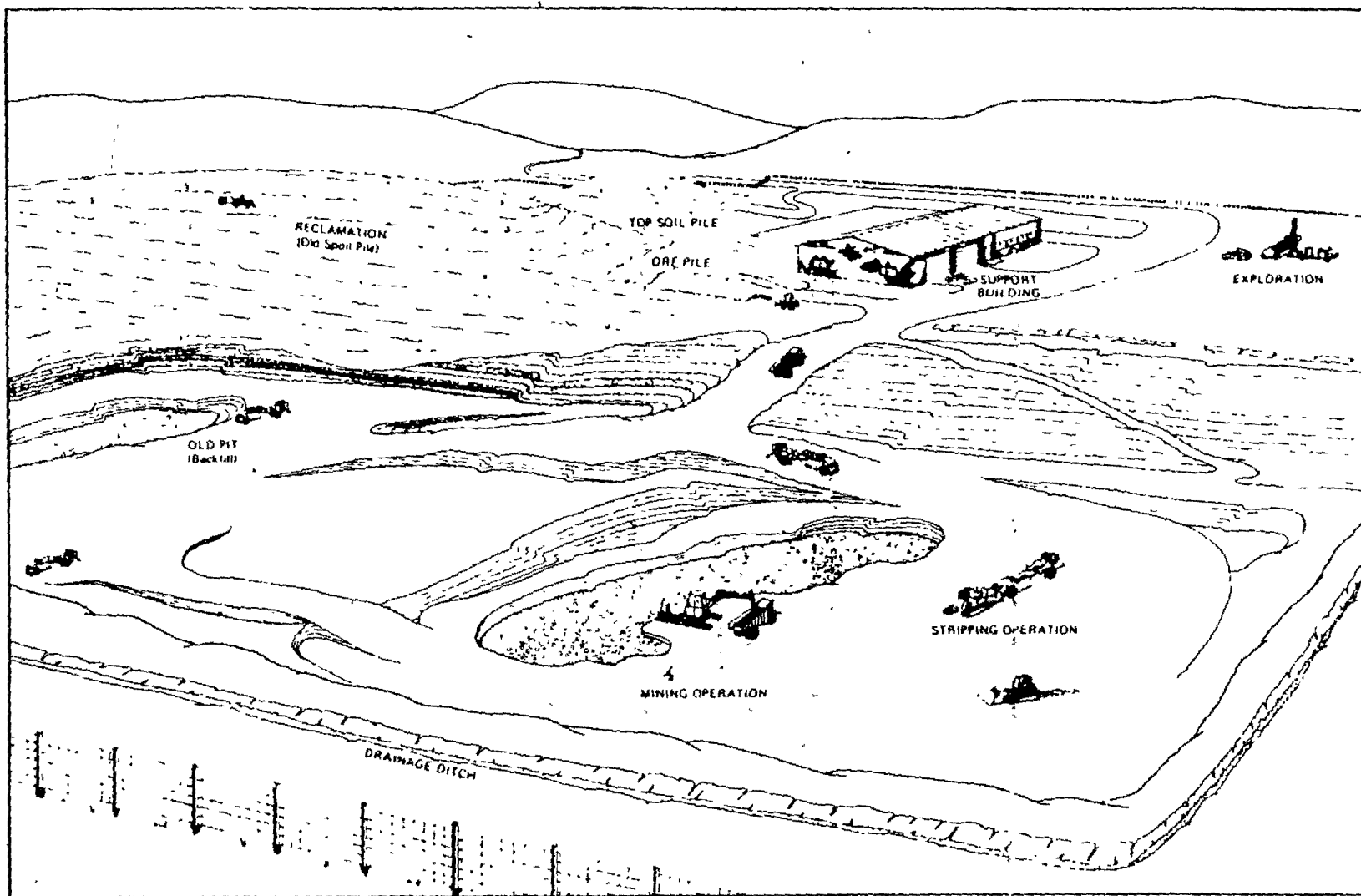


Figure 1 4 Artist's conception of open pit mining operation and support facilities (TVA78a)

Contact with the ore zone is determined by gamma and x-ray detection instruments, usually Geiger-Muller counters that have been calibrated to indicate the uranium content of the rock. Since uranium is usually in sandstone formations, the ore is easily removed. Large backhoes and front-end loaders are often used to remove ore that has been loosened with tractor-mounted rippers. Large ore trucks carry the ore from the pit to stockpiles at the mine or the mill. Uranium ore is usually stockpiled by grade, e.g., high-grade, average-grade, low-grade. Sub-ore grade rock is also usually stored separately in piles. This is rock that contains U_3O_8 at a concentration below what the mill will now accept, but which might later be worth recovering.

Drifts (small tunnels) are sometimes driven into the pit wall to recover small, narrow ore pods. The drifts are generally short, sometimes less than 30 meters. The mining techniques in these drifts are like those used in underground mining (see Section 1.3.3).

Surface mining requires a network of roads from and around the pit area and to the mill. Heavy vehicles operating on these roads, and the digging itself, produce a certain amount of rock and ore-dust. However, the dust can be kept down by routinely sprinkling the roads with water or using other dust suppressants. Treated water from the sedimentation ponds is sometimes used for this purpose.

The ratio of overburden to the ore produced in an open pit mine can vary from 10:1 to as high as 80:1 (St78). One source has estimated the average ratio as 30:1 (Le77). A recent study of eight large open pit uranium mines reported a ratio of 77 (± 36) to 1 (Ni79). Since the latter study did not consider the many smaller surface mines where the overburden to ore ratio is likely to be smaller than 77 to 1, this report will assume an average ratio of 50:1. Considering that the average ore capacity of an open pit mine is approximately 1.2×10^5 MT/yr (see Section 1.3.1), about 6×10^6 MT of overburden must be removed annually and initially stored on the surface until reclamation procedures can be initiated.

Since most uranium deposits lie below the water table, groundwater must be prevented from flooding the mining area. One method is to surround the pit with several large capacity wells to lower the water table near the pit. This water is discharged directly into the natural surface drainage system, in accordance with the National Pollutant Discharge Elimination System (NPDES) discharge permit issued to the mining company. Water that does collect in the pit (mine sump water) is pumped to a sedimentation pond for solids removal and, if necessary, for subsequent treatment prior to discharge into the natural drainage system. Another mine dewatering procedure often used consists of ditches dug along the interior perimeter of the pit floor to channel the water to sumps located at the lowest levels of the pit floor. Water that collects in the sumps is pumped to one or more sedimentation basins for solids removal, possible treatment, and final discharge into the existing natural drainage system in accordance with water quality standards specified in the NPDES permit. The rates at which mines are dewatered range from $0.28 \text{ m}^3/\text{min}$ to $288 \text{ m}^3/\text{min}$ (AEC74, TVA78a, NRC77a, NRC77b).

Barium chloride, to coprecipitate radium, and a flocculent (an agent causing aggregate formation) to remove other contaminants are usually added to pond water before it is discharged. Water with a high concentration of dissolved uranium is often run through ion exchange columns, and the resin regenerant solution containing the uranium is sent to the mill for processing. The precipitated sludge that collects on the pond bottom consists primarily of ferric and calcium hydroxides, calcium sulfate, and barium sulfate with coprecipitated radium. At some sites, this precipitated sludge is transferred to the mill tailings pond at the end of the mining operation.

A small amount of uncontrolled seepage may occur through the bottom of sedimentation ponds and, depending upon soil permeability and direction of flow, may enter the water table. For example, the seepage rate through the bottoms of two settling ponds totaling 4.9 hectares at one site was less than $0.57 \text{ m}^3/\text{min}$ (NRC77a). In addition, seepage can be reduced by lining the ponds (well-compacted bentonite clay is sometimes used for this purpose) and by the sludge that accumulates on the pond bottom.

During active surface mining operations, a total of several thousand hectares of land area will be disturbed (St78, Th79). When all uranium has been mined and the operation is completed, a pit remains. The walls of the pit may be contoured and allowed to fill with water, creating a small man-made lake.

1.3.3 Underground Mining

Underground mining is much less disruptive to the surface terrain than open pit mining. The surface affected generally involves less than 41 hectares, but the mine may extend laterally underground for more than a mile and at several depths. Figure 1.5 illustrates a typical large, contemporary underground mine.

In underground mining, access to the ore body is gained through one or more vertical shafts, generally sunk to a slightly greater depth than the ore body, or through inclines, declines, or adits, all cut through waste rock. The waste rock is removed to a spoils area that may be, but usually is not, surrounded by a ditch to contain runoff, as discussed above.

The sizes of the accesses vary considerably. The vertical haulage shaft may vary from less than 8 feet in diameter, sufficient to accommodate one small ore skip (a large bucket), to a diameter of 20 feet, which will accommodate dual ore skips as well as a man and material skip. In some cases, the near horizontal accesses are sufficiently large to allow passage of large diesel-powered vehicles.

Underground mines are developed in a way that minimizes the removal of waste rock, resulting in much smaller spoil storage piles than those at surface mines. It is estimated that the ore to waste rock ratio generally ranges from 20:1 to 1:1 (ACE74, Th79). At seven presently active mines, the ore to waste rock ratio ranges from 1.5:1 to 16:1 with an average ratio of 9.1:1 (Ja80). Using the average ratio and the average annual ore capacity of an underground mine (see Section 1.3.1), each year the average underground mine will produce about 2.0×10^3 MT of waste rock that is removed and stored on the surface. Initially all waste rock is transported to the surface, but,

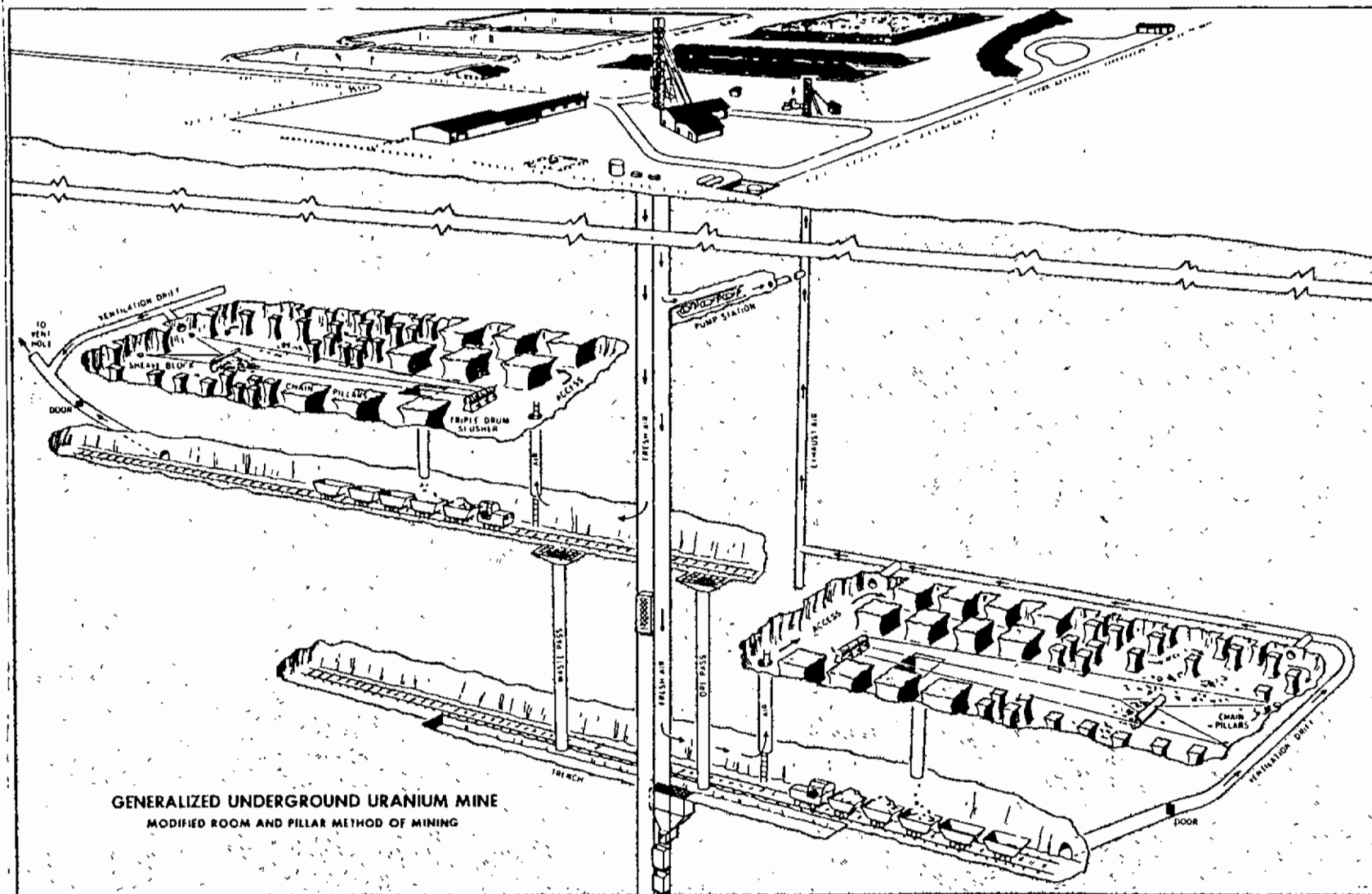


Figure 1.5 Generalized underground mine showing modified room and pillar method of mining (TVA78b)

as mining progresses, it is sometimes transported to mined-out areas of the mine and retained beneath the surface. This practice may diminish as lower grade sub-ore becomes more economical to mill. Since waste rock may contain sub-ore, some waste rock will likely be kept available for milling.

Ore deposits, outlined by development drilling, are followed as closely as possible. When ore lies in narrow, long deposits, drifts are cut through the ore body and raises or stopes are driven from the drift to reach small ore pods. Crosscuts are driven from the haulage drifts when necessary to reach nearby deposits. Large area deposits are commonly mined by the "room and pillar" method. This involves mining out blocks of ore while leaving adjacent pillars of ore or waste as support for the roof. The size of the rooms depends on the roof condition. The roof is usually strengthened by bolts, wire mesh, timbersets, and steel arches. When an area is completely mined, the ore pillars are removed in a systematic sequence that allows safe retreat.

Ore is usually broken by drilling and blasting. The broken ore is removed and transferred to mine rail cars. The ore is then carried by rail cars or wheeled vehicles either directly to the surface or to a skip at the bottom of the haulage shaft and lifted to the surface. Haulage in large area mines is often accomplished by large diesel-powered loaders, haulers, and trucks. When the ore is sufficiently soft, it may be removed with continuous mining machines instead of drilling and blasting techniques; however, most ore bodies are too small and irregular to mine economically this way.

Ventilating systems are required in underground uranium mines to remove blasting fumes and radon-222 ($Rn-222$) that emanates from the ore and mine water and to control temperature. Fresh air is usually forced down the main haulage shaft and along the main haulage drifts to the working areas. The mine air is exhausted through ventilation shafts to the surface. The ventilation air is diverted from inactive areas of the mine to reduce air contamination. Inactive areas are usually sealed with airtight bulkheads to prevent radon gas in those areas from circulating. The ventilation rate should be sufficient to maintain the radon daughter concentration of the mine air at, or below, levels that meet federal and state occupational exposure stan-

dards. The rate will vary depending upon mine size (volume), grade of exposed ore, size of the active working areas, rock characteristics (diffusion rate of Rn-222), effectiveness of bulkhead partitions, atmospheric pressure, and other factors. Ventilation rates in active mines vary from a few hundred m^3/min to over a hundred thousand m^3/min . For example, the ventilation rates for seven uranium mines in the Grants, New Mexico area ranged from 4.4×10^3 to $1.1 \times 10^4 \text{ m}^3/\text{min}$, with an average of $7.4 \times 10^3 \text{ m}^3/\text{min}$ (Ja79).

Because ore bodies often lie in or beneath major aquifers, dewatering operations similar to those practiced in surface mining are required. These operations commence during the initial shaft-sinking process and may continue throughout the working life of the mine. Water is pumped from wells that are driven into the water-bearing strata near the mining operation and discharged either directly into the natural surface drainage system, in accordance with an NPDES permit, or to settling ponds. Water that collects in the mine is diverted to sumps and pumped to a settling pond. The impounded mine water is treated similarly to that described above at surface mines (see Section 1.3.2). The discharge of water from these ponds is in accordance with water quality standards specified in the NPDES permit. (Note.--About one-half of the active New Mexico mine discharges have NPDES permits that are presently under adjudication and, therefore, are not necessarily in accord with discharge limits [Pe79a].)

1.3.4 In Situ Leaching

In situ leaching has less adverse impact on the environment than conventional uranium mining and milling methods. It also may permit economical recovery of currently unrecoverable low-grade uranium deposits (NRC78). Though in situ leach mining currently produces only a small amount of the annual U.S. output of U_3O_8 , variations of this technique are being widely tested for uranium extraction and have potential for becoming commercially significant (La78, NRC78, TVA78b, Ka78). Table 1.6 lists in situ leaching operations for uranium as of January 1, 1978. The operations are concentrated on the coastal plain of southwest Texas and in the Wyoming basin regions. Most commercial sized operations are in southern Texas, where recent expansion is expected to increase the production of U_3O_8 by this

technique to about 900 MT annually (TVA78b). Two Texas sites alone, Bruni and Lamprecht, are expected to produce annually 110 and 230 MT of U_3O_8 , respectively (Wy77). A number of projects are currently testing the effectiveness of the in situ leaching technique. Though these studies usually last about 18 months to 3 years, some feasibility tests require up to 6 years before expanding to full or commercial scale operations (La78). Excellent reviews of this mining method are available (La78, Ka78).

Uranium extraction by in situ leaching probably will not be restricted to one or two geographical areas. Uranium deposits potentially suitable for mining by this method are prevalent in almost all of the established uranium mining areas in the United States. Uranium deposits are potential candidates for in situ mining, if they meet the following criteria: (1) the ore deposit is located in a zone saturated with water; (2) the ore deposit lies above and preferably between geological layers impervious to water; (3) the deposit is adequately permeable to water; and (4) the uranium in the ore deposit is in a leachable state. Colorado and New Mexico already have in situ leaching activities at the pilot scale, and the mining industry has inquired about additional pilot-scale research and development sites in South Dakota, Arizona, Utah, and Montana (La78).

In the in situ leaching method, a leaching solution (lixiviant) is injected through wells into the uranium-bearing ore body. It forms chemical complexes with the uranium, which dissolves in the solution. Production wells bring the uranium-bearing solution to the surface where the uranium is extracted. The barren lixiviant can then be reconstituted and reused. To control groundwater flow, the production (pumped) well operates as a sump or pressure sink in the formation, which produces a flow of groundwater and lixiviant from the injection wells to the production well. Also, some of the barren lixiviant is not reinjected. This reduces the water level in the well field, allowing groundwater to migrate into the mining zone. This inflow prevents the flow of the lixiviant away from the field area.

Lixiviants for in situ mining contain salts of anions (negatively charged chemical groups), such as sulfate, carbonate, bicarbonate, and ammonium, that form stable aqueous complexes with hexavalent (positively

Table 1.6 Summary of current in situ leaching operations as of
January 1, 1978

Name	Location	Well (a) Pattern	Scale of (b) Operation	Flow Rate (c) (m ³ /min)
Sundance Project	Crook County, WY	5-SP	RD-PS	ND ^(d)
Red Desert Site I	Sweetwater County, WY	5-SP	RD-I	ND
Red Desert Site II	Sweetwater County, WY	5-SP	RD-PS	ND
Charley Site	Johnson County, WY	5-SP	RD	ND
Highland Site	Converse County, WY	7-SP	RD-C	4.54
Double Eagle Site	Carbon County, WY	5-SP	RD	ND
North Rolling Pin Site	Campbell County, WY	5-SP	RD-I	ND
Collins Draw Site II	Campbell County, WY	5-SP	RD	0.38-0.57
Bear Creek Site	Converse County, WY	5-SP	RD-I	ND
Nine Mile Lake Site	Natrona County, WY	5-SP	RD-PS	0.38
Red Desert Site	Sweetwater County, WY	5-SP	PS	0.38
Irigaray Site	Johnson County, WY	ND	C	6.06
Site No. 1	McKinley County, NM	4-SP	PS-I	ND
Site No. 2	Sandoval County, NM	4-SP	PS-I	ND
Crownpoint Project	McKinley County, NM	4-SP	RD	ND

Table 1.6 Summary of current in situ leaching operations as of January 1, 1978 (continued)

Name	Location	Well (a) Pattern	Scale of (b) Operation	Flow Rate (c) (m ³ /min)
Grover Site	Weld County, CO	5-SP	PS-C	0.76
Palangana Dome Site	Duval County, TX	ND ^(d)	C	ND
O'Hern Site	Duval County, TX	ND	C	ND
Bruni Site	Duval County, TX	ND	C	ND
Lamprecht Site	Bee County, TX	ND	C	0.76
Zamzow Site	Live Oak County, TX	ND	C	ND
Boots/Brown Site	Live Oak County, TX	ND	C	ND
Clay West Site	Live Oak County, TX	ND	C	ND
Burns Ranch Site	Live Oak County, TX	ND	C	ND
Moser Site	Live Oak County, TX	ND	C	ND

(a) Well pattern: 5-SP indicates one or more 5-spot pattern(s), etc. See Figure 1.6.

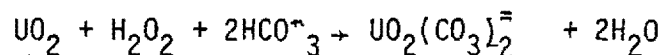
(b) Given are past or present operations - planned future operations: RD - research and development, PS - pilot scale, C - commercial scale, I - presently inactive.

(c) Flow rate of leachate to processing plant in m³/min.

(d) No data.

Source: La78; Du79.

charged in the +6 state) uranium. An oxidant, such as air, oxygen, hydrogen peroxide, sodium chlorate, sodium hypochlorite, or potassium permanganate, is added to oxidize the uranium to the hexavalent state. For example:



Unfortunately, there is no lixiviant specific for uranium. Consequently, other minerals commonly associated with uranium deposits, such as iron, selenium, vanadium, molybdenum, and arsenic, may also be dissolved. This tends to contaminate the leach solution and deplete the lixiviant. Lixiviant agents and their concentrations are selected to maximize uranium recovery and minimize undesirable secondary reactions. Acidic solutions (pH 2) are avoided because they are less selective. Neutral or basic lixiviants (pH 6-10), such as ammonium or sodium carbonate or bicarbonate, are often used.

Many variables affect the accumulation of trace elements in leaching solutions, particularly the chemical and physical nature of the host formation. Table 1.7 illustrates relative contaminant levels in the two lixiviant types in a laboratory experiment. Except for Ra-226, significantly greater trace element concentrations occur in the acid lixiviant; the total dissolved solids is about eight times higher than in the alkaline solution. Hence, it would be necessary to bleed much larger volumes of acidic lixiviant from the system prior to reinjection in order to maintain acceptable levels of these undesirable constituents. Large volumes of liquid wastes containing higher toxic metal concentrations are generally produced when acidic lixiviants are employed. Also, because calcium minerals are abundant in geologic strata and carbonate minerals are highly soluble in acid solutions, particularly calcium carbonate, large amounts of calcium accumulate in recirculated acid lixiviant, and they must be removed by a purification process prior to reinjection. However, acid lixiviants leach more rapidly than alkaline ones, yield higher uranium recoveries -- about 90 percent with sulfuric acid compared to 60 to 70 percent with a bicarbonate solution -- and generally extract less radium (Wy77).

The number of wells, their spacing, and their pattern depend upon the

size and hydrologic characteristics of the formation. Figure 1.6 shows diagrams of some common well patterns. Several hundred injection wells with several recovery wells may be employed. Well spacing may vary from 10 to 60 m. In addition, a number of monitoring wells are driven a short distance from the well field to detect any excursion of lixiviant from the leach field. A commercial-size operation may require a well field area of 20 hectares or more (TVA78b).

The pregnant (containing uranium) leachate from the production wells is filtered through a sand filter to remove suspended particulates, then passed through a surge tank (storage reservoir) to ion-exchange resin beds that selectively remove the uranium complex. The uranium is washed from the resin beds, precipitated, filtered, dried (at most sites), and packaged.

Some processes of solution mining produce liquid and solid wastes. The volume of liquid wastes produced is much smaller, per weight of U_3O_8 produced, than that from the dewatering activities of conventional mining methods. There is also no waste rock. Residues obtained from drilling are a solid waste. Those that traverse the ore zone will contain some uranium ore. If calcium or sulfate control of the lixiviant is necessary, additional solid wastes are impounded in waste ponds under a liquid seal to minimize atmospheric dispersion. Precipitation compounds will also be produced as evaporation concentrates the impounded waste solutions.

Liquid waste streams include lixiviant, filter and resin washes, resin eluant bleed, and water used in cleaning the injection wells. The total production rate of these waste streams may vary between 0.19 to 0.38 m³/min (Ka78, Wy77, TVA78b). At most sites, all liquid wastes flow to waste ponds and evaporate. Pond size depends on the flow rate of the wastes and the evaporation rate. The pond bottoms are usually lined with clay, asphalt, or a continuous plastic sheet to minimize the seepage rate, although some seepage may inadvertently occur. Deep-well injection is also used, principally in Texas, to dispose of liquid wastes from in situ leaching (Du79).

Table 1.7 Trace metal concentrations of recirculated acid and alkaline lixiviants

Trace Metal	Concentrations, mg/l	
	Acidic ^(a)	Alkaline ^(b)
Arsenic	<0.05	<0.05
Chromium	0.15	0.07
Cobalt	0.2	NR ^(c)
Copper	1.0	0.04
Iron	25.4	0.6
Lead	0.7	0.2
Manganese	1.2	NR
Molybdenum	NR	0.9
Nickel	0.6	0.06
Selenium	NR	1.6
Strontium	3.7	1.5
Vanadium	1.0	NR
Zinc	4.3	0.1
Zirconium	3.3	0.9
Radium-226 ^(d)	390	1750
TDS ^(e)	7.8	1.0

(a) Composition - 5 g/l H_2SO_4 and 0.1 g/l $NaClO_3$.

(b) Composition - 8 g/l NH_4HCO_3 and 1 g/l H_2O_2 .

(c) NR - Not Reported.

(d) Units - pCi/l.

(e) Total dissolved solids in grams.

Source: Ka78.

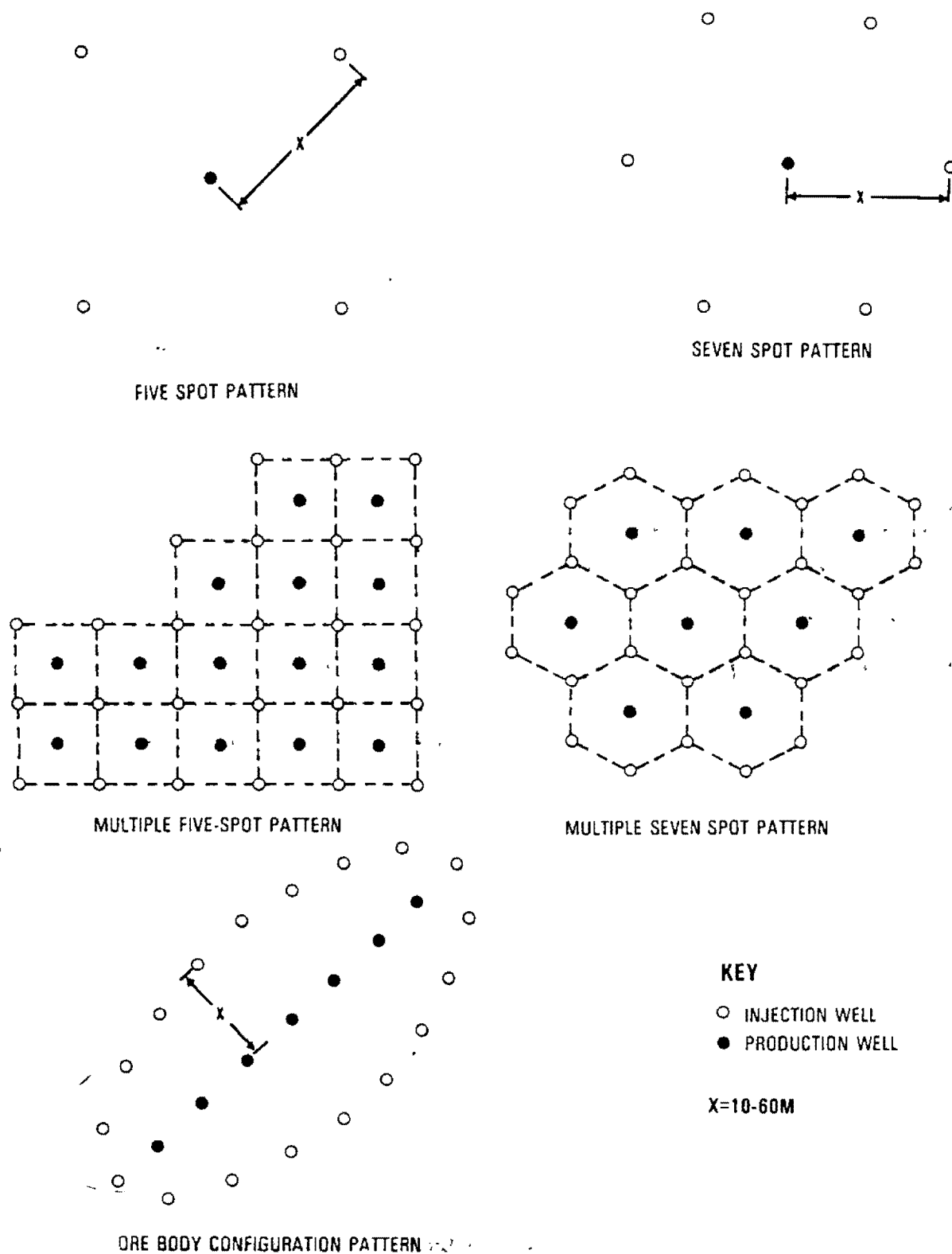


Figure 1 6 Diagrams of some common injection-recovery well patterns used in uranium in-situ leach mining.

Atmospheric emissions from in situ leaching include Rn-222 that is vented mainly from the pregnant lixiviant surge tanks and particulate matter that may escape from the scrubbers (exhaust filters) of the yellowcake (uranium product) drying and packaging units. Radon-222 emanation from the waste ponds is probably negligible, since the sediment which contains the radium remains submerged and little Radon-222 will diffuse through the water and escape to the atmosphere.

When the mining operation is completed, the water volume of the leach zone will be restored to limits set by regulatory agencies. The primary method of aquifer restoration is flushing the zone with groundwater by pumping from the production wells and/or the injection wells. This process may produce up to $1.70 \text{ m}^3/\text{min}$ additional liquid wastes that contain a high concentration of dissolved solids (NRC78). This contaminated water passes through an ion-exchange unit and then discharges to the waste ponds. The barren effluent can be further treated by desalination and reinjected to the formation. Also, a barium chloride solution can be injected into the leach zone to coprecipitate radium from the aquifer water prior to the groundwater sweeps.

1.3.5 Other Mining Methods

1.3.5.1 Heap Leaching

Ore of a grade too low to be economically extracted in the mill operation is sometimes treated by heap leaching. In this process, the low grade ore is placed in large, rectangular, open-air piles on a specially prepared pad. To construct the heap-leach pad, the topsoil is first removed and the cleared area graded to a 2 percent to 3 percent slope. The graded area is then covered by a plastic sheet. Perforated plastic pipe is placed on the plastic parallel to the slope and covered with approximately 30 cm of clean, coarse gravel. A collecting trough is formed at the base of the slope and a berm surrounds the pile area.

When the ore is dumped on the pad, large solution reservoirs are formed on the top. Acidic or alkaline mine water is pumped into the reservoirs and

percolates through the pile. The percolated water is collected in the trough and recirculated until the concentration of uranium in solution is sufficient to be economically extracted. The leaching process may require up to six months to recover approximately 80 percent of the uranium in the ore (NRC77a). The heap-leach pile at one mine contained an annual accumulation of approximately 360,000 MT of low-grade ore (NRC77a). The pile measured 300 m x 90 m x 7.6 m with solution reservoirs of 22 m x 90 m x 1.5 m.

After leaching operations are completed, the leached pile is neutralized with lime to a pH of about 7. The site is then contoured to blend with the surrounding terrain, covered with layers of subsoil and topsoil, and seeded to control wind and water erosion.

Because necessary information is unavailable and the contribution of heap leaching to the total uranium production is very minor and not expected to become significant (NRC79), an assessment of the environmental impact of heap-leached piles has not been conducted. However, the NRC has recently concluded that, although the hazard of tailings produced by heap leaching will be much less than the hazard of tailings at conventional uranium mills, the same tailings management and disposal criteria should possibly apply (NRC79).

1.3.5.2 Mine Water Recirculation

At several sites mine water is recirculated to leach "worked-out areas" of underground mines (Pe79b). In the early uranium mining years, ore with less than about 0.15 percent U_3O_8 was not mined. This grade is relatively high compared to present day markets. Consequently, significant quantities of uranium remain in these abandoned areas. Because the roofs of these areas collapsed during the initial mining retreat, this ore is difficult to retrieve by conventional methods. To recover a portion of this uranium, holes are drilled to the top of the collapsed zone and mine water is sprayed from these holes onto the shattered ore. Water for leaching may be sprayed from the mine floor if the abandoned area is accessible to the workers (Pe79a). The oxidized uranium (uranyl ion) is leached by the slightly alkaline mine water, which flows to collection sumps. The enriched water is pumped to a resin ion-exchange unit to extract the uranium, and then it is recycled.

After the available oxidized uranium has been leached, the process is discontinued for a few weeks to allow more uranium to oxidize. Mine water is then circulated again through the ore.

This process increases the recovery of uranium with minor effort and expense, but it contributes little to the total domestic uranium production (NRC79). In addition, the quality of the stored mine water used will be enhanced after passing through the resin ion-exchange unit. Hence, mine water recirculation has little impact on the environment. It was not assessed in this study.

1.3.5.3. Borehole Slurry Mining

Hydraulic borehole slurry mining is a recently proposed technique for extracting uranium ore (Ka78, St78). As the name suggests, this method uses pressurized water to loosen and combine with ore-bearing material to form a watery mixture known as 'slurry' that is transported from the borehole and then conventionally milled. This method could be applied to sandstone deposits at depths of 30 m to about 100 m. By present estimates, yellowcake from ore containing 0.06 percent U_3O_8 and mined at a 60 m depth by this method would cost \$42 per pound (Ka78). This method presently is not as economical as the more conventional methods of uranium mining.

The process consists of drilling a 45-cm diameter hole to approximately 2 m below the uranium-bearing strata. A cutting jet assembly is positioned in the hole at the end of a rigid service column containing conduits for the pressurized water and slurry transport. The slurry pump is placed at the bottom of the hole. The underground mining operation is started with the jet set at the lowest position. The rotating jet cuts material through an arc of somewhat less than 360° for a distance of up to 25 m, depending upon the design of the jet system. The segment of unmined ore acts to support the overlying strata. After the material is removed as a slurry, the jet is raised to the next level of ore and the process is repeated. After milling, the decanted water from the slurry is recycled for slurrying more ore. The tailings from the milling operation are used to backfill the borehole cavities and minimize subsidence.

A 15 m to 25 m radius borehole can be mined in an 8- to 24- hour period (Ka78). Large ore bodies might be mined by drilling, slurring, processing, and backfilling in a systematic pattern that leaves ore in between boreholes for support. These areas could be mined in a second phase after the original boreholes are backfilled.

Borehole mining for uranium is currently only a proposed method with no pilot or commercial scale units in operation. Thus, the possible environmental impact from this process was not assessed in this study.

1.3.5.4. Uranium as a By-Product

The recovery of uranium as a by-product from other mineral mining and milling operations was discussed briefly in Section 1.3.1. Since recovery is basically from the milling operation, any environmental problem that might exist is associated with milling rather than mining. Therefore, it was not assessed in this study.

1.4 Current Applicable Standards and Regulations

1.4.1 Federal Regulations

Health, safety, and environmental hazards associated with uranium mining are regulated by Federal and State laws. This review focuses on laws and regulations applicable to mine operations. Nuclear Regulatory Commission regulations for milling operations apply to in situ leach extraction and are therefore included. Some laws and regulations on exploration rights also cover the environmental impact of mining operations and wastes.

Prior to the National Environmental Policy Act (NEPA) of 1969, there were few regulations protecting the environment of lands not controlled or owned by the Federal Government. Even with NEPA, much Federal authority on environmental problems was unused until recently. This Act established a national policy concerning the environment. Section 102(2) (C) states that every agency of the Federal Government must "include in every recommendation or report on proposals for legislation and other major Federal actions significantly affecting the quality of the human environment, a detailed statement" of the environmental impact of such an action. Major Federal actions

"... includes actions with effects that may be major and which are potentially subject to Federal control and responsibility ... actions include new and continuing activities, including projects and programs entirely or partly financed, assisted, conducted, regulated, or approved by Federal agencies; new or revised agency rules, regulations, plans, policies, or procedures; and legislative proposals (Sections 1506.8, 1508.17) Approval of specific projects, such as construction or management activities located in a defined geographic area. Projects include actions approved by permit or other regulatory decision as well as federal and federally-assisted activities"(40 CFR 1500).

1.4.1.1 Federal Laws, Regulations, and Guides for Protection of Health and Environment

Table 1.8 provides an overview of federal laws and regulations for the protection of health or environment and the administering agencies. Federal agency responsibilities for water use, conservation laws, and exploration and mining rights are indicated in columns 1-4. Laws and regulations for environmental quality and health and safety are indicated in columns 5-10. See Appendix A for an itemized list of the laws and regulations shown generally in Table 1.8.

1.4.1.1.1 Air Quality

Regulations on air quality have been promulgated pursuant to the Clean Air Act (42 U.S.C. 1857 et seq), which includes the Clean Air Act of 1963 (Public Law 88-206) and amendments by the following: Public Law 89-272, Public Law 89-675, Public Law 90-148, Public Law 91-604, Public Law 92-157, Public Law 93-319, Public Law 95-95, and Public Law 95-190. The Environmental Protection Agency establishes National Ambient Air Quality Standards, New Source Performance Standards, and National Emissions Standards for Hazardous Air Pollutants under the Clean Air Act (CAA). Primary standards are set to protect public health and secondary standards are set to protect public welfare from known or anticipated adverse effects.

National Ambient Air Quality Standards (NAAQS) have been established for seven pollutants in 40 CFR 50. The Administrator of EPA is authorized to set emission standards for hazardous air pollutants for which no ambient air quality standard is applicable. Asbestos, beryllium, mercury, and vinyl

Table 1.8 Federal laws, regulations, and guides for uranium mining

Federal Agency	General		Permits		Mining					Health and Safety
	Water Use	Conservation-Preservation Statutes	Exploration Rights	Mining Rights	Environmental Quality			Land Reclam		
					Air	Water Surf	UG		Solids	
Dept. of Int.	X	X	X	X					X	
BIA ^(a)			X	X					X	
BLM ^(a)		X	X	X					X	
USGS ^(a)			X						X	
Dept. of Energy		X	X	X					X	
Dept. of Agr.	X	X							X	
USFS ^(a)			X	X					X	
EPA	X	X					X		X	X
AIR-OAQPS ^(a)					X					
Water										
Surface OWPS ^(a)							X			
Ground OSW ^(a)								X		
Land-OSW ^(a)							X	X	X	X
Radiation-ORP ^(a)					X	X	X	X	X	X
U.S. Army										
Corps of Engrs.	X	X					X		X	
Dept. of Labor		X								
MSHA ^(a)										X
OSHA ^(a)										X
Nuclear Reg. Comm. ^(b)		X		X		X		X	X	X

(a) BIA-Bureau of Indian Affairs
 BLM-Bureau of Land Management
 USGS-United States Geological Survey
 USFS-United States Forest Service

OWPS-Office of Water Planning and Standards
 OSW-Office of Solid Waste
 ORP-Office of Radiation Programs
 MSHA-Mining Safety and Health Administration

(b) OAQPS-Office of Air Quality, Planning and Standards
 Nuclear Regulatory Commission (NRC) regulations and guides for milling do apply to in situ extraction or mining but not conventional surface or underground mining where NRC has no authority.

chloride emission standards are in subparts, B, C, E, and F of 40 CFR 61, respectively. Section 122 of the CAA directed the Administrator to determine whether emissions of radioactive pollutants, cadmium, arsenic, and polycyclic organic matter (such as benzene) into ambient air will cause or contribute to air pollution and endanger public health. If they do, EPA must propose emission standards for them within 180 days after that decision. The EPA has listed radionuclides as "hazardous pollutants" under Section 112 of the Clean Air Act in December 1979 (44FR76738, December 27, 1979). To date, no standards for radionuclide emissions in air have been promulgated.

The particulate concentration values of the NAAQS apply to mining operations. Emissions (including dust) must be controlled to meet the standards. Dust from mining operations was excluded from any air quality impact assessment for prevention of significant air quality deterioration (PSD) (see 43 F.R. 26395). However, as a result of the court decision in *Alabama Power Company v. Costle*, 13 ERC 1225, EPA has proposed amendment of PSD regulations (44 F.R. 51924, September 5, 1979).

The emission of radioactive substances or gases from gaseous release is controlled by NRC regulations 10 CFR Parts 20 and 40 for uranium milling and in situ leaching. The NRC does not have this authority over mining. There are no Federal regulations for radioactive pollution of air from mining at this time. However, MSHA enforces standards for radioactivity in air inside mines (30 CFR 57.5-37 through 57.5-42). Health and Safety standards of MSHA for Metal and Nonmetallic Mine Safety are given in 30 CFR Parts 55, 57, and 58.

1.4.1.1.2 Water Quality

Standards for water quality are promulgated by EPA under the Federal Water Pollution Control Act (FWPCA) of 1948 (as amended) and the Safe Drinking Water Act (SDWA) (as amended). The FWPCA and SDWA regulate surface water quality and groundwater quality, respectively.

The Federal Water Pollution Control Act Amendments of 1972(Public Law 92-500) established that no one has a right, without permit, to discharge pollutants into navigable waters of the nation. The Act provides for the establishment of both water quality standards and effluent limitations. In addition to requiring effluent standards for existing sources, it required EPA to set new source performance standards for uranium mining. The following standards and guidelines apply to uranium mining and milling: Regulations on Policies and Procedures for the National Pollutant Discharge Elimination System (40 CFR 125), Effluent Guidelines - Mining and Processing (40 CFR 116), Effluent Guidelines and Standards for Mining and Processing (40 CFR 436), and Protection of the Environment-Ore Mining and Dressing - Point Source Category (40 CFR Part 440). Table 1.12 lists other pertinent regulations and guides.

The Safe Drinking Water Act primarily protects municipal water systems. Part C of the Act requires that states establish underground waste water injection programs according to EPA regulations. Most mining operations dispose of waste water through surface discharges subject to the NPDES permit program and to the FWPCA. However, if a mine or mill seeks to dispose of polluted water by injection and such injection may endanger public drinking water supplies, then the Safe Drinking Water Act would apply. Finally, EPA will be developing regulations pursuant to Subtitle C of the Resource Conservation and Recovery Act that will provide controls on hazardous uranium mining wastes, including protection of groundwater resources. Section 4004 criteria, promulgated on September 13, 1979, apply to the nonhazardous portion of the wastes.

The NRC's water quality standards for radioactivity in discharges from uranium milling to the environment are in 10 CFR Parts 20 and 40. These would apply to in situ mining licensed by NRC or an agreement state.

1.4.1.1.3 Land Quality

Federal regulations on solid waste disposal and land reclamation specifically for uranium mining wastes are being developed pursuant to the Solid Waste Disposal Act (as amended). The Surface Mining Control and Reclamation Act of 1977 only applies to coal mining. Uranium mining occurs on Federal

lands, where the Departments of Interior and Agriculture require reclamation. A large part of the western states is Federally owned land: Arizona (43 percent), California (45 percent), Colorado (36 percent), Idaho (64 percent), Montana (30 percent), Nevada (87 percent), New Mexico (34 percent), Texas (2 percent), Utah (66 percent), Washington (29 percent), and Wyoming (48 percent). State laws and local zoning ordinances may affect waste disposal. Many states authorize counties to regulate land use outside incorporated areas. Likewise, many states allow cities, towns, and villages to enact zoning ordinances for land use within their boundaries. Thus, mining operations in each state are subject to different reclamation requirements, depending upon land ownership and location.

Regulations for hazardous uranium mining wastes have been proposed by the EPA pursuant to Subtitle C of the Solid Waste Disposal Act as substantially amended by the Resources Conservation and Recovery Act of 1976 (Public Law 94-580). These were published in the Federal Register (43 F.R. 58946-59028) on December 18, 1978. Waste rock and overburden from uranium mining are listed as hazardous wastes, because they contain radioactive substances that meet the definition of hazardous wastes given in Section 1004 (5) of the Act. Special waste standards (Part 250.46-4) were proposed for the treatment, storage, and disposal of overburden and waste rock.

1.4.1.2 Federal Mineral Leasing and Location/Patent Laws

Some Federal regulations govern mineral exploration and mining rights. The Mining Law of 1872 (30 USC §§ 21-50) permits persons to enter public lands to discover, locate, and mine valuable minerals. The law has no provisions for facility siting, surface protection, or reclamation. Free use of water and timber for the mining operation and land for a mill site are ancillary rights granted by the law. Most subsequent mineral leasing laws are similar, designed to provide an orderly system for locating, removing, and utilizing valuable mineral deposits on federally owned and controlled lands. Pursuant to Section 603 (C) of the Federal Land Policy and Management Act of 1976, DOI has proposed specific environmental protection regulations (43 CFR 3800) for mining activities in potential or identified wilderness study areas (44 FR 2620).

1.4.1.2.1 Prospecting and Mining Rights

Consideration of environmental impacts may be required before obtaining the right to prospect or explore. Depending upon land category, prospectors may have to assess the environmental impact of mineral exploration before being permitted to explore. Table 1.9 summarizes these requirements. Prospectors on private lands simply must have permission from the owner of record of mineral estate. On the other hand, Tribal and Indian lands, National Forest System lands, and public lands (not public domain) all have specific approval systems that require exploration plans or other appropriate considerations.

Obtaining rights to mine usually involves the same Government agency involved with prospecting rights. Table 1.10 summarizes applicable Federal laws and regulations.

1.4.1.2.2 Mining and Environmental Plans

Before mining begins certain operating or mining and reclamation plans must be submitted and approved. Table 1.11 summarizes these. The requirements parallel those for prospecting and mining rights.

1.4.1.3 Laws Having Potential Applicability

Federal laws require regulation for quality of air, water, and land. In addition, though their direct influence has not been evaluated in this report, federal laws protecting wildlife and cultural resources could affect uranium mining activities.

Water use is also of potential concern in regard to uranium mining. However, except for in situ mining, uranium mining operations have modest needs for water. In fact, most mines typically dispose of significant quantities from necessary dewatering. Appendix B lists federal water programs and rights activities and the lead agencies administering them; and Appendix C lists Congressionally approved compacts that apportion water. These compacts apportion water to the affected states, and each state in turn allocates its share of the water among intrastate users on the basis of its own system of water rights.

Table 1.9 Requirements to obtain rights to prospect or explore by federal, state and private lands

Land Category	Requirements
Federal:	
Tribal.....	Prospecting permit issued by BIA with consent of tribe. 25 CFR 171.27a. Technical examination of environmental effects of prospecting by BIA, 25 CFR 177.4. Exploration plan submitted to USGS. Approval of plan by USGS required, 25 CFR 177.6. Enforcement of plan by USGS, 25 CFR 177.10.
Allotted Indian....	No specific provisions for prospecting. Procedure for leasing to prospect is same as for mining. If allotted land has been patented, treat same as private land.
Public Domain.....	No restriction on prospecting. Entry under General Mining Law of 1872 (30 USC 22, 43 USC 1744, 43 CFR Part 3810), uranium included, 43 CFR 37461.
Acquired Public....	Prospecting permit from BLM, 43 CFR 3510.0-3 and 3511.2-1. Acquired lands not subject to prospecting permits are listed in 43 CFR 3501.2-1. If acquired land is not under BLM jurisdiction, consent of governmental entity having jurisdiction is required before permit issued by BLM (43 CFR 3501.2-6).
Withdrawn Public...	Public domain land withdrawn for power development is open to entry and location under General Mining Law of 1872, 30 USC 621. Agency having control of withdrawn land reports any objections to mining activity based on land use for which withdrawal was made. If controlling agency recommends stipulations in the permit, they are included (43 CFR 3501.3-1 (a), (c)).
Reserved Public....	Some Federal lands are disposed of with minerals reserved to the Government; e.g., see 43 USC 299, 43 CFR 3814.1, 30 USC 50. For these lands, permit issued by BLM requires conformance with law under which reservation was made, 43 CFR 35013-2(2). For lands reserved or segregated for particular purpose, special requirements may be made by BLM for protection and use of land for purpose that it was reserved or segregated. Leases from Dept. of Energy may be possible under 42 USC 2097.

Table 1.9 (Continued)

Land Category	Requirements
National Forest System...	Public domain lands inside National Forest System boundaries are subject to General Mining Law of 1872, with the following conditions: (a) If Dept. of Agriculture requires operations plan, it must be submitted. Dept. of Agriculture approves plan, 36 CFR 252.1; (b) Operations must minimize environmental impact on surface resources in System lands, 36 CFR 252.8; (c) Surface inspection and securing compliance with plan is responsibility of Dept. of Agriculture, 36 CFR 252.7. Acquired National Forest System land same as Acquired Public Land.
State.....	Lease obtained from appropriate State Agency according to state law.
Private.....	Permission given by owner of record of mineral estate.

Source: San Juan Basin Regional Uranium Study, Working Paper No. 28, Legal Infrastructure Related to Uranium Mining in the San Juan Basin, United States Department of Interior.

Table 1.10 Requirements to obtain rights to mine ore by
federal, state, and private lands

Land Category	Requirements
Federal:	
Tribal.....	Secretary of Interior has general authority for leases, 25 USC 396a. Tribe must approve. Leases given by bid. Approval of Secretary of Interior required, 25 CFR 171.2. Tribe may negotiate lease if Secretary grants permission. Secretary has discretion to reject lease negotiated by Tribe, 25 CFR 171.2. Secretary may issue charter of incorporation to Tribe which may include authority for Tribe to negotiate mining leases without approval.
Allotted Indian....	Leases given by bid. If Secretary of Interior approves, leases may be negotiated by Indian owners, but negotiated lease subject to rejection by Secretary, 25 CFR 172.4 and 172.6. Approval of allottee required. If patented, treat same as private land.
Public Domain.....	No lease required. Location of mineral deposit (staking a claim) after mineral has been discovered, 43 CFR 3831.1 and 3841.3. File locations with BLM and in accordance with 43 USC 1744. Also record in accordance with State law. Obtain patent for land claimed, 30 USC 29, 43 CFR Part 3860. Mill sites may be claimed by location and patenting, 30 USC 42, 43 CFR Subpart 3844. If claim has been patented, treat same as private land.
Acquired Public....	Mineral estate on acquired lands can be leased by BLM, 43 CFR 3501.3-1, subject to exceptions (43 CFR 3501.1-5 and 3501.2-1). Permittee who prospected and discovered is entitled to preference right lease, 43 CFR 3520.1-1(a)(3). BLM leases land which contains valuable minerals on competitive basis, 43 CFR 3520.1-2(a). If land is not under BLM jurisdiction, consent of governmental entity having jurisdiction is required before lease issues.
Withdrawn Public...	For public domain land withdrawn for power development, laws are same as for land in Public Domain, 30 USC 621. If withdrawal does not preclude mining, BLM can lease mineral estate. Agency having jurisdiction of withdrawn land reports any objections to mining activity, based on land use for which withdrawal was made. If controlling agency recommends stipulations in lease, they are included, 43 CFR 3501.3-1(a)(c). Leases from Dept. of Energy on lands withdrawn for DOE use under 42 USC 2097.

Table 1.10 (Continued)

Land Category	Requirements
6. Reserved Public	Some Federal lands are disposed of with minerals reserved to the government; see e.g. 43 USC 299, 43 CFR 3814.1 and 30 USC 50. For these lands, lease issued by BLM requires conformance with law under which reservation was made, 43 CFR 3501.3-2(2). For lands reserved or segregated for particular purpose, special requirements may be made by BLM for protection and use of land for purpose that it was reserved or segregated.
7. National Forest System	Public Domain land inside National Forest System boundaries are subject to General Mining Law of 1872, with the following exceptions: 36 CFR 252.1, (a) If Department of Agriculture requires operations plan, it must be submitted. Department of Agriculture approves plan; (b) Operations must minimize environmental impact on surface resources on System Lands, 36 CFR 252.8; (c) Surface reclamation required, 36 CFR 252.8(g); (d) Inspection and compliance with plan responsibility of Department of Agriculture, 36 CFR 252.7. Acquired National Forest System Land same as Acquired Public Land.
8. State	Leases obtained from appropriate State Agency according to state law.
9. Private	Lease of mineral estate (or total estate) by private negotiation.

Source: San Juan Basin Regional Uranium Study, Working Paper No. 28, Legal Infrastructure Related to Uranium Mining in the San Juan Basin, United States Department of Interior.

Table 1.11 Requirements for mining and environmental plans by federal, state, and private lands

Land Category	Requirements
Federal:	
Tribal.....	Mining plan must be approved by USGS. If lease requires revegetation, the revegetation work is included in mining plan. Mining plan can be changed by mutual consent of USGS and operator, 25 CFR 177.6. BIA evaluates environmental effect of proposed operations and formulates environmental mitigation requirements. BIA consults with USGS, 25 CFR 177.4.
Allotted Indian.....	Same as Tribal Land, 25 CFR 177.1., unless allotted land has been patented. If patented, treat same as private land.
Public Domain	Plan same as acquired public land.
Acquired Public	Geological survey approval of mining plan to mitigate adverse environmental effects for federal leases, 30 CFR 231.10.
Withdrawn Public.....	Stipulations can be put in the lease by the agency for whom the land was withdrawn. These could affect operations but no formal submission of plans required, 30 CFR 231.10.
Reserved Public	Lessee must conduct operations in conformance with such requirements as may be made by BLM. Requirements will conform to purposes for which land was reserved, 43 CFR 350.3-2(b). Approval of mining plan required, 30 CFR 231.10.
National Forest System...	Operations plan submitted to District Ranger, Department of Agriculture, if he deems it necessary, 36 CFR 252.4. Reclamation of surface required under operator's plan, 36 CFR 252.8(g). Compliance with Federal and State environmental laws, preserve scenic values, wildlife, etc., 36 CFR 252.8. District Ranger, Department of Agriculture, inspects and assures compliance with operations plan, 36 CFR 252.7.
State.....	Mine plan filed with and approved by State.
Private	Same as State Land.

Note.--Some states require submission of mining and reclamation plans for all land.

Source: San Juan Basin Regional Uranium Study, Working Paper No. 28, Legal Infrastructure Related to Uranium Mining in the San Juan Basin, United States Department of Interior.

1.4.2. State Regulations

Federal statutes and regulations control many areas of environmental quality. Most state licensing or regulatory authority is often the result of a Federal-State agreement. However, land reclamation for uranium mining on federal and nonfederal lands is principally under state control. Table 1.12 shows the regulatory scheme for six states with uranium mining, and Appendix D lists the specific laws, regulations, and guides indicated generally in Table 1.12.

Agreement states have made formal arrangements with the NRC to develop programs to issue by-product, source material, and processing licenses. The Atomic Energy Act (Sec. 274), as amended, requires agreement states to provide by 1981 regulatory programs that are equivalent to or more stringent than the federal requirements for mill operations. Much of the environmental regulation of mining operations outside of federally controlled lands, especially for reclamation activities, currently depends upon state or local requirements. No NRC licenses are required for mining, except in situ.

The Federal Water Pollution Control Act amendments of 1972 give EPA National Pollution Discharge Elimination System (NPDES) permitting authority. However, Section 402 provides for approval of a state or interstate program to permit. The Administrator has established guidelines specifying procedural and other elements that must be present to obtain approval (40 CFR 124). Where states have not been approved, applicants apply for discharge permits from EPA. However, EPA asks what state requirements should also be certified so that state standards are met. Column 2 of Table 1.12 lists states that are approved to issue NPDES permits.

The Clean Air Act (CAA) amendments of 1970 and 1977 require, under Section 110, that State Implementation Plans (SIP's) must be submitted for approval to EPA for implementation of CAA on a local level. The approval and implementation of State plans are given in 40 CFR 52. In areas where NAAQS are violated, SIP's must produce compliance by 1982. If a state fails to enforce its plan, EPA may enforce it. There are currently no emission standard regulations specific for uranium mining by State governments.

Table 1.12 State laws, regulations, and guides for uranium mining

State	General			Mining							
	NRC Agreement State	NPDES Permit State	Water Use	Permits		Environmental Quality					Health and Safety
				Exploration Rights	Mining Rights	Air	Water		Land		
							Surf	UG	Solids	Reclam	
COLORADO	Yes	Yes	-	-	-	-	-	-	-	-	-
Department of Health											
Water Quality Control Div.	-	-	-	-	-		x	x	-	-	-
Air Quality Control Div.	-	-	-	-	-	x	-	-	-	-	-
Department of Natural Resources											
Div. of Water Reserves (State	-	-	x	-	-	-	-	-	-	-	-
Board of Land Commissioners	-	-	-	x	x	-	-	-	-	x	-
Mined Land Reclam Bd	-	-	-	x	x	-	x	x	-	x	-
Division of Mines	-	-	-	-	-	-	-	-	-	-	x
NEW MEXICO	Yes	No	-	-	-	-	-	-	-	-	-
State Land Commission	-	-	-	x	x	-	-	-	-	-	-
Dept. of Energy and Minerals	-	-	-	-	-	-	-	-	-	x	x
Dept. of Natural Resources	-	-	x	-	-	-	-	-	-	-	-
Env. Improvement Div.	-	-	-	-	-	x	x	x	x	-	x
TEXAS	Yes	No	-	-	-	-	-	-	-	-	-
Dept. of Water Resources	-	-	x	-	x	-	x	-	x	x	-
R.R. Commission of Texas	-	-	-	x	x	x	x	x	-	x	-
General Land Office	-	-	-	x	x	-	-	-	-	x	-
Dept. of Health	-	-	-	-	-	-	x	-	-	-	x
Air Control Board	-	-	-	-	-	x	-	-	-	-	-
UTAH	No	No	-	-	-	-	-	-	-	-	-
State Engineer	-	-	x	-	-	-	-	-	-	-	-
Dept. of Social Services	-	-	-	-	-	-	-	-	-	-	-
Division of Health	-	-	-	-	-	x	x	x	-	-	x
Water Pollution Control Bd.	-	-	-	-	-	-	x	x	-	-	-
Dept. of Natural Resources	-	-	-	x	x	-	-	-	-	x	-

Table 1.12 (continued)

State	GENERAL			Mining							Health and Safety
	NRC Agreement State	NPDES Permit State	Water Use	Permits			Environmental Quality				
				Exploration Rights	Mining Rights	Air	Water		Land		
							Surf	UG	Solids	Reclam	
WASHINGTON	Yes	Yes	-	-	-	-	-	-	-	-	-
Dept. of Natural Resources	-	-	-	x	x	-	-	-	x	x	-
Dept. of Ecology	-	-	x	-	-	-	-	-	-	-	-
Office of Water Programs	-	-	-	-	-	-	x	(No)	-	-	-
Dept of Social Services & Health	-	-	-	-	-	-	-	-	-	-	-
Health Services Division	-	-	-	-	-	-	-	-	-	-	x
Air Quality Division	-	-	-	-	-	x	-	-	-	-	-
WYOMING	No	Yes	-	-	-	-	-	-	-	-	-
State Inspector of Mines	-	-	-	x	x	-	-	-	-	x	x
State Engineers Office	-	-	x	-	-	-	-	-	-	-	-
Dept. of Env. Quality	-	-	-	-	-	-	-	-	-	-	-
Air Quality Div.	-	-	-	-	-	x	-	-	-	-	-
Water Quality Div.	-	-	-	-	-	-	x	x	-	-	-
Land Quality Div.	-	-	-	x	x	-	-	-	x	x	-
Solid Waste Management	-	-	-	-	-	-	-	-	x	-	-

Note.--An "x" indicates the existence of one or more controlling laws, regulations, or guides. See Appendix D for a list of the specific laws, regulations, or guides.

Applicable laws, regulations, and guidelines that apply to uranium mining in Colorado, New Mexico, Texas, Utah, Washington, and Wyoming are discussed below. Laws and regulations of other previously mined or potential uranium mining states, such as Arizona, California, Idaho, Montana, and South Dakota, are not reviewed. However, the basic environmental considerations of uranium mining should not be significantly different for other states.

1.4.2.1. Colorado

Colorado is an NRC "Agreement State" and has been approved by the EPA to issue NPDES discharge permits. Both radiation and water quality regulatory activities are under the jurisdiction of the Colorado Department of Health. The Health Department's Radiation and Hazardous Wastes Control Division administers radiation control activities and the control of hazardous wastes disposal. However, there are no operable rules or regulations for mining. Water quality is the responsibility of the Water Quality Control Commission (affiliated with the Health Department), which promulgates water quality standards and control regulations, and the Health Department's Water Quality Control Division, which administers and enforces the Commission's regulations and issues NPDES permits, as well as being responsible for numerous other water quality activities.

Colorado's permitting of discharges to "navigable" waters has been approved by EPA. Unlike most states, Colorado has promulgated specific "Guidelines for Control of Water Pollution from Mine Drainage" (November 10, 1970). These guidelines have the status of regulation since the State does not issue the NPDES permit unless the guidelines will be met. Colorado also has "Rules for Subsurface Disposal Systems" that, in conjunction with other rules, may assure protection of groundwater. These "Rules" cover all wastes that are disposed of underground, whether by direct or indirect means. "Wastes" include "any substance, solid, liquid, or gaseous, including radioactive particles thereof, which pollute or may tend to pollute any waters of the State." Solid waste and other land disposals are covered by Section 25-8-501, CRS 1973, as amended. In cases where these regulations do not control, the rules for subsurface disposal systems may apply.

The Colorado Department of Natural Resources administers water use in Colorado. As with most Western States, water is not in great abundance in Colorado. Determination of priority of water rights to surface and tributary groundwater is under the jurisdiction of a system of Water Courts, while the Division of Water Resources (State Engineer) administers and controls the allocation of actually available waters on an annual basis according to water rights priorities.

There are no State air quality standards or regulations that apply specifically to uranium mining. However, the National Ambient Air Quality Standards and various State emission control regulations apply to uranium mining activities as they do to all other types of emission sources. Colorado's air quality activities are the responsibility of the State Health Department's affiliated Air Quality Control Commission and its Air Quality Control Division. The Commission defines State air quality policy and promulgates air quality ambient standards and emission control regulations, while the Division administers and enforces the air quality regulations and issues emission permits.

The Board of Land Commissioners, affiliated with the Colorado Department of Natural Resources, issues permits for prospecting and controls leases for mining on State lands. The Board has policies and regulations concerning environmental impacts on prospected or leased lands.

The Colorado Mined Land Reclamation Board was created in 1976. It is administered by the Department of Natural Resources. The Board issues permits for all mining operations on all Federal and non-Federal lands in the State. The stated intent for Colorado Mined Land Reclamation Law is "to allow for the continued development of the mining industry in this State, while requiring those persons involved in mining operations to reclaim land affected by such operations so that the affected land may be put to a use beneficial to the people of this State. It is the further intent...to conserve natural resources, aid in the protection of wildlife and aquatic resources, and establish agricultural, recreational, residential, and industrial sites and to protect and promote the health, safety, and general welfare of the people...." The Board has established rules and regulations to implement the

law. Rule 5 (Prospecting Notice and Reclamation Requirements) considers prospecting a separate activity, but still covered by certain reclamation requirements. The reclamation performance standards of Rule 6 have specific requirements for grading, hydrology and water quality, wildlife safety and protection, topsoiling, and revegetation. Rule 7 ("Surety") assures reclamation. Before the Board issues any permit and before any Notice of Intent to Prospect is valid, the applicant must post surety with the Board. The amount of surety, established by the Board, is to be sufficient to fully reimburse the State for all expenses it would incur in completing the reclamation plan in the event of default by the operator.

Colorado also has regulations that apply for health and safety in mining operations. For each individual employee of any mining operation within the state a lifetime history is maintained on exposure to radon daughter concentrations when certain minimum values are reached. The State Department of Natural Resources Division of Mines administers these.

1.4.2.2 New Mexico

In New Mexico, a mine plan must be filed with and approved by the State Mining Inspector before he will issue a permit. The State Mining Inspector does not review the plan for environmental impact. Groundwater use rights are established by the State Engineer, and the Land Commission handles exploration and mining rights. The engineer's office issues a permit for beneficial use of any water pumped from uranium mines. However, the Navajo tribe claims jurisdiction of the State's groundwater in the northwest region of New Mexico. It is likely that the Departments of Interior and Justice will eventually become involved in this dispute as Trustees for the tribe.

Approval status has been given, with some exceptions, by EPA to New Mexico's plan for the attainment and maintenance of national air standards (40 CFR 52.1622). However, neither Federal nor State regulations include specific emission standards for uranium mining. But "Ambient Air Quality Standards" (40 CFR 50.6) on suspended particulates apply to all sources of air pollution.

New Mexico is an NRC agreement state, but it is not an approved NPDES state. Part 2 of the amended Water Quality Control Commission regulations applies to any discharge that is not subject to a permit under the NPDES system. The State requires approved discharge plans for discharges that could contaminate groundwater. However, the applicable NPDES regulations (Subpart E-Uranium, Radium and Vanadium Ores Subcategory, 40 CFR 440.50) have been challenged by some mine operators. They claim that discharges to a dry arroyo do not constitute "the discharge of pollutants into the navigable water, water of the contiguous zone, and the oceans." Because more than half of active New Mexico mine discharges have NPDES permits that are now under adjudication, there is no enforcement and discharges may not be in accordance with Standards. If the NPDES challenge is sustained, then New Mexico's Part 2 regulations could be applied, even though they are not particularly suitable for uranium mining discharges. Possibly only the regulations on chemical oxygen demand and settling of heavy metal solids would apply to uranium mine wastes. The Part 3 "Regulations for Discharges onto or below the Surface of the Ground" (3-100) that are designed to "protect all groundwater" would also be important. A discharge plan is required for effluent discharges that move directly or indirectly into groundwater, if the effluent contains any of the contaminants listed in Section 3-103 a, b, and c, or toxic pollutants. Since the list of contaminants includes uranium and radium, New Mexico can approve only discharge plans meeting the drinking water standards.

There are no state regulations for solid wastes and land reclamation for mining operations. The mining plan and bonding requirements associated with mining permits determine the extent of mining reclamation.

Radiation safety requirements (Sections 74-3-1 et seq NMSA 1978) apply to both mining and milling. Air quality monitoring in underground mines currently involves potential duplication of effort by the New Mexico Mine Inspector (69-5-7 et seq NMSA 1978) and the Federal Mine Safety and Health Administration (30 CFR 57.5-37).

1.4.2.3 Texas

Texas is an NRC agreement state, but not an EPA approved NPDES permit state. The Department of Water Resources controls water use. Even though much of the water used in Texas comes from wells, there are no regulations on pumping groundwater. However, some counties have regulations that limit groundwater withdrawal to control subsidence.

Specific regulations for in situ uranium mining are enforced by the Texas Department of Health (TDH). Since Texas is an agreement state, its regulations reflect all appropriate NRC regulations. The TDH also implements the Safe Drinking Water Act (SDWA) and monitors groundwater to assure that its provisions for radium and selenium concentrations are met.

The General Land Office (GLO) issues prospecting permits and mining leases on state-owned lands. Mining and reclamation plans for uranium mining on state-owned lands are reviewed for approval by GLO. The "Texas Uranium Surface Mining and Reclamation Act" exempts state-owned lands from regulation by the Railroad Commission.

Surface mining is regulated by the Railroad Commission. All requirements of state and federal laws must be fulfilled before a permit is issued. Mining and reclamation plans must be submitted and approved. A bond is required to assure reclamation after mining.

The Texas Air Control Board administers provisions of the Clean Air Act. Except for suspended particulates, there are no applicable standards, i.e., there are no state source standards, for uranium mining.

The Texas Guides and Regulations for Control of Radiation (TRCR) do not apply to surface uranium mining. They do apply to in situ mining due to NRC agreement state licensing. The radioactive content of water discharged from all mines to the environment must not exceed TRCR limits.

1.4.2.4 Utah

Utah is neither an NRC agreement state nor an NPDES permit approved state. The Utah State Engineer's Office is responsible for approval of water

use rights. The Department of Natural Resources oversees exploration and mining rights on State lands.

The Division of Oil, Gas, and Mining of the Department of Natural Resources issues permits for uranium mining operations, except in situ mining licensed by the NRC. A mining and reclamation plan must be approved. Rule M-10 standards include consideration of land use, public safety and welfare, impoundment, slopes, high walls, toxic materials, roads and pads, draining, structures and equipment, shafts and portals, sediment control, revegetation, dams, and soils. Bonding requirements assure reclamation.

Discharges to surface waters are regulated under the EPA administered NPDES system and the Utah Water Pollution Committee. Utah does have separate regulations administered by the Department of Social Services. These are applied to mining operations such as non-discharging waste water systems and in situ mining where no NPDES permit is required.

No sources of pollution will be allowed to cause groundwaters to exceed drinking water standards. The applicable standards for classes 1A and 1B domestic water sources are given in Wastewater Disposal Regulations, Part II.

Utah is developing radiation safety regulations. We do not expect that they will apply to uranium mining, since they are based on the model state suggested regulations.

1.4.2.5 Washington

Washington is an NRC agreement state and an NPDES approved permit state. The Department of Ecology regulates water use and water quality. Washington has no regulations for groundwater. These waters could be protected under the Safe Drinking Water Act.

The Department of Natural Resources controls exploration and mining rights for state-owned lands only. The mineral lease law covers both surface and underground mining but not in situ or heap leaching. The State Reclamation Act applies to state and private lands only. A mining and reclamation bond is required before a permit is issued. Reclamation is assured

Washington has a Clean Air Act under which regulations have been promulgated consistent with the Federal Clean Air Act. No source emission standards have been issued for uranium mining. National Ambient Air Quality Standards could apply to suspended particulates.

Washington has rules and regulations for radiation protection, but they do not apply to uranium mining.

1.4.2.6 Wyoming

Wyoming is an approved NPDES permitting State but not an NRC Agreement State. The State Engineer's Office controls water use rights. Control is primarily on the quantity of water used, but there is some statutory responsibility regarding sedimentation. Discharges to surface waters are regulated by the Water Quality Division of the Department of Environmental Quality. The construction of any water or waste water facility requires a construction permit. Groundwater regulations have been proposed. These include groundwater quality standards for any activity. Permitting requirements specific to in situ uranium operations is one of a group of special process discharges.

The Land Quality Division of the Wyoming Department of Environmental Quality is the principle agency responsible for enforcing environmental protection standards and reclamation standards with respect to uranium mining operations. The Division also enforces mineral exploration regulations that afford protection to groundwater and restoration of significant surface disturbances.

Wyoming law requires that uranium mined land must be restored to a use at least equal to its highest previous use (W.S. 35-11-402(a)(i) and (ii)) and mining operations must be conducted to prevent pollution of waters of the State (W.S. 35-11-402(a)(vi)). Before a mining operation receives a permit it must submit to the Department a mining and reclamation plan that demonstrates compliance with the law and associated rules and regulations. The plan must contain a plan for the disposal of all acid-forming, toxic materials or materials constituting a fire, health, or safety hazard uncovered

or created by the mining process: radioactive material is included (W.S. 35-11-406(b)(ix)).

An operator must also, in accord with his approved mine and reclamation plan, cover, bury, impound, contain, or dispose of toxic, acid-forming, or radioactive material determined to be hazardous to health and safety or constitute a threat of pollution to surface or subsurface waters (W.S. 35-11-415(b)(iv)). A required surety bond assures that the operator will reclaim the land according to his approved plan. If the bond is forfeited, the State is responsible for reclamation.

Wyoming has legislated authority for a position on radiological restoration of mined lands. It is described in the Division's Guideline No. 1, Section III. The Division is presently drafting regulations for radiation protection on uranium mined lands and handling of uranium mine wastes. These regulations shall set standards.

Wyoming also has a solid waste management program that presently regulates only refuse generated at mines. Solid waste disposal sites are permitted at these facilities. Solid waste regulations could be promulgated that affect mining.

In Wyoming, Ambient Air Quality Standards are applied to mining operations, and fugitive emissions are controlled to the extent that these standards are met. An Air Quality Permit is required for the construction of a uranium mining and/or processing facility, and the applicant is required to demonstrate that applicable ambient and PSD (Prevention of Significant Deterioration) provisions are met.

Wyoming has radiation protection regulations for the safety of mines while they are actually in process. These regulations are under the jurisdiction of the State Inspector of Mines. According to Wyoming Law, the protection of miners from hazardous exposure to radioactivity must conform to the American Standards Association revised Publication N 13.8, "Radiation Protection in Uranium Mines and Mills." The uranium regulations (94-R-11) are found in Chapter 3, Article 4 of Title 30 - Mines and Minerals.

1.5 References

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SECTION 2

INVENTORY OF URANIUM MINES

2.0 Inventory of Uranium Mines

To inventory the numbers, types, and locations of uranium mines in the United States, we used data from the Department of Energy, Grand Junction Office (DOE-GJO). We produced the inventory of uranium mines presented in this section and Appendixes E and F of this report from the DOE-GJO master data file (DOE79a) and personal communications with DOE-GJO (Ch80, ME80a, ME80b). These two sources combined yielded our own EPA master data file, which we divided into two parts - active and inactive mines.

Table 2.1 classifies active and inactive U.S. uranium properties according to the method of uranium production (mine type) based on data that were current as of 1978 (Me80a). The major mining methods are surface and underground mines (DOE79b). The remaining mining methods are only minor contributors to the total uranium ore production (DOE79b).

Table 2.1 shows a total of 340 active mines. This final total, which is 52 less than the original total of 392 active mines provided by DOE-GJO (Me80a), was derived, in consultation with DOE-GJO (Me80b), by eliminating 43 mines that were duplicated on the list and 9 that were small producers (i.e., producing only a few tons of ore for the entire year of 1978). Most (if not all) of the 52 eliminated mines were either underground or surface mines.

The original totals of 305 active underground mines and 63 active surface mines (DOE79b), whose combined total of 368 mines accounts for the later eliminated 52 mines that were duplicate listings or small producers, were the totals we used in modeling the average underground and surface mines in this study. The differences between these totals and the smaller Table 2.1 totals of 256 active underground mines and 60 active surface mines are insignificant compared with other uncertainties in predicting health effects. The smaller totals for underground and surface mines would introduce differences of less than 17% and less than 5% for the active average underground and average surface model mines, respectively.

Table 2.2 gives locations and types of active uranium mines by state. With respect to the number of mines, Colorado and Utah dominate the inventory, especially for underground mines. However, since New Mexico and

Wyoming have large mines (underground in New Mexico, and surface in Wyoming) and dominate ore production, New Mexico is the site of our model active underground mines and Wyoming is the site of our model active surface mines. Our model in situ leaching operation is also sited in Wyoming, which is one of two states mining uranium with that method. Appendix E gives a complete inventory of active uranium mines.

The numbers of inactive uranium mines according to state and mining method are given in Table 2.3. Colorado and Utah have the greatest number of inactive mines, but Arizona, Wyoming, New Mexico, and South Dakota also contain significant numbers. Since New Mexico and Wyoming have dominated ore production over the past 10 years (DOE79b), New Mexico (because of its large underground mines) is our model site for inactive underground mining and Wyoming (because of its large surface mines) is the site of our model inactive surface mine. Appendix F gives a complete inventory of inactive uranium mines.

Figures 2.1 through 2.9 are maps showing the locations, status, and types of uranium mines in Colorado, New Mexico, Texas, Utah, and Wyoming (Ch77, Co78a, Co78b, Co78c, Ea73, G175, Hi69, Pe79, Ut77). Since it is not always possible to show all the mines in a given district, the maps indicate only the area and number of mines in some major mining districts, particularly for Colorado and Utah. The maps do not show the location of many small mines started during the uranium boom of the 1950's because their exact locations are unknown. In Colorado alone there are over a thousand such mines.

Table 2.4 shows total ore production through January 1, 1979 for active and inactive surface and underground mines. The larger mines (>910 MT ore production) dominate the list of active mines, and the smaller mines (<910 MT ore production) dominate the inactive list. If remedial action becomes necessary for inactive mines, the information in Table 2.4 could help estimate the magnitude of such an action, at least affording a way to make rough estimates of waste rock, sub-ore, and overburden that are present at the inactive site. A recent DOE report (DOE79c) contains additional information on mining waste tonnage and acreage of specific properties.

Table 2.1 Type of U.S. uranium properties

Uranium Production Method ^(a)	Number of Active Properties	Number of Inactive Properties
Surface mine	60	1252
Underground mine	256	2036
Mine water production	2	1
Heap leach - dumps	1	7
Heap leach - ores	0	1
Dumps	1	42
Sub-ore	1	12
In-situ leaching	11	2
Miscellaneous	0	23
Tailings dump	2	0
Unknown	6	13
TOTAL	340	3389

(a) Categories listed in this column are modifications of the originals (DOE79a). Copper by-product and surface-underground combination categories were eliminated because they contained no properties. The miscellaneous-phosphate by-product category was reduced to miscellaneous because most phosphate by-product properties were not included in the DOE-GJO master data file (DOE79a). The low grade or protore category was changed to sub-ore to be consistent with the rest of this report.

Table 2.2 The location and type of active uranium properties

State	Surface Mine	Underground Mine	Mine Water Production	Heap-Leach Dumps	Heap-Leach Ores	Dumps	Sub-ore	In-Situ Leaching	Miscellaneous	Tailings Dump	Unknown	Total
Arizona	1	1	0	0	0	0	0	0	0	0	0	2
Colorado	5	106	0	0	0	0	0	0	0	1	3	115
New Mexico	4	35	2	0	0	1	0	0	0	0	0	42
Texas	16	0	0	0	0	0	0	8	0	1	0	25
Utah	13	108	0	0	0	0	0	0	0	0	3	124
Washington	2	0	0	0	0	0	0	0	0	0	0	2
Wyoming	19	6	0	1	0	0	1	3	0	0	0	30
TOTAL	60	256	2	1	0	1	1	11	0	2	6	340

Table 2.3 The location and type of inactive uranium properties

State	Surface Mine	Underground Mine	Mine Water Production	Heap-Leach Dumps	Heap-Leach Dues	Dumps	Sub-ore	In-Situ Leaching	Miscellaneous	Tailings Dump	Unknown	Total
Alaska	0	1	0	0	0	0	0	0	0	0	0	1
Arizona	135	189	0	1	0	0	0	0	0	0	1	326
California	13	10	0	0	0	0	0	0	0	0	0	23
Colorado	263	902	0	0	0	35	1	0	10	0	6	1217
Florida	0	0	0	0	0	0	0	0	1	0	0	1
Idaho	2	4	0	0	0	0	0	0	0	0	0	6
Minnesota	0	0	0	0	0	0	0	0	1	0	0	1
Montana	9	9	0	0	0	0	0	0	0	0	0	18
Nevada	9	12	0	0	0	0	0	0	0	0	0	21
New Jersey	0	1	0	0	0	0	0	0	0	0	0	1
New Mexico	34	142	1	0	0	0	8	0	1	0	2	188
N. Dakota	13	0	0	0	0	0	0	0	0	0	0	13
Oklahoma	3	0	0	0	0	0	0	0	0	0	0	3
Oregon	2	1	0	0	0	0	0	0	0	0	0	3
S. Dakota	111	30	0	0	0	0	0	0	0	0	0	141
Texas	38	0	0	0	1	0	0	2	0	0	1	42
Utah	378	698	0	0	0	7	1	0	6	0	3	1093
Washington	13	0	0	0	0	0	0	0	0	0	0	13
Wyoming	223	32	0	6	0	0	2	0	2	0	0	265
Unknown	6	5	0	0	0	0	0	0	2	0	0	13
TOTAL	1252	2036	1	7	1	42	12	2	23	0	13	3389

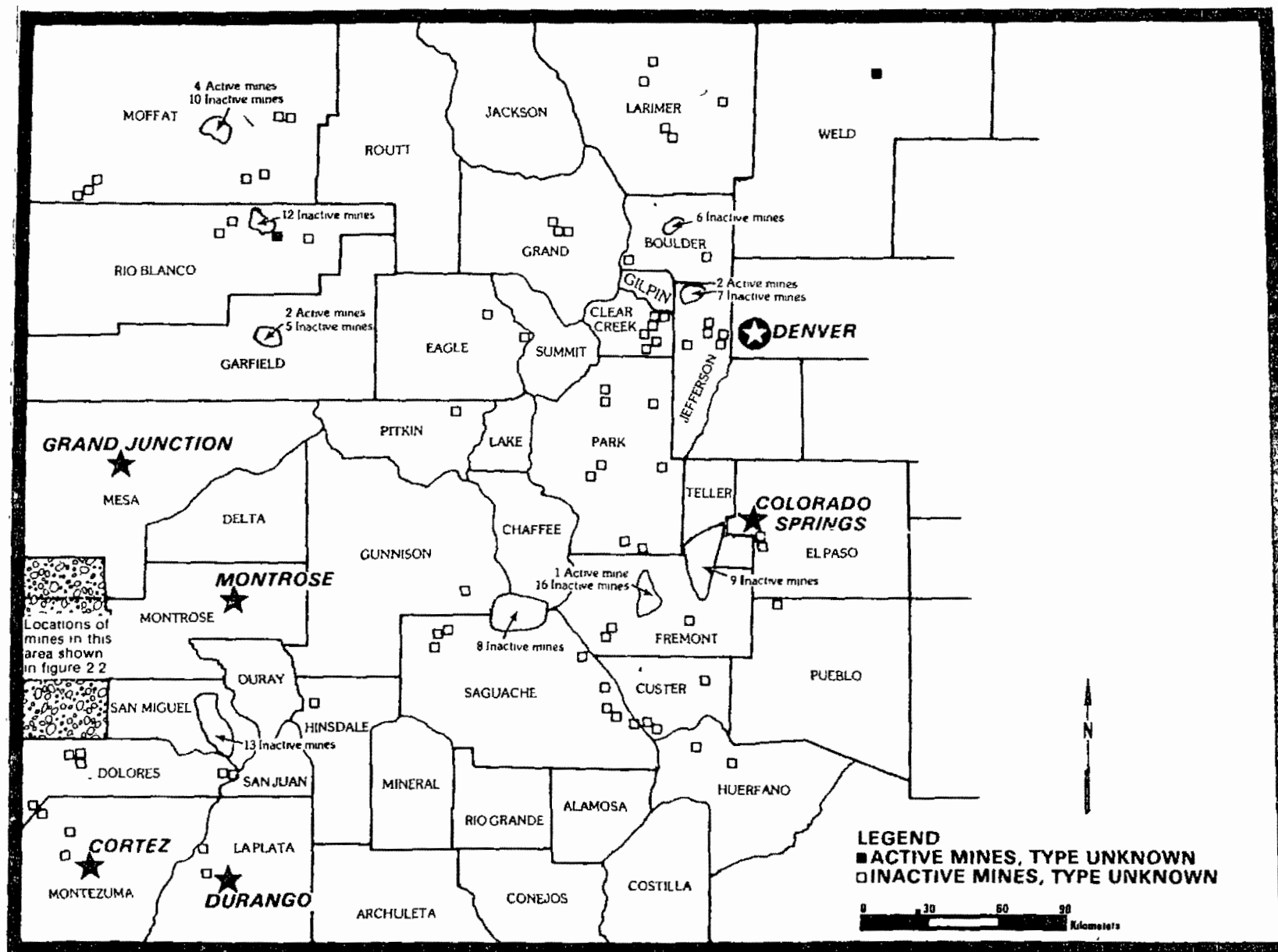


Figure 2.1 Location of active and inactive uranium mines and principal uranium mining districts in Colorado

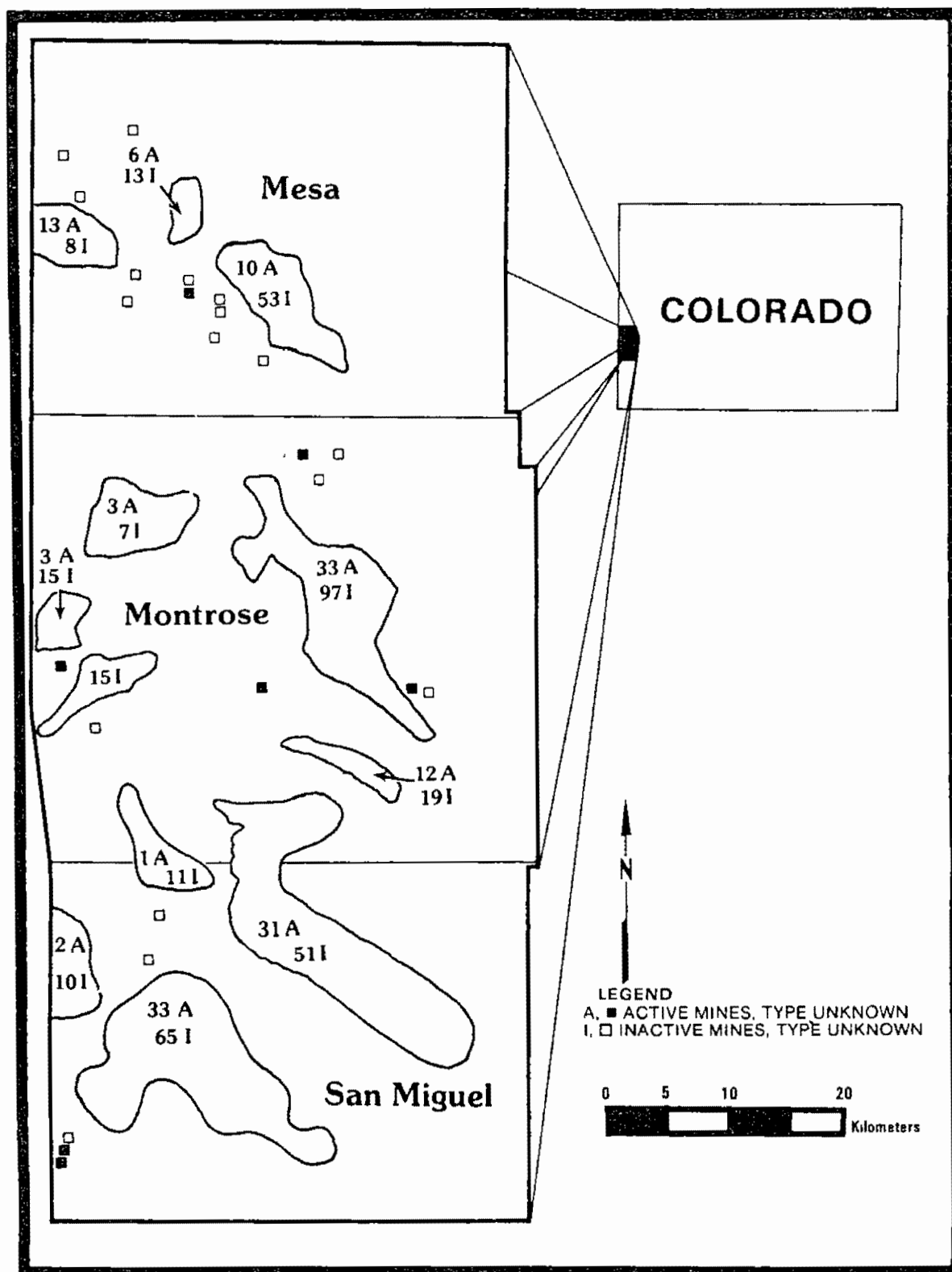


Figure 2.2 Location of active and inactive uranium mines and principal uranium mining districts in the Urvan Mineral Belt of western Colorado

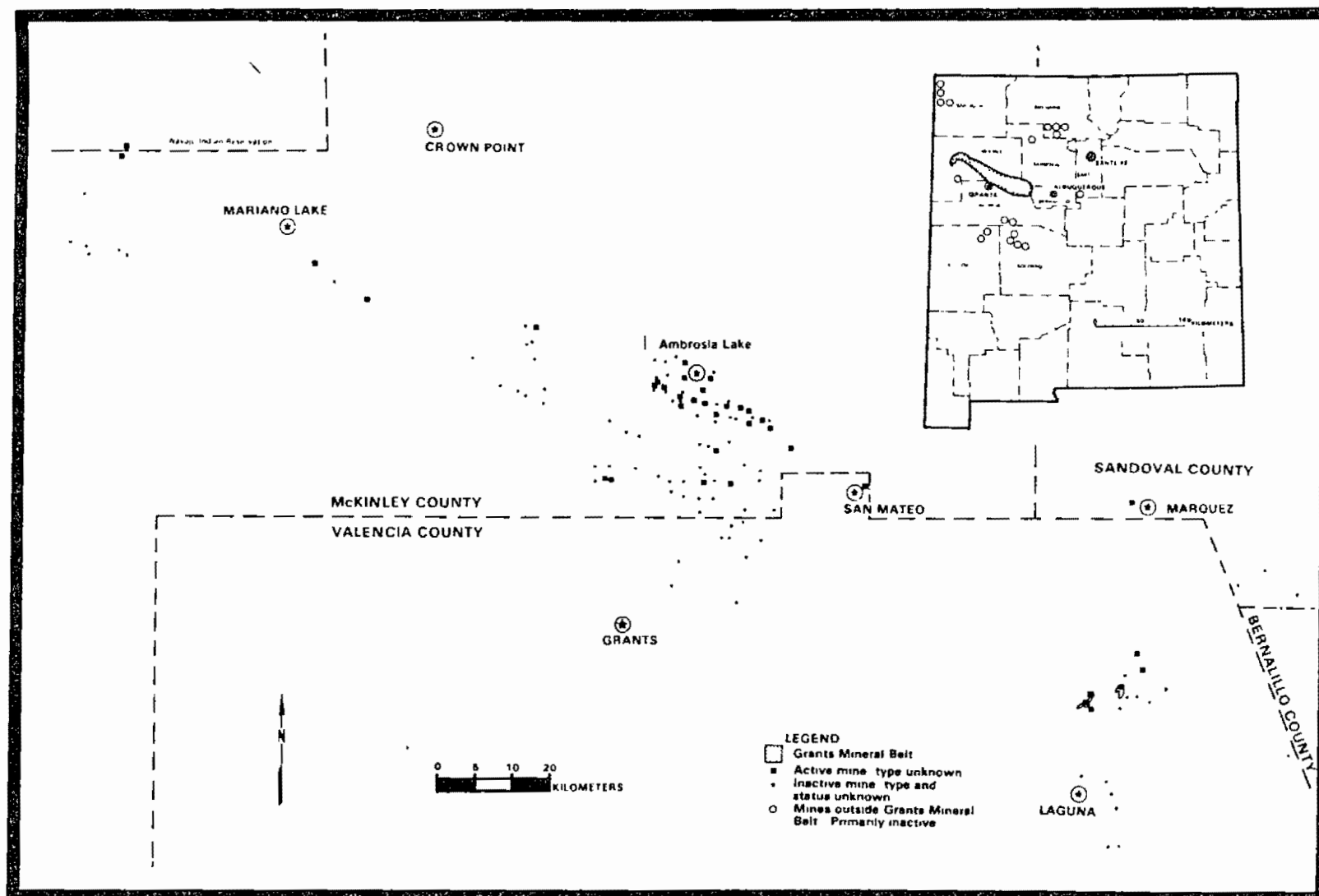


Figure 2.3 Location of active and inactive uranium mines in the Grants Mineral Belt and other areas of New Mexico

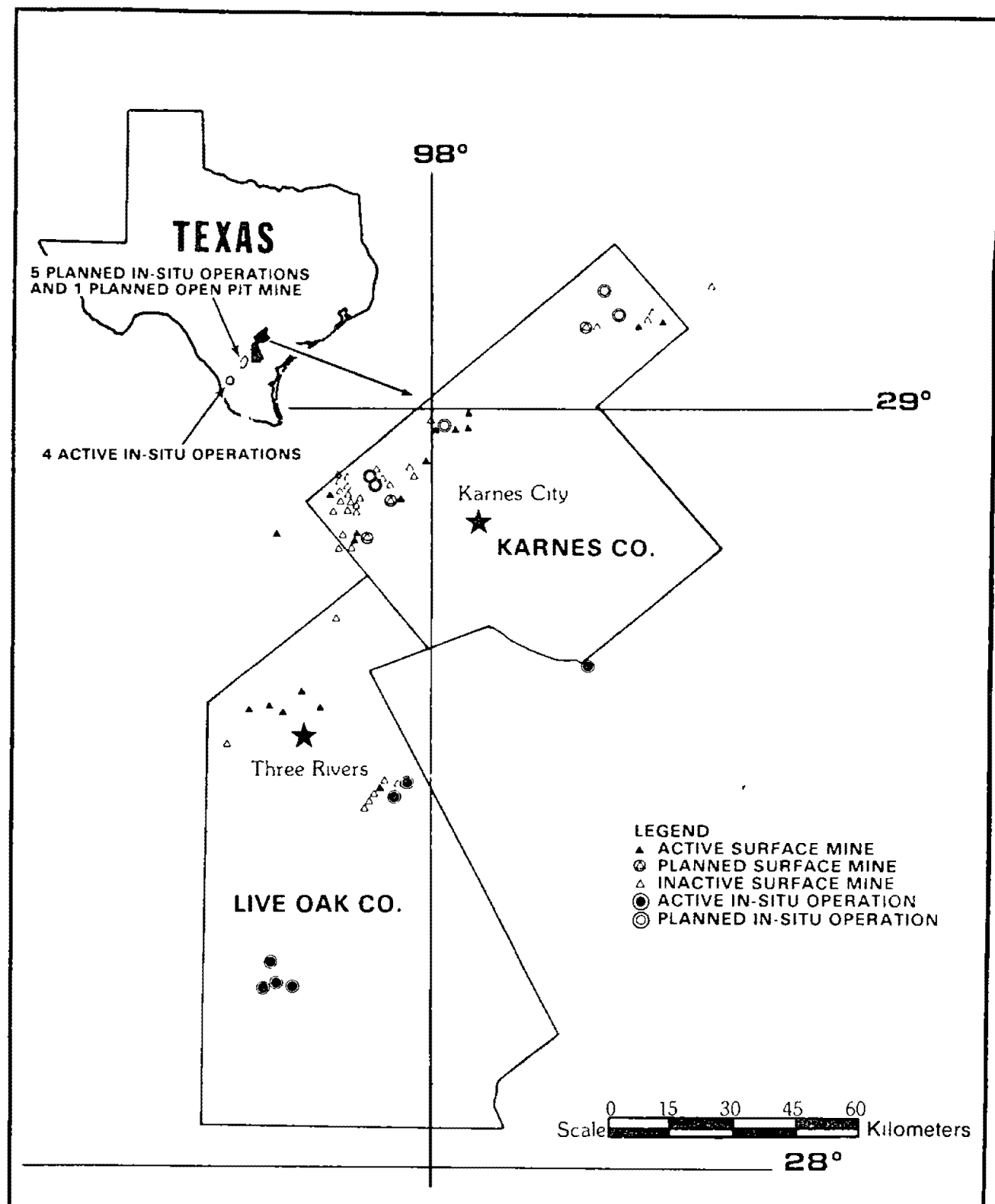


Figure 2.4 Location of active, inactive, and proposed surface and in situ uranium mines in Texas

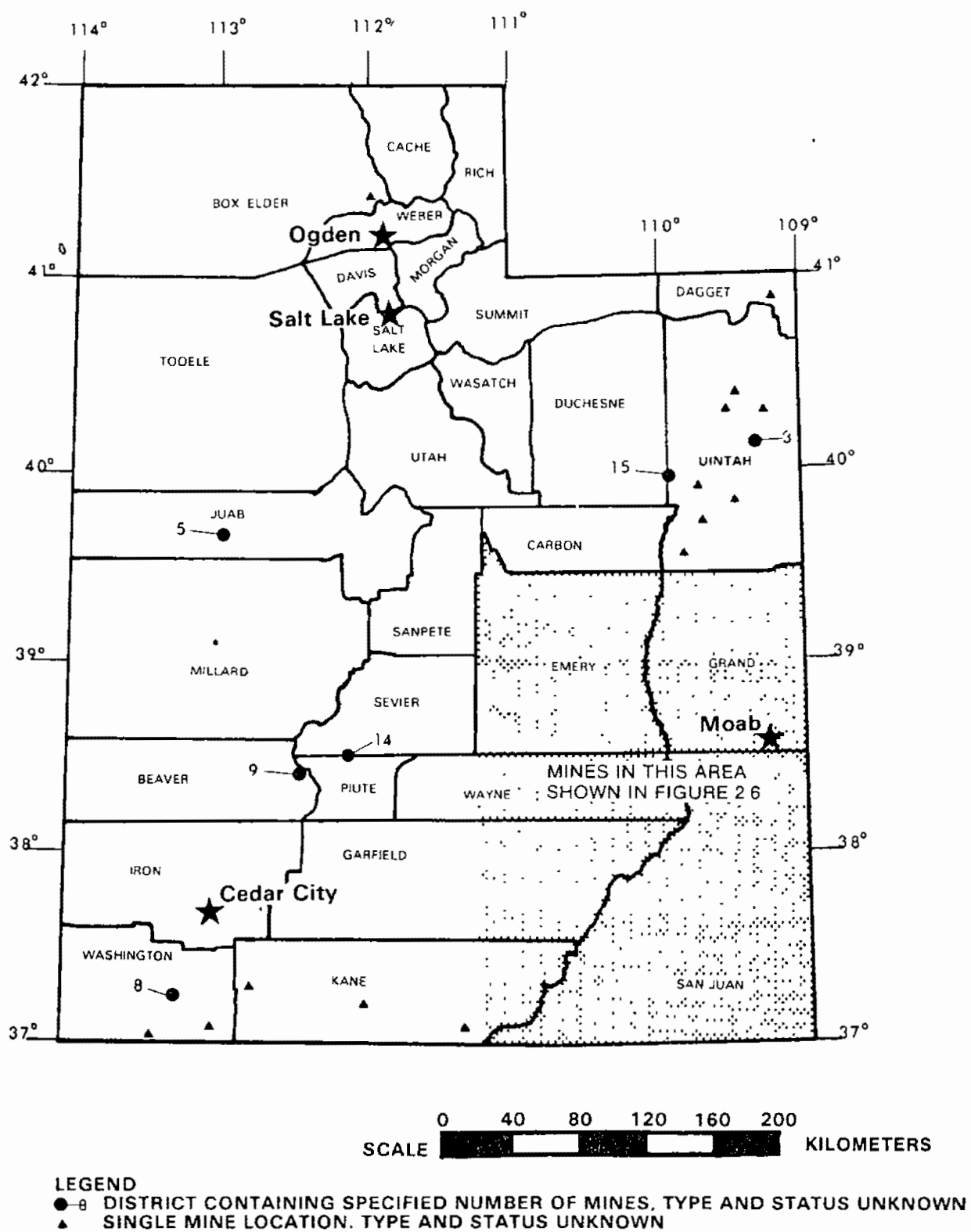


Figure 2 5 Location of uranium mines and mining districts in Utah

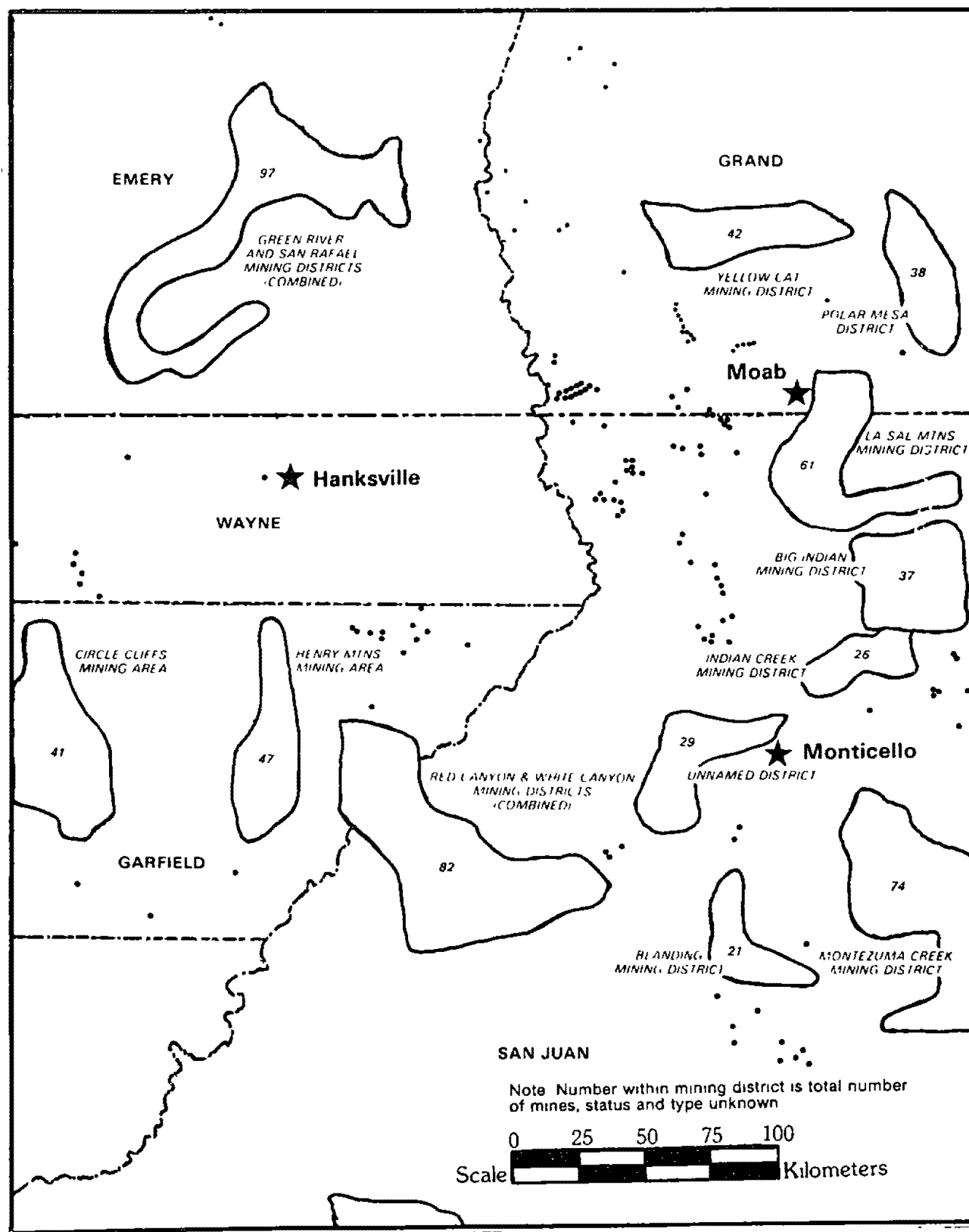


Figure 2-6 Location of uranium mines and principal uranium mining districts in southeastern Utah

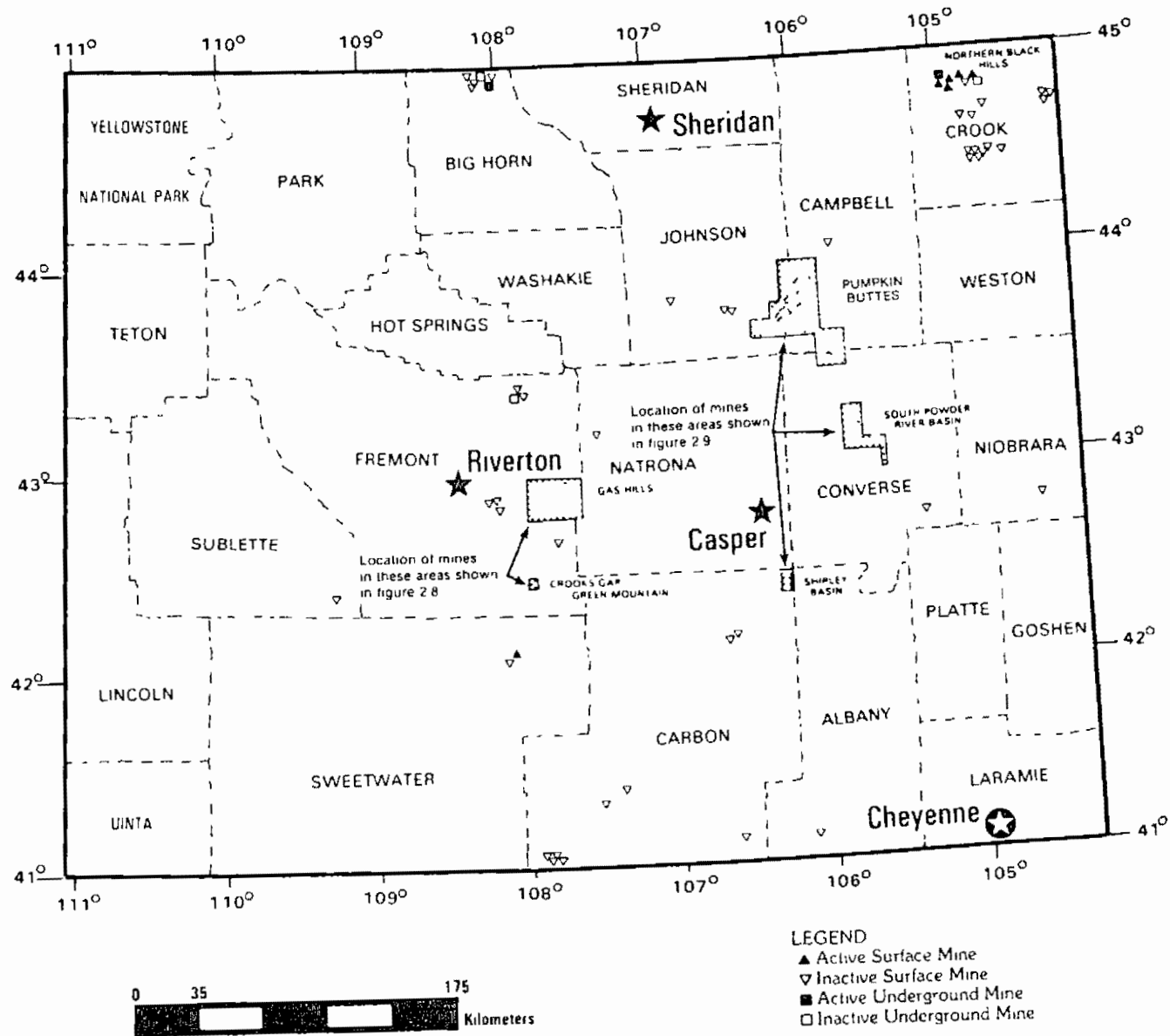


Figure 27 Location of active and inactive uranium mines and principal uranium mining areas in Wyoming

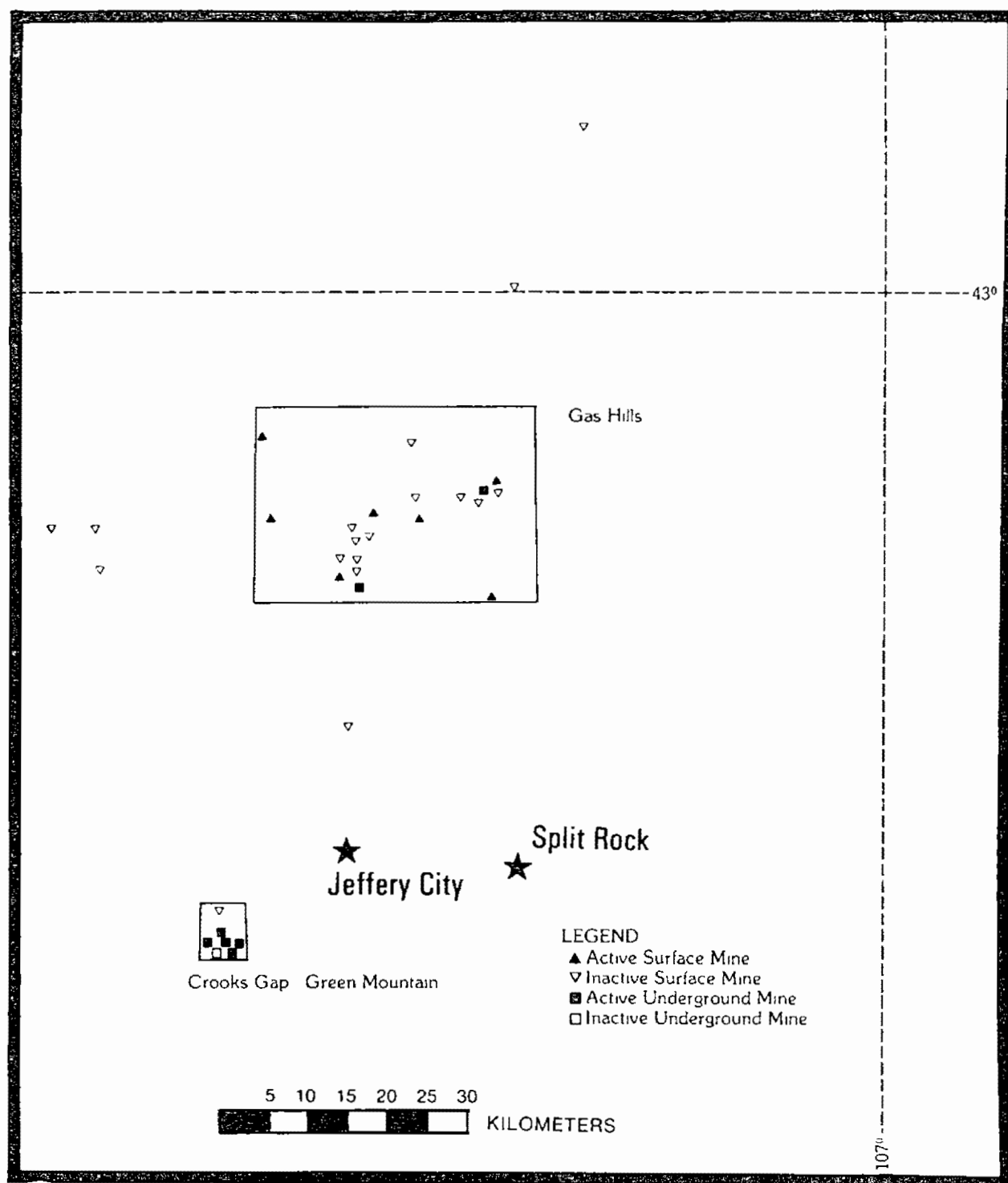


Figure 2 8 Location of active and inactive uranium mines in the Gas Hills and Crooks Gap-Green Mountain areas of central Wyoming

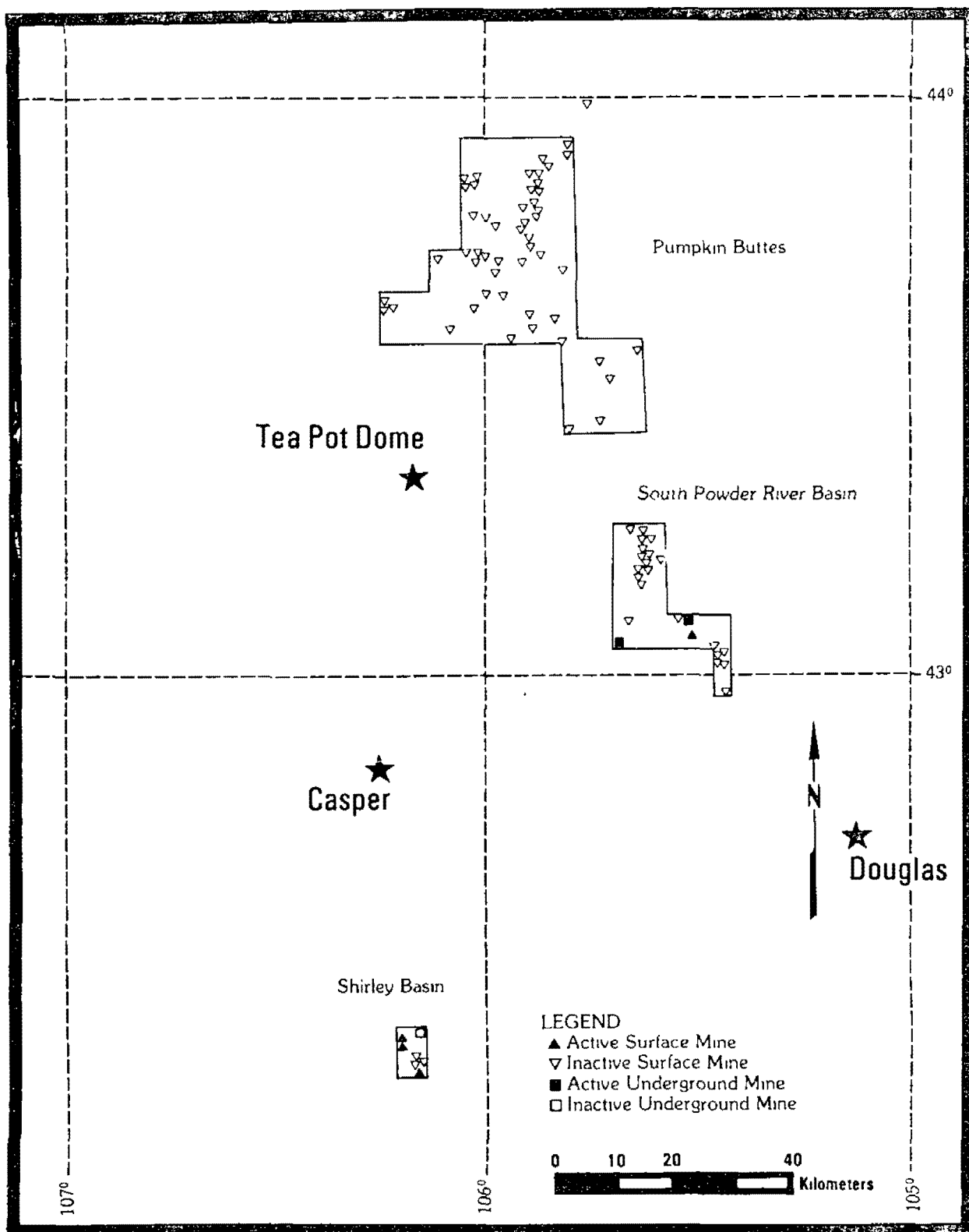


Figure 2 9 Location of active and inactive uranium mines in the Shirley Basin, South Powder River Basin, and Pumpkin Buttes areas of Wyoming

Table 2.4 Cumulative Ore Production
through January 1, 1979

Ore Production MT	Active				Inactive			
	No. Mines	(% of total)	Under-		No. Mines	(% of total)	Under-	
			Surface ground				Surface ground	
< 91	16	(4.7)	8	3	1553	(45.8)	899	628
91-910	33	(9.7)	5	24	753	(22.2)	134	588
910-91,000	188	(55.3)	15	165	986	(29.1)	180	766
> 91,000	103	(30.3)	32	64	97	(2.9)	39	54
Total	340	(100.0)	60	256	3389	(100.0)	1252	2036

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SECTION 3

POTENTIAL SOURCES OF CONTAMINANTS TO THE ENVIRONMENT AND MAN

3.0 Potential Sources of Contaminants to the Environment and Man

3.1 Background Concentrations of Radionuclides and Trace Metals

3.1.1 Naturally Occurring Radionuclides

Potassium-40 and radionuclides in the decay chains of uranium-238 and thorium-232 are the principal sources in the earth's crust of background radiation. Figures 3.1 and 3.2 show the uranium-238 and thorium-232 decay chains. Potassium-40 constitutes 0.0118 percent of naturally occurring potassium. Its half-life is 1.26×10^9 years and, upon decay, potassium-40 emits a 1.46 MeV gamma ray in 11 percent of its disintegrations. Table 3.1 lists the average concentration of and gamma-ray energy released by these radionuclides in one gram of rock. Table 3.2 lists the radionuclide content and dose equivalent rates from common rocks and soils. Potassium-40 and the thorium-232 decay chain each contribute about 40 percent of the dose rate at 3 feet above the ground while the uranium-238 decay chain contributes approximately 20 percent of the total dose rate.

Radon-222 occurs in the uranium-238 decay chain and has a half-life of 3.8 days. It is a noble gas and, upon decay, produces a series of short-lived, alpha-emitting daughters (see Fig.3.1). The average atmospheric radon concentration in the continental U.S. is 0.26 pCi/liter (0a72). Under most conditions, the radon daughters contribute less than 10 percent (a few tenths of a μ rem/hr) to the terrestrial external dose equivalent rate. However, inhaled radon daughters contribute a large fraction of the total dose equivalent rate to the respiratory tract: about 50 percent (90 mrem/yr) to the lung and nearly all of the dose (450 mrem/yr) to the segmental bronchioles (NCRP75).

Eighty-five percent of the surface area of the United States, and nearly all of its population, is underlain by rocks and soils of sedimentary origin. However, the correlation between the bedrock activity and the aboveground activity is not clear.

In most soils, the amount of water varies from 5 to 25 percent. The soil moisture attenuates gamma radiation from the soil. The potassium-40 dose equivalent rate can decrease by 30 percent when the soil water content increases from 0 to 30 percent (0A72). Moisture can retard the diffusion of radon into the atmosphere and reduce the exposure to airborne radon daughters. Since radon daughters account for 95 percent of the gamma-ray energy from the uranium-238 series, their accumulation in the ground increases

URANIUM — 238 DECAY SERIES

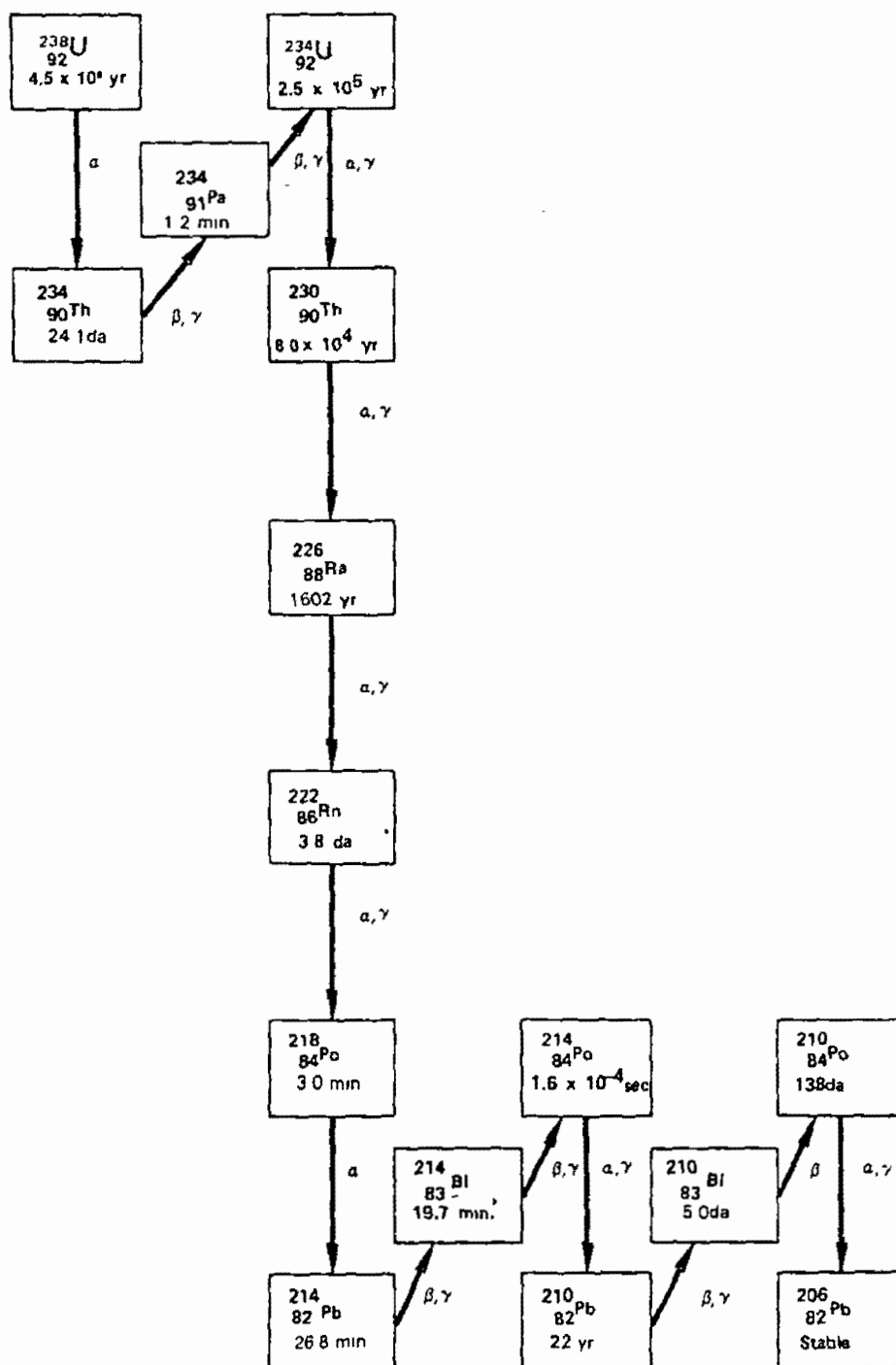


Figure 3 1 The uranium decay series showing the half lives and mode of decay

THORIUM - 232 DECAY SERIES

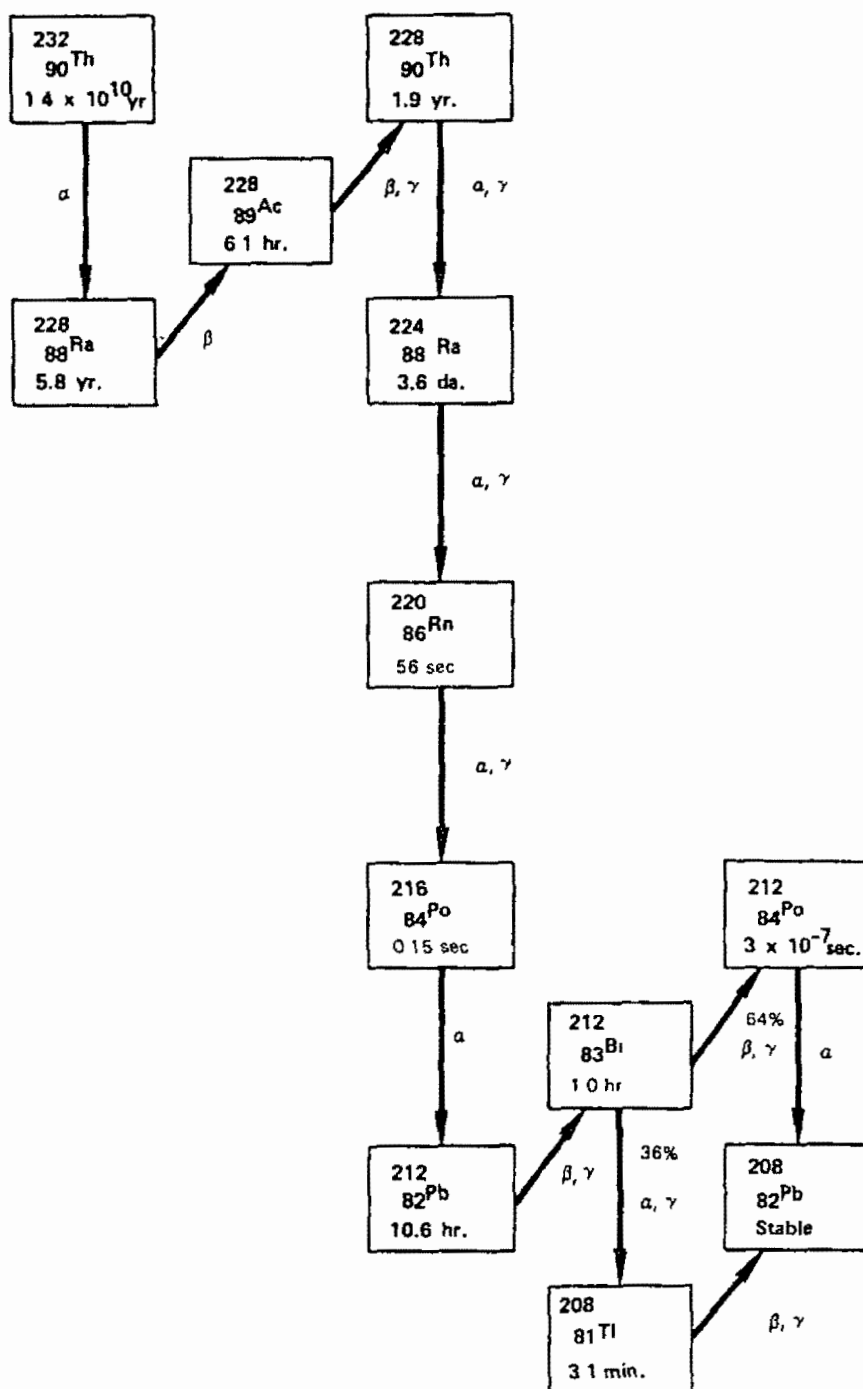


Figure 3.2 The thorium decay series showing the half lives and mode of decay

Table 3.1 Gamma-ray energy released by one gram of rock

Isotope	Average Concentration, Percent	Energy, KeV/sec.
Uranium-238 (in equilibrium with decay products)	2.98×10^{-4}	68.2
Uranium-235 (in equilibrium with decay products)	0.02×10^{-4}	1.53
Thorium-232 (in equilibrium with decay products)	11.4×10^{-4}	87.8
Potassium-40	3.0	149
Other Elements	---	2.7

Source: Oa72.

Table 3.2 Radionuclide content and dose equivalent rates from common rocks and soil

Rock Type	Uranium		Thorium		Potassium-40		Total mrem/yr
	ppm	mrem/yr ^(a)	ppm	mrem/yr ^(a)	ppm	mrem/yr ^(a)	
Igneous ^(b) basic	0.9	5.2	2.7	7.3	1.2	14.7	27.2
Silicic (granite)	4.7	26.9	20.0	53.8	5.0	61.3	142.0
Sedimentary ^(b)							
Shale	3.7	21.2	12.0	32.3	3.2	39.2	92.7
Sandstone	0.45	2.6	1.7	4.6	1.1	13.5	20.7
Limestone	2.2	12.6	1.7	4.6	0.32	3.9	21.1
Upper crustal average ^(c)	2.8	16.0	10	26.9	2.4	29.4	72.3
U.S. surficial average ^(d)	1.8	10.3	9.0	24.2	1.8	21.8	56.3

(a) mrem/yr/ppm. uranium, 5.73; thorium, 2.69; potassium-40, 12.3 (Be68).

(b) Source: C166.

(c) Uranium and thorium averages (Ph64); potassium (He69).

(d) Source: Lo64.

the exposure from this series. Thus, soil moisture decreases the potassium-40 and thorium-232 dose equivalent rates and increases or leaves unchanged the uranium-238 series dose equivalent rate.

Snow cover also affects the terrestrial dose equivalent rate and the radon emanation rate. Gamma radiation attenuates exponentially as a function of the density and thickness of the snow cover (0a72). However, the overall influence of snow on population exposure is negligible since, in most populated areas, there is relatively little snowfall that remains for long periods of time.

Table 3.3 shows the average dose equivalent rate due to radiation in some Western mining states (0a72). Terrestrial radiation in the Western uranium mining states is higher than in the rest of the nation due to the greater concentration of the uranium-238 series.

Table 3.3 Average dose equivalent rates due to terrestrial radiation in western mining states

State	Terrestrial Dose, mrem/yr
Arizona	45.6
Colorado	65.8
New Mexico	51.7
South Dakota	45.6
Texas	29.0
Utah	45.6
Wyoming	45.6

Concentrations of radionuclides measured in surface and groundwater samples collected on a proposed uranium project site are listed in Table 3.4 (NRC79a). The large variations among concentrations at different collection

sites are typical of surface water concentrations. (Concentrations in sea water are more uniform.) Hence, generalizations about background concentrations of radionuclides in fresh water systems are impractical. Extensive, site-specific studies over an extended period of time are necessary to obtain meaningful background concentrations for a site.

3.1.2 Stable Elements

Concentrations of metals occurring in the earth's crust generally range from several parts-per-billion (ppb) to a few parts-per-million (ppm). Measured concentrations vary widely from site to site and often in different samples taken from the same site. Table 3.5 lists the results of measurements for selected elements. It should be emphasized that these are general estimates of element composition of rocks in the United States and do not reflect large variations that occur within the different rock types.

Concentrations of metals measured in surface and groundwater samples collected from different locations on a proposed uranium project site are listed in Tables 3.6 and 3.7, respectively. There are large differences in the composition of surface and groundwaters. Table 3.8 shows the average concentrations of three trace metals that are sometimes associated with mine discharge water. These values, which were taken from the results of an extensive study (Tu69), approximate average concentrations in United States streams. Background concentrations at any specific site could be much different.

Table 3.8 Estimated average concentrations (ppb) of three metals in U.S. streams

Element	Turekian's Results	Other results from Literature
Chromium	1.4	1.0
Molybdenum	1.8	1.0
Selenium	0.2	0.2

Source: Tu69.

Table 3.4 Radionuclide concentrations in surface and groundwater
in the vicinity of a proposed uranium project

Radionuclide	Concentrations, pCi/l					
	Location 1	Location 2	Location 3	Location 4	Location 5	Location 6
<u>Surface Water</u>						
U-238	9.8	4.0	2.5	1.8	1.6	---
Ra-226	0.4	0.5	<0.1	0.08	<0.1	---
Rn-222	145	108	42	<4	<4	---
Th-230	<0.1	0.2	0.3	<0.1	<0.1	---
Th-232	<0.1	<0.1	<0.1	<0.1	<0.1	---
<u>Groundwater</u>						
U-238	2.0	4.5	2.3	3.2	3.8	1.2

Source: NRC79a.

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Table 3.7 Concentrations of selected elements in groundwater at six locations
in the vicinity of a proposed uranium project

Element	Concentrations, mg/ℓ					
	Location 1	Location 2	Location 3	Location 4	Location 5	Location 6
Aluminum	<0.1	1	0.8	14.7	0.2	10
Antimony	<0.01	<0.01	<0.01	<0.01	<0.01	<0.01
Arsenic	<0.01	<0.01	<0.01	<0.01	0.01	0.04
Barium	<0.1	0.5	0.5	0.2	0.3	0.7
Beryllium	0.001	<0.001	<0.001	0.003	0.001	<0.001
Cadmium	<0.001	<0.001	<0.001	<0.001	<0.001	<0.001
Cobalt	<0.01	<0.01	<0.01	0.01	<0.01	<0.01
Chromium	<0.001	0.003	0.003	0.004	0.001	0.003
Copper	0.003	0.003	0.006	0.027	<0.001	0.006
Lead	0.003	<0.001	<0.001	0.003	<0.001	0.002
Mercury	<0.0004	0.0012	0.0005	<0.001	<0.0004	0.0017
Molybdenum	0.001	0.002	0.002	0.001	0.001	0.003
Nickel	<0.01	<0.01	<0.01	0.02	<0.01	<0.01
Selenium	0.06	<0.01	<0.01	<0.01	0.13	<0.01
Vanadium	0.01	0.01	0.02	0.01	0.02	0.08
Zinc	4.4	0.07	0.21	0.07	0.01	0.02

Source: NRC79a.

3.2 Water-Related Aspects of Uranium Mining

3.2.1 Previous and Ongoing Hydrologic and Water Quality Studies Related to Uranium Mining

In the late 1950's and early 1960's, the U.S. Public Health Service conducted field studies to determine the water quality impacts of the uranium mining and milling industry. The studies emphasized uranium milling rather than mining. The Federal Water Pollution Control Administration conducted extensive stream surveys to assess the effects of uranium milling (but not mining) on the main stem and principal tributaries of the Colorado River. Subsequent stream survey work in Colorado by the Water Pollution Control Commission and the U.S. Geological Survey (Mo74, We74) mentioned a portion of the Uravan Mineral Belt and uranium mines therein, but the work did not emphasize uranium mines. Significant amounts of acidity and total trace metal concentrations were found in streams from 18 different mining areas. Dilution and chemical precipitation below mine drainages decreased concentration and increased the pH. Given enough time and distance, the streams recover naturally, but the accumulations of trace metals in the sediments increase. Field observations in 1971-72 of streams in most of Colorado indicated that approximately 724 km of streams in 25 different mining areas were adversely affected by mine drainage (We74).

Discussions of the impacts of uranium mining on water quality or quantity are incidental in numerous impact statements and environmental reports prepared by industry and (or) the U.S. Nuclear Regulatory Commission as an integral part of licensing or relicensing uranium mills. Coverage on mining is usually minor as the principal focus is on milling impacts. The same is true for the recently prepared generic EIS on regulation of uranium milling (NRC79b).

Radiochemical assessment of surface and groundwater in uranium mining districts of New Mexico is done by self-monitoring programs associated with NPDES permits. Also, radiochemical assessment studies have been funded recently by the New Mexico Environmental Improvement Division and U.S. Environmental Protection Agency. Self-monitoring, particularly in the pre-operational phase, characterizes mining and milling operations in all of the concerned States. These, together with results of surveys by State personnel, have resulted in extensive files of water quality data, flow measure-

ments, field observations of mine conditions, and exchanges between industry, regulatory agencies, and the public. Rarely are the data assembled and interpreted for dissemination outside a given agency. States experiencing rapid growth in uranium mining and milling are undoubtedly placing first priority on activities directly related to licensing, monitoring, and otherwise implementing regulations. Unfortunately, there is no concerted effort to prepare broad assessments of the cumulative impacts of mining and milling. Texas, New Mexico, and Wyoming are cases in point. Critical review and synthesis of these types of data can produce rather useful information. For example, the publication "Water Quality Impacts of Uranium Mining and Milling Activities in the Grants Mineral Belt, New Mexico" (EPA75) addresses the groundwater and surface water changes as the result of extensive uranium mining and milling production in a relatively confined area.

Some states have since initiated review of their data files, conducted field studies, and, in some cases, contracted study teams to investigate similar water quality changes. For example, a recent report by the Wyoming Department of Environmental Quality summarizes 16 years of aqueous radium and uranium data. The study reports that significant amounts of Ra-226 and uranium were present in surface water in the Shirley Basin as a result of inadequate mine water treatment (Ha78).

In Texas, surface water and groundwater monitoring conducted by industry, as well as by State and Federal agencies, reveal little or no change of chemical quality attributable to uranium mining and milling (Ge77, Ka76). This conclusion is based on 586 samples collected from 198 stations over a period of 39 years but primarily from 1961 to 1975. The State monitoring program by several agencies is continuing, but either summary reports are not issued or are two years overdue, depending on the agency. Not all of the findings exonerate the industry. One survey (It75) showed that none of the mine water from the 10 lakes that were sampled was suitable for human use. The lakes were also unsuitable for irrigation due to mineralization of the water by sulfate, chloride, and TDS. One of the 10 was suitable for stock watering.

The conditions or limits in the NPDES permits consider the quality of water being discharged, the quality of receiving water, and available, practical, treatment technology. Industry is required to monitor the discharges

on a periodic basis, usually daily, weekly, or monthly, and report results to EPA. The NPDES permits and related monitoring data help to estimate the quantity and quality of discharge allowed to enter the off-site environment.

Intensive studies of the influence of uranium mining on water quality and availability have not been conducted. Most investigations to date have been site-specific, of relatively short duration, and focused on the influence of surface discharge or subsurface seepage on water quality. Baseline studies from specific projects typically consist of quarterly or semi-annual sampling and are oriented toward milling instead of mining activities. The effects of dewatering on depleted water supplies or on water quality shifts in the aquifers of an area are rarely considered, and, then only on a mine-by-mine basis. Rarely are soil, stream sediment, and biologic samples collected in the preoperational period for radiologic analysis.

3.2.2 Mine Water Management

Figure 3.3 shows a scheme for considering the fate of water discharged from underground and open pit mines, including principal sources and sinks, most of which affect both water flow and quality. Broken lines in Fig. 3.3 indicate less important sequences with respect to water quality. For example, those mines that handle all water by on-site evaporation are likely to involve small volumes of water, and impacts on groundwater as a result of seepage are also likely to be small.

Mine drainage is surface water or groundwater flowing from a mine or an area affected by mining activities. Mine related point and nonpoint pollution sources can contaminate both surface water and groundwater throughout all phases of mining, that is, during mineral exploration, mine development, mineral extraction, processing, transport, and storage, and waste disposal. While mine-related point pollution sources usually include only milling and processing plant discharges and mine dewatering discharges, nonpoint sources can occur during any or all phases of mining. The chemical and physical characteristics and the mode of transfer of these nonpoint sources are variable and depend upon, among other things, the mineral being mined, its geo-

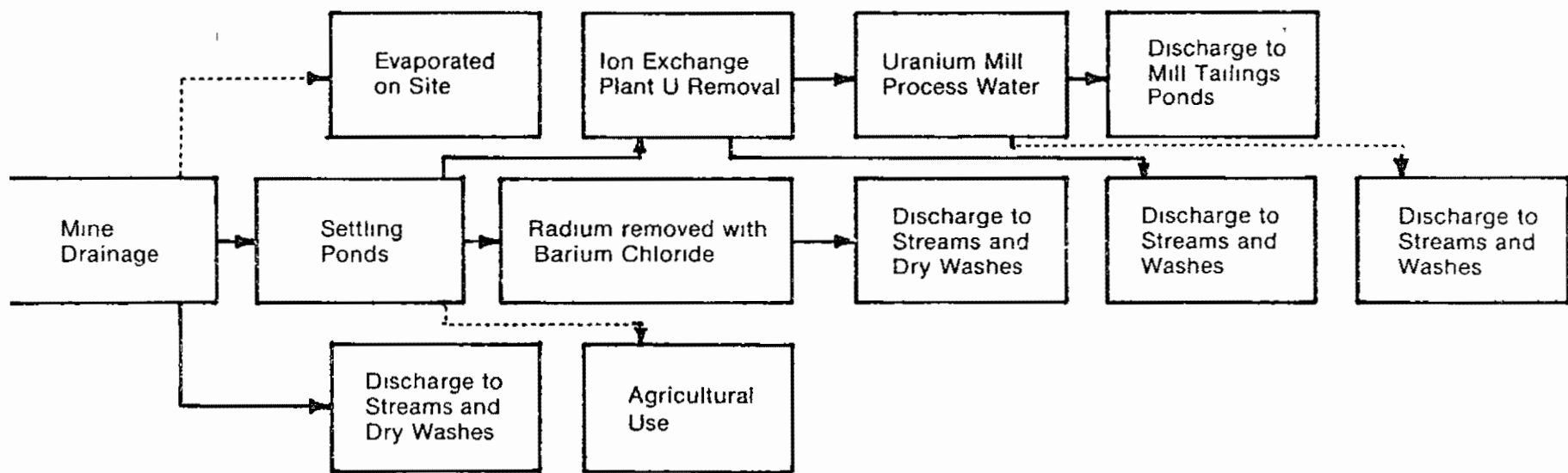


Figure 3.3. Disposition of drainage water from active surface and underground uranium mines

logic environment, the interrelations of all associated hydrologic systems (both surface water and groundwater), and the type of processing, transportation, storage, and waste disposal methods. Some mine-related nonpoint pollution sources are as follows (EPA77a):

1. suspended solids carried by immediate surface runoff
2. dissolved solids carried by immediate surface runoff
3. suspended and dissolved solids in proximate subsurface water seepage
4. dissolved solids in groundwater recharge
5. dissolved solids in groundwater discharge
6. uncontrolled contributions from mine-related point sources:
 - a. high instantaneous concentrations of regulated pollutants in excess of effluent discharge guidelines, but falling within the NPDES instantaneous and daily average discharge limitations
 - b. unregulated minor contaminants in point source discharges which are not specifically included under NPDES effluent limitations
 - c. untreated mine dewatering discharges during or following major storm events (NPDES point source treatment systems may be bypassed during storm events of greater than a 10-year, 24-hour intensity)
7. reclaimed mine area and undisturbed area drainage diversion discharges
8. surface water and groundwater contamination and degradation induced by mine-related hydrologic disturbances and imbalances

Typically, waters affected by mine drainage are chemically altered by an increase in iron, sulfate, acidity (or alkalinity), hardness, TDS, and various metals, and are physically altered by an increase in suspended solids such as silt and sediment (Anon69,Hi68).

Many but not all uranium mines dewater at rates of 1 to perhaps 20 m³/min. Typically, the water from the mine goes to settling ponds and then either to the mill or a nearby stream, dry wash, river, etc. Depending on the amount of mine water recycled in the mill and the amount of water pumped from the mines, there may or may not be any release to streams or arroyos. In at least one instance in the Grants Mineral Belt, mine water is totally recycled through the mine to enhance solubilization of uranium which is removed with ion exchange columns. Large evaporation ponds and some seepage losses help maintain a water balance and minimize releases to streams, arroyos, etc. Increasing competition for water in the western states is likely to induce maximum mine water reuse (in the mill), reinjection, or use

for potable supplies (Hi77) or power plant cooling.

When mill tailings ponds are used for final disposal of mine water, there is significant addition of chemical and radiochemical contaminants to the water in the course of milling. After treatment to reduce suspended solids, mine water may be recycled for use in the milling process or released to nearby streams, necessitating radium removal and reduction of suspended solids. Dissolved uranium in mine water, if present in concentrations exceeding about 3 mg/l, is recovered by ion exchange columns. Settling ponds at the mines remove suspended solids. The water is then conveyed to receiving streams or to the mills for uranium recovery and (or) to satisfy mill feed water requirements. There are rather rigid requirements for release to surface water compared to groundwater.

A recent survey of 20 U.S. uranium mills (Ja79a) found large variation in the degree of water recycling. Where mine water is readily available, it probably is reused less than in water-short areas. More efficient water use by uranium mills possibly could increase the amount of (relatively) high quality mine water being discharged to the environment; lessen adverse impacts of mill tailings disposal by reducing the amount of liquid; and make mine water available as a source of potable water (after treatment) in water-short areas such as Churchrock and Gallup, New Mexico (Hi77). To date, water quality deterioration related to seepage and accidental release of tailings to surface streams has received the most study and has been the focus of regulatory programs. In the future, it is likely that water quantity issues will become increasingly important, particularly in areas where water supplies are already limited and where extensive dewatering necessarily accompanies mining.

Of 20 uranium mills surveyed, 6 reported part or all of the mill feed water came from mine drainage (Ja79a). In New Mexico, 19 of 30 mines surveyed by the State Environmental Improvement Division (J. Dudley, written communication) had off-site discharge to arroyos ranging up to 19 m³ per minute. Those mines with no discharge utilized evaporation ponds or used the water for dust control. Most of the mines discharged to arroyos. In several instances, however, water was piped to a nearby mill at flow rates of 5 to 8 m³ per minute. Relatively small quantities of mine water were used for sand backfill of mines, in-situ leaching of old workings, and irrigation of grasslands. In summary, New Mexico mines discharge 66 m³ per minute off-site.

Of this, 12 m^3 per minute is routed to mills, and the balance is discharged directly off-site from the mine. Average discharge to streams and arroyos for the active underground mines in the Grants Mineral Belt on the whole was $1.8 \text{ m}^3/\text{min}$, whereas 12 mines in the Ambrosia Lake District averaged $1.7 \text{ m}^3/\text{min}$. In New Mexico, all mines discharging to an arroyo practice radium removal with approximately 90% efficiency. Uranium removal from mine water discharge occurs in all but two active mines. Future trends are likely to reflect increased discharge from the mine to the environment. Settling ponds, radium removal, or both will be used to meet discharge permit requirements.

In Wyoming, discharge from both surface and underground mines may be used as process water for uranium mills, discharged to surface streams, or used for irrigation. For example, at the North Morton underground mining operation, approximately 2 m^3 per minute of mine water discharge will be used to irrigate 800 hectares of alfalfa. At the South Morton surface mine operation, a like amount of discharge will become mill feed water.

A survey of all active U.S. uranium mills showed that 14 of 20 make no use of mine water (Ja79a). This may reflect mines where water simply is not encountered or the fact that mines and mills are not co-located. Most mills depend on deep wells, except in New Mexico where mine water is the main mill water supply. Table 3.9 summarizes water sources for U.S. uranium mills. Proposed NRC regulations on mill tailings disposal (44 Fed. Reg. 50012-59) purport to make long-term tailings isolation the primary consideration in mill siting. In areas subject to severe natural erosive or dispersive forces, this may mean that mills cannot be sited in the vicinity of mines. This may have effects on use of mine water for milling.

Table 3.9 Summary of feed water sources for active U.S. uranium mills

Water Source	No. of Mills
Rivers, Reservoirs	4
Wells	8
Springs	1
Unknown	1
Mine Water	3
Mine Water and Wells	3
	<u>20</u>

Source: Ja79a.

Although underground mining is now dominant in the Grants Mineral Belt, the greatest number of mines are small stripping operations that have long been inactive. This type of mining activity has apparently had little adverse impact on water resources. Few data are available on drainage assessment of large open pit mines such as the Jackpile-Paguate. The St. Anthony pit discharges about 0.076 m^3 per minute. Usually, the ore is above the water table. Any water present on the mine floor presumably is flood runoff or discharge from a nearby underground mine. Other strip mines in the Mineral Belt were not studied; hence, no conclusions were drawn (J. L. Kunkler, USGS, in preparation).

Mine dewatering is done either by pumping the mine pit/shaft directly or by drilling high capacity wells peripheral to the mine and pumping a sufficient volume of water to at least partially dewater the sediments. Because of the great volume of water that must be removed from an aquifer, the latter method is impractical for deep underground uranium mines. This is particularly true for the artesian aquifers of most of the deeper mines in the Grants Mineral Belt. More commonly this method is reserved for shaft sinking and open pit mines to depths of several hundred feet. Most underground mines are dewatered by pumping the water that collects in the mine itself. Borings ("longholes") made into the ore body for assay work and explosives facilitate drainage. There is considerable difference in the quality of water depending on the dewatering method used. Water removed from wells adjacent to the mine typically is representative of natural quality, but water removed from the mine can be high in radionuclides, stable elements, and suspended solids. In large part this is due to the disruptive nature of mining. However, more subtle, chemical processes of oxidation and bacterial action, aided by evaporation and free flow of air in the mine, are also operative.

The extent to which uranium exploration adversely impacts water resources is not well understood. Land surface disruption from drilling pads and access roads obviously affects erosion rates and results in mud pits and piles of contaminated cuttings on or near the land surface. Subsurface effects are less obvious. A potentially serious one is interaquifer connection via exploratory boreholes. In Wyoming, 6 million meters of exploratory drilling took place in 1979. Although State law requires mining companies to plug the holes after drilling, it is common practice to install only a surface plug and to rely on the drilling mud to effect a seal at depth.

Similar situations are likely in New Mexico and Texas. Shortages of funds and personnel to oversee proper completion and abandonment exist at the State level.

Hydraulic effects of water released from mines, whether from pumping or gravity flow, include increased surface discharge, recharge of shallow aquifers by infiltration, and decline of static water levels in formations intersected by the mine or related cone of depression. Of most concern are the effects relating to mine water discharge on downstream users and any influences, direct or indirect, of pumping/dewatering on water quality in the ore body and contiguous strata. In some locations, the Grants Mineral Belt, for example, the ore body is also a major regional artesian aquifer; hence, dewatering affects present water levels and will affect water levels at least to the year 2000, with complete recovery taking much longer. The extent and significance of uranium mine dewatering are as yet poorly documented. Recent studies have been made in New Mexico where dewatering is of concern because of the influence on regional groundwater availability for municipal use and in relation to return flows to the San Juan River (NRC79b; Ly79). In Wyoming, static water levels in wells on ranches adjacent to uranium mines owned by Exxon, Kerr-McGee, Rocky Mountain Energy Co., and other companies southwest of Douglas and between Pumpkin Buttes and Douglas have reportedly dropped 7 to 10 meters (Anon79). Water quality changes associated with dewatering generally are unknown and not specifically monitored regardless of the mining area location.

Water quality associated with dewatering is generally good, although suspended solids may be high, as expected. Discharge from dewatering wells will be low in suspended solids because of filtering by soil and rock aquifers. Overall water quality from dewatering wells, particularly for underground mines, is likely to be representative of ambient conditions in the ore body and, to a lesser extent, the adjacent formations that may also be dewatered.

Recent USGS work on groundwater in the San Juan Basin Region has indicated that mining expansion will have a significant impact on the water yield of the Morrison Formation (Ly79). In this study, although no water quality data are derived, the recharge and mine dewatering parameters that impact the expected drawdowns in the aquifer imply that a total of 7.03×10^8 m³ of water will be produced by the 33 planned or announced mines by the year

2000. If the projected development of 72 mines occurs, dewatering would exceed $1.48 \times 10^9 \text{ m}^3$. The model also estimates that flow in the San Juan River will decline very slightly ($0.05 \text{ m}^3/\text{min}$). Similarly, flow in the Rio Grande Valley would be reduced by $0.85 \text{ m}^3/\text{min}$. The impacts will continue after mining and dewatering cease.

Table 3.10 summarizes New Mexico uranium mine discharge in relation to mine type, depth, and status (active or proposed). Note that projected mining is primarily underground and represents an average increase in mine depth of 275 percent and an increase in dewatering rate from 2.4 to 13.8 m^3 per minute. One would expect numerous water quality and quantity issues to arise if these projections materialize. For example, competition for water supply is likely to be widespread throughout the Upper Colorado River Basin, and uranium mines/mills are already relatively large water users. Dewatering and discharge require no water rights under State water laws in New Mexico, but the water is essentially wasted. Use of water in mills constitutes a beneficial use of water, and state water laws therefore require filing for water rights. Such filings may be denied upon protest from existing water users.

Inactive uranium mines and related wastes also influence water quality, particularly as a result of chemical and physical transport by surface water runoff. The main reasons why mine waste piles erode more quickly than undisturbed soils are lack of topsoil, steep angle of slopes, presence of toxic elements and buildup of salt in the near surface, and poor water retention characteristics. Usually, inactive surface and underground uranium mines are not a source of direct discharge of water, be it contaminated or of ambient quality, because of the low rainfall-high evaporation characteristics of the western uranium regions, static groundwater levels deep below the land surface in mining areas, and, in a few instances, recontouring of mined lands such that drainage is internal. Whether mines contaminate groundwater by groundwater leaching or by recharge contacting exposed oxidized ore bodies is poorly documented. Preliminary feasibility studies by the U.S. Geological Survey (Hi77) indicate generally good quality water from one inactive underground mine in the Churchrock area of New Mexico. It is possible that this water may be used as a municipal water supply for Gallup, New Mexico.

Table 3.10 Current and projected uranium mine discharges
in the Grants Mineral Belt, New Mexico

Mine Type	Number of Mines	Average Depth (m)	Average Discharge (m ³ /min)
<u>Active</u>			
Underground	33	248	2.42
Open pit	3	48	0.045
<u>Proposed</u>			
Underground	46	681	13.8
Open pit	0	N/A	N/A

Source: Environmental Improvement Division, State of New Mexico.

For most uranium regions, the volume of discharge from inactive mines to surface water bodies, though poorly documented, is believed to be less significant than that from active mines. The degree to which inactive mines contribute contaminants, directly or indirectly, to adjacent water resources can only be qualitatively assessed. The significance of inactive mines is highly dependent on regional setting and mine type.

Inactive surface mines in Texas are, with rare exception, not a source of direct discharge to surface water. It is unknown if there is any adverse impact from standing water in the mine pits, the most recent of which have been final-contoured with an internal drainage plan. Various observers suspect that water quality deteriorates when overland flow crosses mine spoils associated with overburden piles (It75 and He79). Water in the mine pits is unsuitable for potable and stock use due to high stable element contents, but it is generally acceptable in terms of radioactivity. Water in Texas open pit mines is a combination of runoff and groundwater. Before release from a mine, water is put in retention ponds to reduce total suspended solids. Holding ponds are used for storing mine water, and discharge is not allowed unless such discharge does not adversely affect the receiving

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Environmental problems associated with alkaline and saline drainages are not well documented (Hi73).

Water quality impacts from uranium mining are a function of both quality and quantity of discharge. Underground and surface mines commonly require dewatering prior to or during the ore removal phase, although there is considerable variation from one area or mine to another. In New Mexico, large and medium sized mines are essentially dry, whereas mines in Texas and Wyoming require extensive dewatering. Regardless of location, underground mines rarely are dry and many require extensive dewatering. Considering the variety of water management measures, regional differences in contaminants and receiving waters, and geochemical characteristics of ore bodies, detailed discussion of the effects of mine drainage or mining, in general, on water quality must await further site - or area - specific study. It is questionable if sufficient data on mine drainage exist to assess effects on biota and the fate of contaminants in surface or sub-surface water bodies.

Limited data from Colorado, Texas, Wyoming, and New Mexico suggest adverse impacts on water quality from discharge of mine water. Effects of dewatering on deep groundwater quality are very poorly documented; hence, no conclusion as to relative significance is drawn. With respect to surface water resources, discharge of mine water and overland movement of water and suspended or dissolved contaminants may be significant. Because of the dearth of data on overland flow, emphasis herein is on contaminants discharged via mine drainage water. Substantial studies to evaluate sediment yield and quality from lands mined for uranium, particularly from areas of surface mining, have not been conducted, although recent work in Texas (He79) is a notable exception.

Elements such as uranium, radium, molybdenum, selenium, zinc, and vanadium may be enriched in point and nonpoint discharges from uranium mines. The dispersal, mobility, and uptake of such elements are directly relevant to the subject of this report. We reviewed selected literature and field data to at least qualitatively understand what processes and elements are most significant and to thereby strengthen some of the underlying source term assumptions in the transport and health effects modeling. Despite the annual chemical load introduced to ephemeral streams by both dissolved and suspended constituents in mine effluent and overland flow from mined lands, waste piles etc., a number of processes affect the concentrations in the ambient environ-

ment. These include dilution, suspended sediment transport, sorption and desorption, precipitation, ion exchange, biological assimilation or degradation, and complexation.

3.2.3.1.1 Dilution and Suspended Sediment Transport

In many uranium mining regions there is flooding or flash flooding. Such storms may well be the only runoff event for a year or more at a time. It is worthwhile then to consider some of the effects of such events on mobilization of contaminants associated with uranium mining wastes. Typically there is significant interdependence between the physical and chemical processes.

A principal physical process is dilution. This will reduce concentrations of pollutants released to surface waters, but is considered to play a relatively minor role over the short term for water percolating through the soil to sources of groundwater in arid or semiarid regions.

Transport of suspended sediments in floods is another dominant process. Suspended load is largely a result of physical, hydraulic processes, hence elements that are rapidly and thoroughly removed from solution as a result of solubility limits, precipitation with other ions, ion exchange, and sorption may well be transported in the suspended load. In metal mining areas of central Colorado, total and dissolved metal loads in streams are greater during high flow periods, apparently a result of flushing from mines and tailings piles and scouring of chemical precipitates from stream substrates (Mo74). Typically, total and dissolved loads decrease downstream, regardless of discharge. Increase in iron in the downstream direction reflects scouring of precipitate--an amorphous, hydrated ferric oxide. Dispersal occurs for quite a distance downstream.

3.2.3.1.2 Sorption and Desorption

Sorption can play an extremely important role in purifying waters, particularly if infiltration or percolation is involved. This is especially true when contaminant concentrations are too low to undergo precipitation reactions. Virtually every ionic species will be sorbed and removed to some extent except for chloride and, to a lesser extent, sulfate and nitrate. These seem to pass through soils and alluvium without significant sorption (Ru76). Sorption processes can be highly specific, depending on the type of

contaminant and the physical and chemical properties of both the solution and the porous medium.

There have been numerous laboratory studies on the sorption, leachability, and mobility of stable elements in various types of soil. A recent review of the literature on radionuclide interactions in soils specifically discusses radium, thorium, and uranium, which are especially pertinent to uranium mining (Am78). Distribution coefficients, the ratio of concentration in soil to that in water, ranged between 16 and 270 for uranium in various soil-river water systems, between 200 and 470 for radium, and on the order of 10^5 for thorium at pH6. These observations appear consistent with the generally accepted ideas that uranium is relatively mobile, thorium extremely immobile, and radium somewhere in between in natural water systems (NRC79b; Ku79; Ga77a).

Adsorption is believed to be important for cadmium, copper, lead, and nickel, insofar as these are transported in the suspended fraction, whereas manganese and zinc are primarily in the dissolved fraction. Adsorption of metals onto precipitated manganese oxides or hydroxides at elevated pH is probably insignificant in the case of most uranium mine discharges insofar as these are alkaline and, furthermore, discharge to or co-mingle with other streams that are alkaline.

Radium sorption and desorption tests done on uranium mill waste solids and river sediments collected from several locations in the Colorado Plateau (Sh64) and in Czechoslovakia (Ha68) showed that leaching is primarily controlled by the liquid-to-solid ratio, i.e., the volume of leaching liquid per unit weight of suspended solids. Natural leaching from mining and milling waste solids freely introduced to rivers in the past is one of the major factors in radium contamination of rivers (Ru58). Although settling ponds are now used to remove or at least reduce suspended solids from active mine discharges, the dissolved radium load sorbed on sediments presents a source term that may be somewhat analogous to river sediments contaminated by dissolved and suspended milling and mining wastes. The manner in which the radium is mobilized and the significance is poorly understood and bears further investigation. Apparently there is a "leap frog" transport mechanism involving combined chemical and physical weathering processes. There should be a marked downstream attenuation of both dissolved and sorbed/precipitated radium inventories insofar as sediment burial and dilution take place and leachability limits are reached, i.e., no more radium can be removed regard-

less of the duration, frequency, or intensity of agitation. Should the stream eventually discharge into a reservoir, it is unlikely that renewed leaching will take place. Shearer and Lee (Sh64) did not account for some factors that may be locally significant, such as bio-uptake along the stream/river, use of water for irrigation, number of uranium facilities discharging, and other local factors.

Experiments were conducted in Japan (Ya73) to determine uranium adsorption and desorption using carbonate solutions and three soil types (alluvial, sandy, volcanic ash). Very high adsorption ratios and very low desorption ratios of uranium characterized the various soil types in contact with stream water and help explain the decrease in soluble uranium with flow distance from mines (Ma69). When wastewater flows into streams at the maximum permissible concentration (1.8 mg U/l) recommended (ICRP64), Yamamoto et al. (Ya73) conclude that the uranium behaves as a uranyl carbonate complex anion and that essentially complete sorption readily occurs in the presence of (Japanese river) water which contains 15 to 39.9 mg/l bicarbonate. Since this is similar to concentrations in surface waters of uranium regions in the western states, similar results are expected.

Sorption or desorption of heavy metals such as Co, Ni, Cu, and Zn in soils and fresh water sediments occurs in response to the aqueous concentration of metal, aqueous concentration of other metals, pH, and amount and strength of organic chelates and inorganic complex ion formers in solution (Je68). Other controls on the heavy metal concentrations in soil and fresh water include organic matter, clays, carbonates, and oxide and hydroxide precipitates.

To what degree solubility acts as a limit on stable element concentrations in natural waters is unclear. The crystallographic form or even the chemical composition of a precipitate are often unknown. Elements such as iron, aluminum, manganese, and titanium form insoluble hydroxides and are likely to exceed equilibrium solubility limits (An73). Hem (He60) partially disagrees, saying "it is not unreasonable to assume equilibrium for the iron species in water." Whether mine discharges or overland flow from mined areas are in equilibrium is unknown, but it is doubtful considering the underground or flash-flood origin of such waters. The non-equilibrium aspects of certain peak runoff events has been documented for major streams of the world (Durum and Haffty, 1963). Metals such as iron, aluminum, manganese, and titanium,

which readily form rather insoluble hydroxides as particulates or colloids, may be dissolved from suspended minerals during high flow conditions. Organics present in such flood waters may assist through formation of soluble complexes. Resulting metal concentrations may be higher than solubility and redox relationships alone would indicate.

3.2.3.1.3. Precipitation

Probably one of the most significant processes affecting stable element solubility in natural water systems is adsorption on hydrous ferric and manganese oxides. Jenner (Je68) believes this is the principal control on the fixation of Co, Ni, Cu, and Zn (heavy metals) in soils and fresh water sediments. For example, ferric hydroxide adsorbs one to two orders of magnitude more SeO_3 per unit weight than clays, and 90 to 99 percent adsorption is possible at a pH of seven to eight typical of most western streams (Ho72). At neutral or slightly alkaline pH, both iron and manganese are poorly soluble in oxidizing systems and, in general, exhibit very similar chemical behavior, although manganese is slightly more soluble. Fixation of selenium in soils, particularly by iron oxide or as ferric selenite, renders it unavailable to agricultural and forage crops, although specific selenium-accumulating plants can remove the element and, upon decomposition, release it in water soluble forms, such as selenate and organic selenium compounds, available to other plants (Ro64). The behavior and mechanism of selenium adsorption (as selenium oxyanion) by hydrous ferric oxides is readily extended to the interpretation of other similarly bound minor elements (Ho72). Mobile selenium oxyanion in slightly alkaline waters might be carried to streams by surface runoff or in groundwater. Selenite selenium sorbed upon ferric hydroxide should be transported in surface waters at neutral or slightly acid pH. Other metals forming highly insoluble hydroxides in the pH range of 6 to 9 include copper (above pH 6.5), zinc (above pH 7.5), and nickel (above pH 9). Molybdenum is thought to hydrolyze to the bimolybdate ion under acid conditions and precipitate with iron and aluminum. Aerobic or oxidizing conditions in the vadose zone are favorable for the development of many of these oxides (Cu, Fe, Mn, Hg, Ni, Zn, Pb). Reducing conditions deep in saturated zones generally lead to increased mobility of these metals.

Reducing conditions that can exist in the presence of organic material (bituminous or lower ranking coals, anaerobic bacteria, fluidized humates)

can lead to precipitation reactions favorable for removing contaminants from mine waters. Reduction of uranium to the quadrivalent state and its fixation on clays would play the major role in protecting groundwater supplies from uranium if the appropriate reducing agents were present in soils (Ga77a; Ku79). Inorganic reducing agents could include ferrous iron and hydrogen sulfide produced by the action of an aerobic bacteria on sulfates. Natural reducing conditions can also, theoretically, cause the formation of such native elements as arsenic, copper, mercury, selenium, silver, and lead, which are all quite insoluble in their elemental form (Ru76). Hydrogen sulfide or other sulfides, if available, will serve to reduce the concentrations of such metals as arsenic, cadmium, copper, iron, lead, mercury, molybdenum, nickel, silver, and lead.

The metal-scavenging of hydrated iron oxide precipitate has been documented in a mined area of Colorado where relatively acid schists and gneisses give rise to acid runoff that dissolves large quantities of aluminum, magnesium, and zinc. Runoff from a nearby drainage basin underlain by basic rocks containing base and precious metal veins carries considerably less metal. However, manganese oxide precipitated with iron oxide contains large quantities of metals. Ferric hydroxide precipitates from aerated water solutions containing more than 0.01 ppm iron at pH values of 4.5 and above, aluminum hydroxide precipitates in the pH range of 5 to 7, and manganese hydroxide precipitates above pH8 (He60; Ch54). Considering the alkaline pH of most uranium mine discharges and overland flow from non-point sources such as mine waste piles, precipitation of iron and possibly manganese seems certain. The scavenging effect of iron hydroxide at neutral to alkaline pH is considerably less than that of manganese hydroxide precipitate.

The extensive studies of mine drainage in Colorado by Morgan and Wentz (1974) revealed the effects of solubility on stable element transport. In the downstream direction, dilution and neutralization of the acid mine drainage by bicarbonate caused dissolved metal to decrease due to dilution, chemical precipitation, and probably adsorption onto ferric hydroxide precipitate. The latter creates a coating on the stream substrate for a considerable distance during low flow periods. Subsequently, flood events scour and transport the precipitates. Manganese and zinc remain primarily in the dissolved phase for a considerable distance, whereas cadmium, copper, iron, lead, and nickel concentrate in the suspended fraction and, when turbulence

decreases, precipitate. The mobility sequence for the metals studied in Colorado generally follows the order $Mn \approx Zn > Cu > Cd > Fe > Ni > Pb$. Ferric hydroxide precipitation and scavenging seems to be more important at neutral than at acidic pH's (Je68).

3.2.3.1.4 Biological Assimilation and Degradation

Biological uptake and the role it has on stable element concentrations in water is not predictively understood (An73). Plant uptake of stable elements and resulting phytotoxicity is not merely a function of how much is present in the soils or water. In the case of arsenic, the chemical form of arsenic appears more important than the total soil arsenic (Wo71). For example, water-soluble arsenic in soil created more phytotoxic effects than those with no detectable water-soluble arsenic. Soils high in reactive aluminum remained less phytotoxic, despite heavy applications of arsenic, than soils with low reactive aluminum. Selenium in soils can be present as elemental selenium, selenates, pyritic selenium, ferric selenites, and organic selenium compounds of unknown composition. Selenates and organic compounds are most available to plants, although slow hydrolysis of the other forms can occur such that they become available for plant uptake. The importance of water soluble selenium versus total selenium as the major factor affecting plant uptake has been demonstrated (La72; Gr67). Where sufficient selenium is present in plant-available form, all species will take it up in sufficient amounts to be harmful to animals (La72). Naturally occurring soils containing such available forms are geographically confined to semiarid regions or areas of impeded drainage. Such soils are not hazardous to humans and only locally are they a threat to animals.

Despite numerous examples of high selenium (up to 2.7 ppm) in surface water, particularly that associated with drainage from seleniferous soils in agricultural areas, Rosenfeld and Beath (Ro64) reported only a few cases of water-related selenosis in man or livestock. Water high in selenium is typically unpalatable to livestock and certainly to man. Lakin (La72) concluded that environmental contamination due to selenium is increasing, but hazardous concentrations are unlikely; mining and industrial wastes may cause local problems; and the effect of added selenium in waters in combination with other contaminants bears further study.

Uranium uptake by several species of native plants in the southeastern

Utah portion of the Colorado Plateau varied, sometimes strikingly, with the species, time of year, part of plant, availability of uranium in the soil, and chemical composition of the underlying rocks (Ca57). The type of rooting system and the soil moisture conditions also were influential. In some cases, there was no consistent relationship between the amount of uranium in the soil versus that in the plant ash. Plants are much less selective with respect to cadmium uptake, and it has been conclusively demonstrated that plants absorb cadmium from cadmium containing solutions and soils (Pa73; Fu73). Phytotoxic effects vary considerably with plant species. Cadmium and zinc sulfides tend to concentrate in the organic matter of soils. Upon oxidation to sulfate, plant availability increases along with solubility. Under alkaline conditions (pH8), cadmium is taken up rapidly by biota and by sediments. However, modeling of cadmium transport and its deposition in aquatic systems is very complex and encompasses many variables, most important of which are pH, carbonate content, chemical form, and competing ions.

3.2.3.1.5 Complexation

Published data on Gibbs free energies, enthalpies, and entropies of 42 dissolved uranium species and 30 uranium-bearing solid phases were recently reviewed (La78). Uranium in natural waters is usually complexed with carbonate, hydroxide, phosphate, fluoride, sulfate, and perhaps silicate. Such complexes greatly increase the solubility of uranium minerals and increase uranium mobility in groundwater and surface water. In waters with typical concentration of chloride, fluoride, phosphate, and sulfate, intermediate Eh's, neutral to alkaline pH's, and the presence of phosphate or carbonate, uranyl phosphate or carbonate complexes form and increase mineral solubility by several orders of magnitude. Sorption of the uranyl minerals carnotite, tyuyamunite, autunite, potassium autunite, and uranophane onto natural materials is greatest in the pH range of 5 to 8.5. Uranium content of small streams, in particular, can exhibit wide spatial and temporal variations due to pH and oxidation state of the water, concentrations of complex-forming species such as carbonate or sulfate, and presence of highly sorptive materials such as organic matter, certain metallic hydroxides, and clays (La78). Whereas sorption is probably a dominant control on stable element concentrations in low temperature aqueous conditions, there is insufficient information concerning specific sorbents to allow accurate prediction.

3.2.3.2 Results of Field Studies in Uranium Mining Areas

3.2.3.2.1 Colorado

Extensive studies of the effect of mine drainage on stream water quality and biota were done in central Colorado (Mo74). Although uranium was mined in 14 of the 25 areas studied, other metals were the principal products. Most of the ores were high in iron sulfides, and associated drainage was acidic. Also studied, but less intensely, was the Uravan district of western Colorado where the principal products are uranium and vanadium from Mesozoic sandstone. The Uravan Mineral Belt is different in terms of principal product and geologic features from other mining areas studied in Colorado. For these other areas, the drainage is acidic and heavily enriched in heavy metals and, therefore, somewhat atypical of most Colorado uranium mines in the Uravan area.

After a preliminary field survey of the temperature, specific conductivity, pH, stream-bottom conditions, and aquatic biota at 995 stream sites, 192 were chosen for detailed sampling and analysis during 1971-1972. The data indicate the contamination of approximately 711 kilometers of streams in 25 different areas, mostly in the Colorado Mineral Belt. The water quality effects in these areas arise from many varied causes, including active and inactive mine drainage, tailings pond seepage, drainage tunnels, and milling operations. The length of the streams affected is not absolute as it varies with the time of the year and flow conditions (Mo74).

The general findings indicate that Mn, Se, and SO_4 concentrations, and specific conductivity are poor indicators of mine drainage as natural sources can cause high values for these parameters even in undisturbed areas. Uranium mines make at least some contribution to problems of contaminated streams in central Colorado. In central Colorado, the exact impact of uranium mining on stream water quality is unknown but believed to be less important or significant in most areas as compared to impact from other mining, with the possible exception of the Boulder-Jamestown area (J. Goettl and D. Anderson, Colorado Game, Fish, and Parks Division and Water Pollution Control Commission, respectively, personal communication). Cadmium, As, and Pb exceed the U.S. Public Health Service toxicity limits, respectively, 12.5 percent, 1.4 percent, and 2.1 percent of the time. Mercury and Ag limits were never exceeded, and Cr was never detected. Iron and Mn standards were

frequently exceeded by large percentages, however, these limits are only based on aesthetics. Concerning the negative impacts of the various constituents, Cu and Zn (exceeding the limits 7.8 and 9.0 percent of the time, respectively) pose the greatest threat to resident aquatic life. Mining operations in the Uravan area are a relatively minor source of metals for the San Miguel River (Mo74). Potential problem areas are settling ponds and tailings piles associated with the mining operation. Although not a source of acid drainage, these sources did cause increased concentrations of copper, iron, manganese, nickel, vanadium, and zinc in the river. Only manganese exceeded the standard for drinking water, and no metal concentrations exceeded the biological criteria. Seepage ($0.003 \text{ m}^3/\text{s}$, pH 6.8, $3300 \text{ mg}/\ell \text{ HCO}_3$) from a mine tailings area into Atkinson Creek, a tributary of the San Miguel River, observed in December 1972 caused no adverse impacts.

Because of its size, proximity to population, and effects on surface water quality, extensive surface water quality investigations to assess the impacts of mine water discharge from the Schwarzwaldor mine have been made (EPA72). Grab samples of the mine effluent taken in 1972 revealed $15 \text{ mg}/\ell$ uranium and $80 \text{ pCi}/\ell$ radium-226. As of 1972, overflow and seepage from the settling ponds used to treat the mine effluent significantly degrade the radiochemical quality of nearby Ralston Creek. This was confirmed by both EPA and the State/Denver Water Boards monitoring program. With 20-fold dilution, Ralston Creek downstream of the mine contained $3 \text{ pCi}/\ell$ and $82 \text{ }\mu\text{g}/\ell$ dissolved radium-226 and uranium, respectively. With no dilution, as during July, concentrations were $81 \text{ pCi}/\ell$ and $20,300 \text{ }\mu\text{g}/\ell$. Influx of contaminated stream water to nearby Long Lake raised dissolved radium-226 to $0.8 \text{ pCi}/\ell$ (4-fold increase over background) and uranium to $230 \text{ }\mu\text{g}/\ell$ (20 times background). From these data, conclusions were reached that the mine water caused a 5 percent increase in the radiation dose to consumers in a local water system (based on FRC and NCRP standards and daily consumption of 1.0 liter water). If the $4.5 \text{ mg}/\ell$ uranium limit proposed by ICRP was used, the estimated dose increases to nearly 40 percent of the dose limit for a population group. Since 1972, the effluent has been treated for radium-226 and uranium removal. Trace metals analysis of water samples collected from the creek and the water treatment plants revealed concentrations comparable to or greater than those in the effluent as of July 20, 1978. Concentrations

($\mu\text{g}/\ell$) were as follows:

	<u>As</u>	<u>F*</u>	<u>Pb</u>	<u>Se</u>	<u>Zn</u>
Mine Effluent	5	1	15	<2	18
Ralston Creek (avg)	5	1.3	32	<2	56
Water treatment plants (avg)	5	0.55	97	2	146

*mg/ ℓ

3.2.3.2.2 Wyoming

We assessed the effects of mine drainage by literature review and a limited field study in the Spring of 1979. Results of the latter conclude this section of the report. The effects of mine dewatering, in situ leaching, and mill tailings seepage on surface water quality in the Shirley Basin were previously studied by the Wyoming Department of Environmental Quality (Ha78). Sixteen years of data on aqueous radium and uranium indicated significant amounts of radium-226 and uranium reached streams because of inadequate mine dewater treatment, mill tailings pond seepage, and improper operation of a precipitation treatment unit. Uranium concentrations in stream water increased 60-fold because of mine water discharge and possible tailings pond seepage. The effects of past loadings of uranium and radium on fish propagation or migration are not clear, although biologic uptake of uranium and radium has occurred. Phytoplankton, algae, and bottom fauna organisms also do not appear to have been adversely affected, but no studies have been conducted since 1962. Long-term effects of increased radioactivity levels are known and merit further study: "There exists real need for additional studies to determine the mechanisms involved in the dispersion and ultimate disposal of uranium loaded into the drainage basin...Only after additional studies have been completed, may we understand the total and long range impact that the company's activities have had on the aqueous environment" (Ha78). This latter finding related specifically to the current (1978) loading of uranium from treated mine discharge.

Previous studies by the State of Wyoming (Ha78) found that solution

mining by the Pathfinder Uranium Company noticeably affected ambient uranium concentrations in the study areas. A 1968 survey by the Department of Environmental Quality (Ha78) indicated relatively high loadings of soluble uranium and radium on stream sediments near the mine dewatering outfall. Analysis of fish skeletons indicated radium uptake corresponding to dissolved radium-226 concentration exceeding 1 pCi/g. Resampling in 1970 showed a decrease in radioactivity values in sediment but a tenfold increase in fish uptake of uranium relative to other fish populations in the basin. Radioactivity concentrations in fish tissue were highest near the mine effluent outfalls but did not constitute a major source of radioactive intake by consumers.

In June 1971, the EPA Radiological Activities Section of Region VIII (Denver) made a field reconnaissance of uranium mining and milling activities in the Shirley Basin area. Radiological analyses of water and sediment samples in the Shirley Basin and in the Bates Hole drainage basin to the north unquestionably indicated significant increases in radioactivity levels in water, sediment, and fish because of effluent discharge from mines and mill tailings. Concentrations of dissolved radium-226 and uranium in mine effluent were well above background. The discharges were not considered a source of radiation dose to the populace (residents and transients) because of remoteness and lack of water use, but toxic effects on fish were of concern (M. Lammering, written communications, 1979). Monitoring in 1972 by the Wyoming Game and Fish Department showed water quality effects as far as seven miles downstream. From 1970 to 1972, radium-226 concentrations remained stable, but uranium increased. Fish samples collected in 1972 showed increased amounts of radium in the flesh compared to the 1970 results. Soil samples from a creek that received mine effluent indicated relatively large transport and enrichment of uranium and radium. Radium in particular was enriched in the sediments and showed temporal variations indicative of successive scouring and removal, presumably in flood flows. Precipitation of uranium compounds was not apparent, probably because oxidized uranyl species are quite soluble in natural water.

To further our understanding of the role aqueous pathways play in contaminant dispersal, we monitored stable and radioactive trace elements in the Spring of 1979 in surface runoff from ore, sub-ore, and overburden piles from the Morton Ranch area of active mining in Wyoming. EPA personnel selected

the basic study areas and assisted in sample collecting. Most of the sampling, analysis, and interpretation was done under contract with Battelle, Pacific Northwest Laboratories and is the subject of a draft report (Wo79). Appendix G contains a more complete discussion and listing of the data. Since runoff samples were unavailable, the sampling program emphasized the collection of soil samples in well-defined runoff gullies originating at surface stockpiles of minerals. Where the drainage systems intersected flowing water, upstream and downstream samples were collected. Most of the samples consisted of the top five centimeters of soil in the bottom of the drainage channels and three core profiles. Samples of the source material were also collected.

Trace element and radionuclide analyses of runoff from the Morton Ranch area are primarily based on surface sediments and vertical sediment profiles from the dry stream beds, since few streams or other forms of runoff were encountered. Figure 3.4 shows the waterways surrounding the inactive 1601 pit area and the semi-active 1704 pit area.

Three vertical soil profile samples were collected, two in an erosional drainage area from the ore and waste pits south of the 1601 pit and one in an erosional drainage bed on the east side of the waste pile of the 1704 pit. The radionuclide and chemical constituents of these samples along with analyses of other soil samples are reported in Appendix G. The results indicate aqueous leaching based on the radium/uranium ratios of about 12 in redeposited material in the alluvial fan area of the drainage, compared to a corresponding ratio of 0.9 in the undisturbed (sub-surface) material in the top alluvium profile.

Trace element data also indicate limited transport of mine contamination with respect to uranium, and to a lesser extent, Se and V. The profile samples containing 140 and 21 ppm U resulted from material transported from the adjacent ore piles. The aqueous samples similarly showed no unusual characteristics indicative of mine wastes.

An aqueous sample and the soil profiles collected near the 1704 pit similarly showed no evidence of mine-related pollution or leaching of uranium. This observation is based on the water and particulate analyses and the radionuclide analyses. These samples constitute a worst case, since the sediment samples were collected in the redeposited material of the waste pile drainage and should show greater levels of pollutants there than would fin-

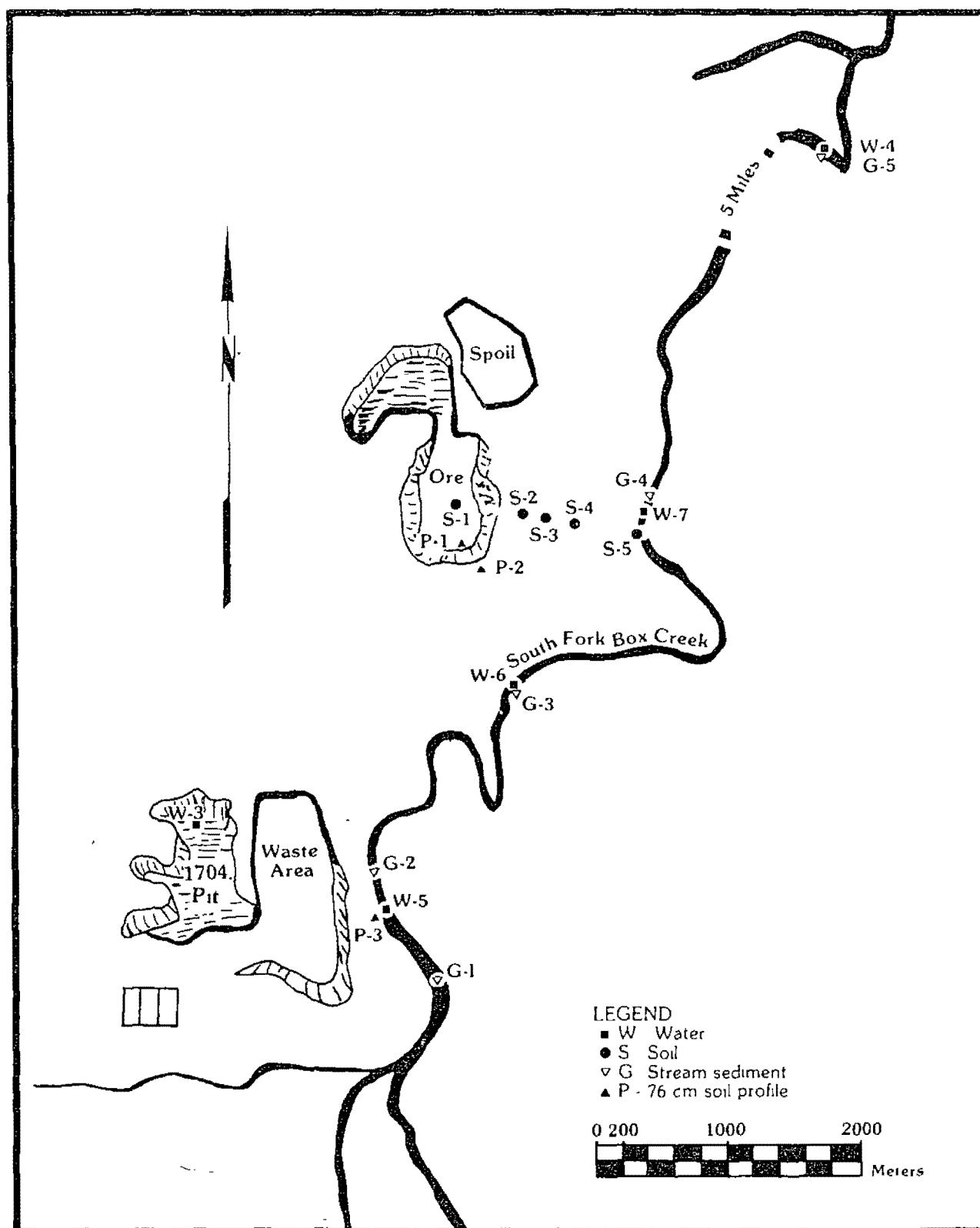


Figure 3 4 Location of mines, ore and waste storage areas and monitoring stations at the Morton Ranch mine, South Powder River Basin, Wyoming

ally reach the South Fork Creek bed.

Pollutant releases from the Morton Ranch, Wyoming uranium mining operations were not observable in water drainages of the surrounding area. The only significant movement of mine-related wastes was the transport of the stockpiled ore in erosional drainage areas on and immediately adjacent to the waste pile of the 1601 pit. Long-distance transport of these pollutants (primarily uranium) into the South Fork of Box Creek was not observable. The strongest evidence that mine wastes are a source of local soil and water contamination is the radiochemical data, and uranium in particular. Pollutant transport is almost entirely confined to the immediate area of the mines, although there has been some dispersal via water in the ephemeral streams. There is considerable disequilibrium between radium and uranium which may indicate leaching and remobilization of uranium. The possibility of natural disequilibrium in the ore body should not be overlooked.

3.2.3.2.3 Texas

A very comprehensive field and literature survey of elements associated with uranium deposits in south Texas (He79) revealed high to very high concentrations of molybdenum, arsenic, and selenium in areas of shallow mineralization; drainages adjacent to older, abandoned mines; and in some reclaimed areas. Areas of shallow mineralization have concentrations of several tens of ppm molybdenum and arsenic and up to 14 ppm selenium. Near surface material exposed by mining may have several hundred ppm molybdenum and arsenic. Waterborne transport of suspended or dissolved solids away from open pit mines resulted from mine water discharge and (or) surface runoff and erosion of abandoned spoil piles. Molybdenum from the mining areas could potentially aggravate natural soil problems leading to molybdenosis (Kab79). Additional careful study is suggested, particularly of areas receiving mine drainage as pumped water or overland flow.

Lakes or ponds associated with 10 mine locations in Karnes and Live Oak counties contained water unsuitable for drinking without prior treatment (It75). Generally, mineralization was also excessive and rendered the water unfit for irrigation. Air and terrestrial sampling revealed no health hazards from mining wastes and mined lands, but insects and other bottom fauna in the lakes concentrated radium-226 400 to 800 times the water concentration based on dry weight of the organisms.

3.2.3.2.4 New Mexico

The principal investigations of the influence of uranium mining on water quality includes EPA and contracted (Wo79) work by the U.S. Environmental Protection Agency, the Department of Interior (Ku79), and ongoing studies by the State of New Mexico (J. Dudley, oral communication, 1979). Because of the co-location of mining and milling facilities, it is difficult to identify impacts from one versus the other.

Survey of groundwater and surface water quality in close proximity to the Jackpile-Paguate, Ambrosia Lake, and Churchrock mining areas (EPA75) revealed extensive discharges of mine water to the ambient environment, use of unlined ponds for settling suspended solids from mine dewatering, use of contaminated mine water as a potable supply (one facility), and failure of all facilities discharging to streams to have a valid NPDES permit. The volume of mine discharge, particularly in the Churchrock area and from a mine near Mount Taylor, led to use of the water for irrigation and stock. In other areas of Ambrosia Lake and near Churchrock, infiltration of mine water mixed with seepage from mill tailings ponds is causing local contamination of shallow, potable aquifers, but the problem is not considered serious and ongoing studies are underway. The State of New Mexico has installed a monitoring well network to determine temporal and spatial trends in groundwater quality. The U.S. Geological Survey, in particular, is monitoring surface flows and water quality in the Ambrosia Lake and Churchrock areas.

As part of the San Juan Basin Regional Uranium Study, the Department of Interior (DOI79) assisted by the U.S. Geological Survey (Ku79) examined selected water quality impacts from mining and milling and concluded that much of the mine effluent is suitable for irrigation, stock, and industrial use. Locally, it supports aquatic life and wildlife. Additional data on stream sediments are needed to evaluate the impact on water resources of erosion of waste rock from mines and mill tailings. It is preliminarily suggested (Ku79) that such erosion may be difficult to detect at distances of more than a few miles from the source because of the large amount of (natural) regional soil erosion. The results of the study are presently in draft form and may therefore be revised.

As part of the present study on uranium mining wastes, two New Mexico areas containing inactive mines were surveyed in the Spring of 1979. Stable and radioactive trace elements were monitored in surface runoff from sub-ore

and overburden piles in Ambrosia Lake and in the nearby Poison Canyon area. EPA staff selected the basic study areas and assisted in sample collection. The bulk of the sampling, analytical, and interpretation phases was done by Battelle, Pacific Northwest Laboratories (Wo79). Appendix G contains the data and discussion. As in the case of the Wyoming study area, the sampling program emphasized stream sediment sampling, cores, and shallow (5 cm thick) grab samples at the land surface. Samples of the source material were also collected. Figures 3.5 and 3.6 show the location of the study areas and sampling stations in New Mexico. Samples of the source material were collected at one of the two New Mexico drainage systems investigated. One system in Poison Canyon, New Mexico, was adjacent to several small surface operations as well as an underground mine site. For this system, no single source could be defined for the runoff constituents.

The Poison Canyon mine drainage system is a dry creek bed. The course of this creek passes an abandoned underground mine site from which it can receive runoff water. It then passes through a dirt roadway and follows a course adjacent to some small open pit mines. After a distance of several hundred kilometers, it joins a second branch drainage that originates next to a waste pile from one of the open pit mines. Samples were collected along this waterway starting with a background sample upstream of the underground mine about 200 m from the road. The first downstream sample was collected about 130 m downstream from the roadway. This was upstream of the runoff source originating in the open pit mine. The remaining samples were collected along the drainage way (Fig. 3.5), below contamination sources from the open pit operations.

A second site, the San Mateo Mine and environs, is located in the southeast portion of the Ambrosia Lake mining district. Large mine waste piles, a heap leaching operation, and a mine drainage pond are prominent at the site, which drains northward to San Mateo Creek.

Soil composites were collected at the waste pile and heap leach pile. These represent the source term for possible contamination of the watershed. The drainage samples were collected following one channel down the waste pile face to the intersection with San Mateo Creek, which was followed for a distance of 500 to 600 m from the site. Additional samples were collected in the gullies leading from the heap leach area and one of the off-site gullies. The latter represents blank soil upstream of the drainage water. Sampling sites are noted in Fig. 3.6. No significant contamination from the under-

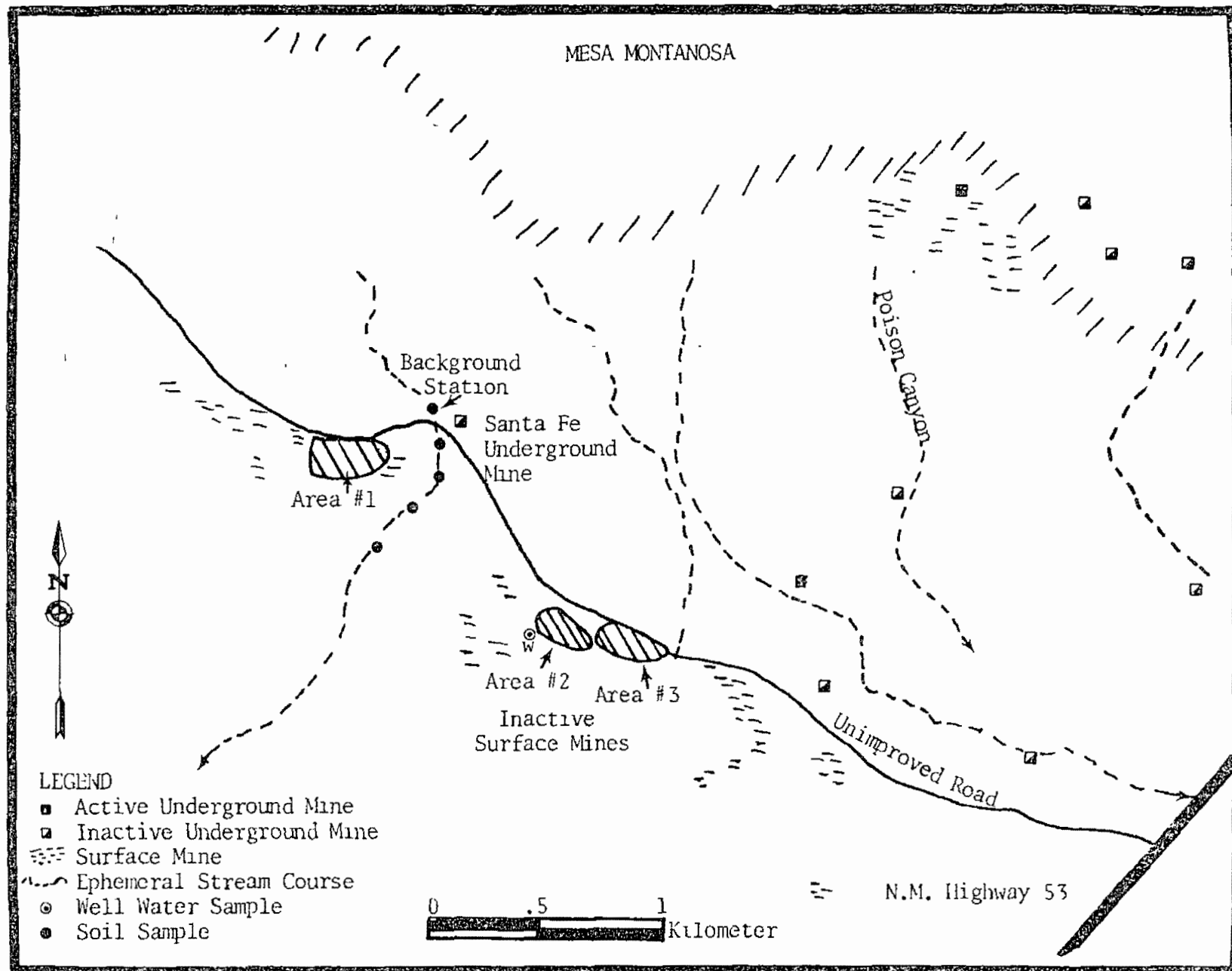


Figure 3.5 Location of study areas, sampling stations and uranium mines, Poison Canyon area, McKinley County, New Mexico

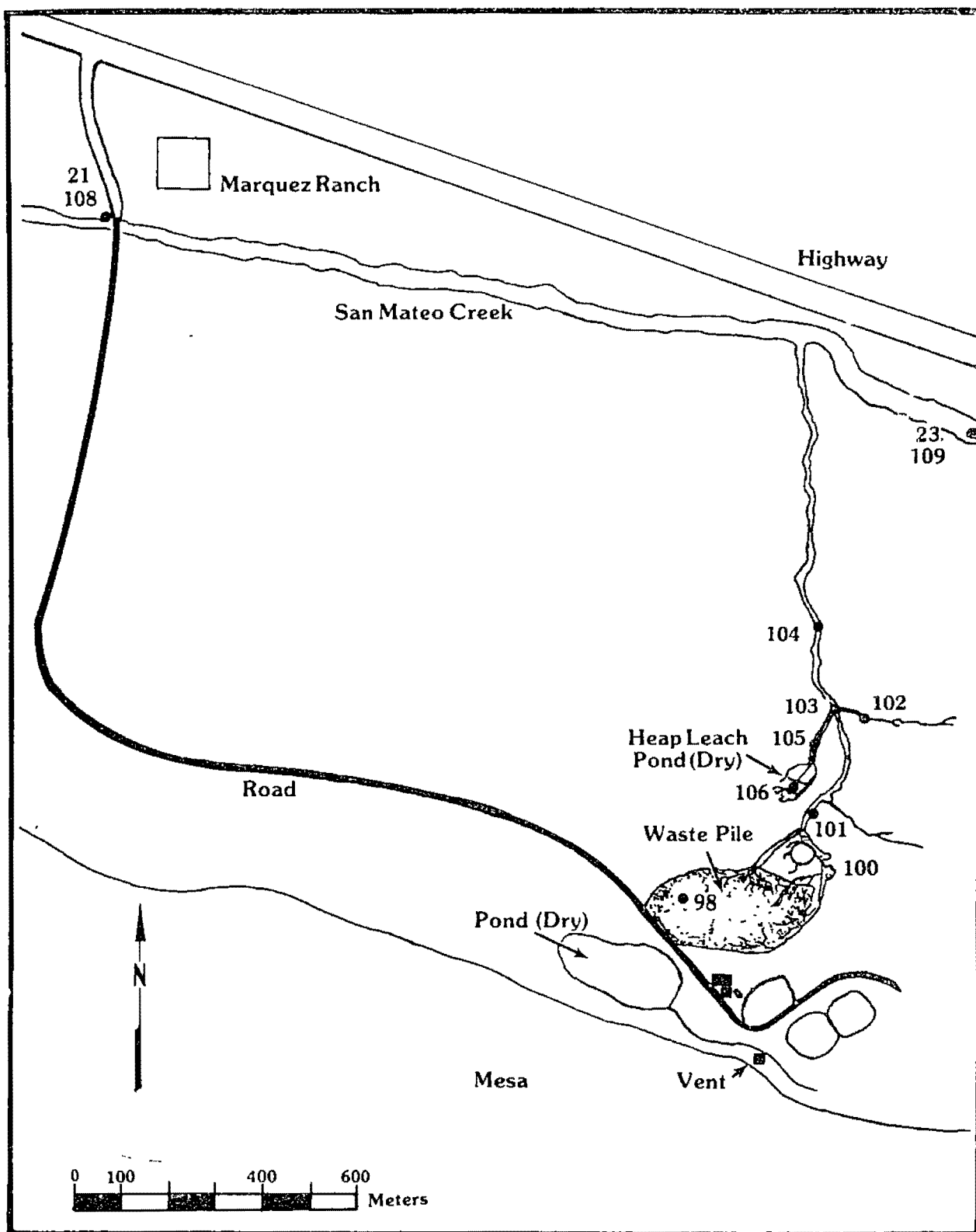


Figure 3.6 Sample locations for radionuclides and select trace metals in sediments, San Mateo mine, New Mexico.

ground site was detected. The Ra-226 content of the soil was about two times the background level at the furthest downstream site. This sample was collected about 130 m from the apparent source.

In summary, at the New Mexico inactive mine drainages the most prominent indicator of runoff from above-ground mineral storage is radium-226 in stream bed sediments. Concentrations in the source material are almost two orders of magnitude higher than those measured in the background soils. Elements such as uranium and selenium also have as large a concentration gradient, with concentrations decreasing downstream. At Poison Canyon, the radium-226 concentration diminished to two times background in a distance of approximately 100 m, while at the San Mateo site the distance was about 400 to 500 m. This may reflect either a more rapid transport by faster flowing water at the San Mateo site or, more likely, the larger source term there relative to background. At the San Mateo mine, radium-226 concentrations in water and sediments are significantly elevated downstream relative to upstream of the mine drainage.

3.2.3.3 Summary

The field studies conducted to date on the impacts of uranium mining on water quality are somewhat contradictory. Although no cases of gross, widespread contamination of groundwater or streams can be documented for uranium mining, there are cases of local contamination of water and sediments. From standpoints of theory and field data, there is need for cautious optimism in the use of local soil and water resources as sinks for waste discharge. Although numerous studies indicate that considerable reliance can be placed on the various physical and chemical processes to protect natural waters from contamination, investigations generally warn against using such studies to predict what may happen in other situations (Ru76; NRC79b; Ku79; Fu77; Am78). Laboratory results are highly dependent on the chemical properties of the fluid matrix and the physical and chemical properties of the particular soil studied. Results of field studies are site and time specific and have often suffered from inconsistent and undefined sampling and sample preservation techniques and questionable analytical measurements (Ku79; Ha78; Si77).

Our analyses reveal that there have been local water quality problems from mine water and wastes. Although widespread hazards have not been identified, this may be false security insofar as the present status of knowledge concerning trace element mobility in aqueous settings representative of

uranium mining areas is rather unclear from both theoretical and real-data standpoints. Most often, effects of mining are interspersed with and masked by impacts from uranium milling. This complicates or renders impossible any meaningful interpretation of the mining-related data. Despite the attempt to sort out some of the information on trace element mobility, there is insufficient understanding at this time to dismiss or otherwise reduce the significance of trace element contributions (from mining activities) to surface streams and, to a lesser extent, to groundwater.

We conclude that there is considerable information on the topics of trace element chemistry. It is also clear that trace element concentrations in natural fresh water are highly variable on both macro and micro geographic scales. There is great difficulty in correlating concentrations with such characteristics as streamflow or lithologic environment. Accurate prediction of the behavior and cycling of trace elements through water and sediments first involves characterization of physical states such as particle size and form (chelate, colloid, complex ion, precipitate, etc.), speciation, and availability to plants and animals (An73). Andelman concludes "...that there can be large differences in trace element concentrations [in water], on both a macro and micro geographic scale, and that such variations often occur in an unsystematic and nonpredictable fashion."

We recommend additional studies of spatial and temporal variations, sources and sinks of trace elements, chemical interactions within the hydro-geologic system, interactions between surface and groundwater systems, effects on aquatic biota, and effects on water use (human consumption, stock watering, irrigation). Periodic monitoring in certain areas would allow for the detection of the long-term trends of potential changes that would accompany anticipated increases in future mining activity -- during a period of increased competition for scarce water resources.

3.3 Surface Mining

3.3.1 Solid Wastes

Surface mining consists of removing materials, separating them into ore, sub-ore, and overburden, and storing them in separate piles on the surface near the mine for various periods of time (Section 1.3.2). The various storage piles are managed differently, vary in size and level of contaminants, and exist for varying periods of time. All are potential sources of

contamination to the environment via dusts suspended and transported by the wind, precipitation runoff, and Rn-222 emanation (Fig. 3.7).

3.3.1.1 Overburden Piles

Surface mining produces spoils at a rate of millions to tens-of-millions of tons per year. Unless this material is used to backfill the pit, large surface areas -- 40 hectares to over 400 hectares -- are covered to depths varying from a few meters to over 100 meters (Ka75, NRC77a, NRC77b, DOA78, Pe79).

Most of the mines begun since the early to mid 1970's use overburden to backfill mined-out areas of the pit (Ka75). Since older mines usually did not, erosion of their storage piles by water and wind may present an environmental problem (Ka75). In addition, the large amounts of overburden that past and present mines have used for road and dike construction and backfill also may present an environmental problem.

The annual average ore production of the 63 surface mines operating in the United States in 1978 was 1.2×10^5 MT (Section 1.3.1). Assuming an overburden to ore ratio of 50:1 (Section 1.3.2), the average annual production of overburden was about 6.0×10^6 MT per mine. A recent study of the eight large mines that accounted for 68 percent of the total 1977 United States U_3O_8 production from surface mines recommends the following average production parameters (Ni79):

1. ore production = 5.1×10^5 MT/yr
2. average ore grade = 0.11 percent U_3O_8
3. overburden:ore ratio = 77
4. overburden production = 4.0×10^7 MT/yr
5. mining days/yr = 330 d/yr

Surface areas of hypothetical overburden piles were computed using the above 63-mine and 8-mine overburden production rates and the following assumptions:

1. an average density of 2.0 MT/m^3 - reported values vary between 1.6 and 2.7 (Ro78, DOA78, NRC78a, Ni79)
2. the dumps are on level terrain
3. a rectangular waste dump with the length twice the width, and sides that slope at 45° angles (Fig. 3.8a)

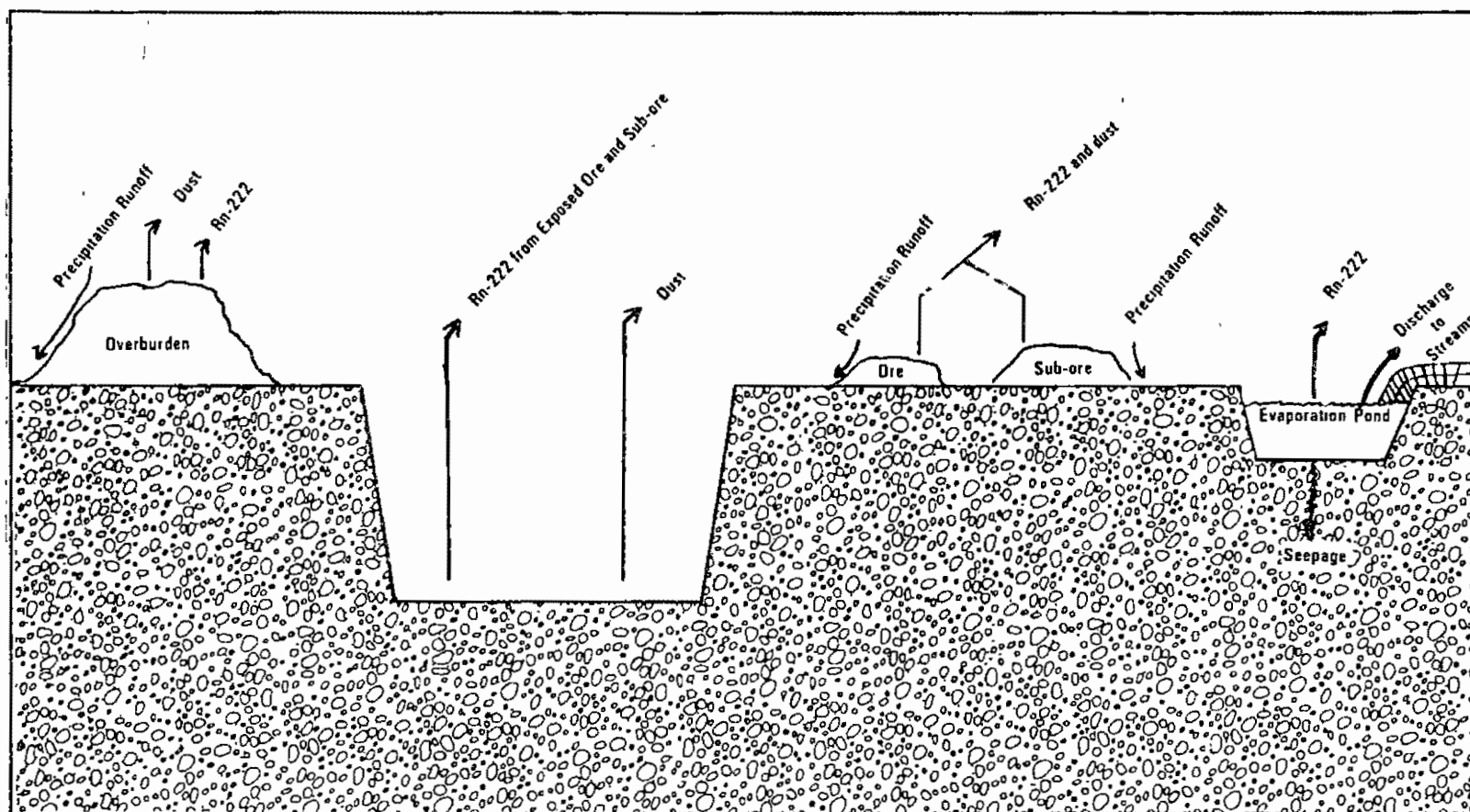


Figure 37 Potential sources of environmental contamination from active open pit uranium mines.

4. a waste dump in the shape of a truncated right-circular cone with 45° angled sides (Fig. 3.8b)
5. a bulking factor of 25% or 1.25 (Burris, E., Navajo Engineering and Construction Authority, Shiprock, NM, 2/80 personal communication)

Table 3.11 lists the surface areas of the hypothetical overburden piles in the following three cases:

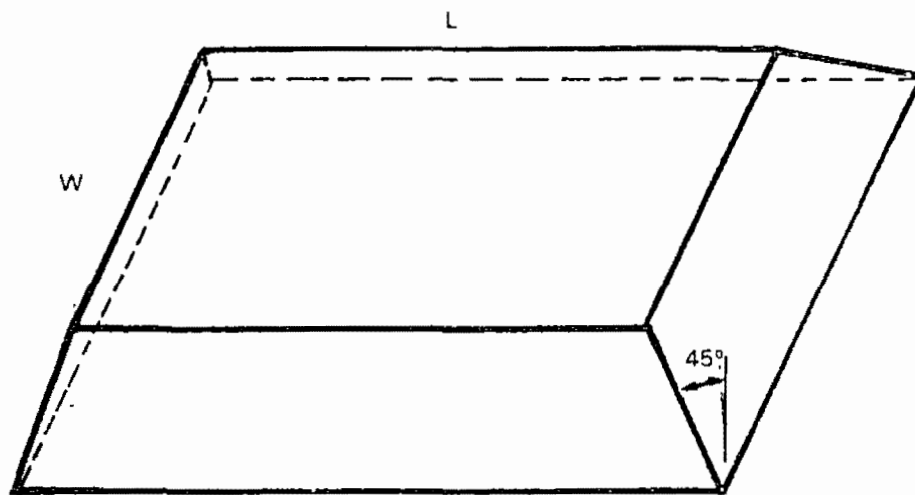
Case 1 - one year production with no backfilling

Case 2 - backfilling concurrent with mining - assumes 7 pits opened in a 17-year mine life with overburden from each successively mined pit used to backfill a previously completed pit, resulting in an equivalent of one pit of overburden (2.4-yr production) stored on the surface (Ni79)

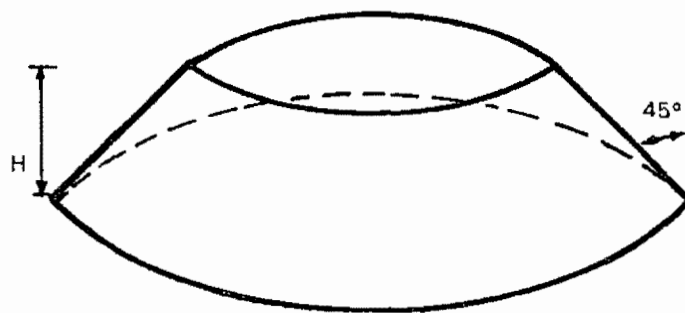
Case 3 - no backfilling during the 17-year mine life

The quantities of dust and Rn-222 that become airborne are directly proportional to the surface areas of waste piles. Table 3.11 shows the large variations possible between surface areas of waste piles at some active mines. Waste piles also cover various areas of terrain. However, for the same volumes, there are no significant differences in surface area or area of terrain covered for the two configurations of waste piles used in this study. Case 2 approximates recently activated mines, and Case 3 approximates older mines.

The type of rock in overburden spoil piles depends on the locations of the ore zones. Common rock types of New Mexico, Wyoming, and Texas mines include sandstone, claystone, siltstone, shale, and limestone, and unconsolidated silt, gravel, and sand (Co78, Pe79, Wy77, Ri78). In Texas, there are also lignite beds, tuffaceous silts, and some nearly pure volcanic ash



a) A rectangular pile with length twice the width and 45 degree sloping sides



b) A frustum of a regular cone with 45 degree sloping sides

Figure 3 8 Storage pile configurations assumed at surface and underground mines.

(Ka75). Coal veins are often present in Wyoming and New Mexico (Wy77, Ri78). However, the most abundant material in waste rock dumps will probably be clastic sedimentary rocks: sandstone, siltstone, and shale.

There is great variation in the particle size of material in waste piles, and this variation is important. Large particles ($>30\mu\text{m}$)*, because they usually settle within a few hundred feet of their origin, do not contribute to the airborne dust concentration (EPA77b). The potential for human respiration of the wind suspended dusts is also strongly influenced by the mean particle diameter (ICRP66).

Overburden rock is as large as available equipment can load and haul to the storage area. Rocks too large to handle with available equipment are broken into manageable sizes by small, explosive charges. Hence, rock particles will vary from less than a μm to a meter or more in diameter. Since weathering eventually breaks down the larger stones, the fraction of smaller particles increases over time.

Particle size distributions of material in waste rock piles at uranium mines have not been determined. It is likely that this material has a greater fraction of larger particles than that associated with crushed uranium mill tailings. Table 3.12 shows an example of the particle size distribution in the latter and the mean particle size distribution from a study of shale overburden removed from a surface mine in Pennsylvania (Ro78). Although the distribution fractions differ, a gross comparison can be made between the particle size of mill tailings and overburden waste. About 28 percent of the tailings were less than $50\mu\text{m}$ in diameter, and only about 12 percent of the particles in the overburden pile had similarly small diameters. Because only particles smaller than $30\mu\text{m}$ are likely to remain suspended by the wind for any significant distance (EPA77b), probably less than 10 percent of the overburden is a potential source of environmental contamination via wind erosion.

Table 3.13 shows the natural radionuclide concentrations in common rock types in the United States. In sedimentary rocks, which are common in the major uranium mining regions, the U-238 concentrations vary from less than 1 ppm** to about 4 ppm. Natural radioactivity usually is somewhat higher in the western states, and the uranium content in overburden prior to mining

* μm = micrometer = 10^{-6} meters.

**ppm = parts-per-million = 10^{-6} grams per gram of rock.

Table 3.11 Estimated surface areas associated with overburden piles

Management ^(a)	Pile Height, m	Overburden Volume ^(b) , m ³	Surface Area of Pile, m ²	Terrain Covered, Hectares
Rectangular Pile ^(c)				
		Average Large Mine ^(d)		
Case 1	65	2.5×10^7	5.2×10^5	48
Case 2	65	6.0×10^7	1.1×10^6	106
Case 2	30	6.0×10^7	2.2×10^6	209
Case 3	65	4.2×10^8	7.1×10^6	682
		Average Mine ^(e)		
Case 1	65	3.8×10^6	1.0×10^5	10
Case 2	65	9.0×10^6	2.2×10^5	20
Case 2	30	9.0×10^6	3.6×10^5	34
Case 3	65	6.4×10^7	1.2×10^6	113
Truncated Cone ^(f)				
		Average Large Mine ^(d)		
Case 1	65	2.5×10^7	5.2×10^5	46
Case 2	65	6.0×10^7	1.1×10^6	104
Case 2	30	6.0×10^7	2.1×10^6	208
Case 3	65	4.2×10^8	7.1×10^6	683
		Average Mine ^(e)		
Case 1	65	3.8×10^6	1.1×10^5	9
Case 2	65	9.0×10^6	2.2×10^5	18
Case 2	30	9.0×10^6	3.5×10^5	33
Case 3	65	6.4×10^7	1.2×10^6	110

Table 3.11 (continued)

(a) Management:

Case 1 - one year production with no backfilling

Case 2 - backfilling concurrent with mining - assumes 7 pits opened in a 17-year mine life and equivalent of one-pit overburden (2.4 year production) remains on surface

Case 3 - no backfilling during 17-year mine life

(b) Volume = production (MT/yr) x production years x bulking factor (1.25) ÷ by density (2.0 MT/m³).

(c) Length of pile is twice the width and the sides slope at a 45° angle (Fig. 3.8a).

(d) Overburden production = 4.0×10^7 MT/yr.

(e) Average 1978 overburden production of all 63 surface mines, assuming an overburden:ore ratio of 50/1, 6.0×10^6 MT/yr per mine.

(f) A frustum of a regular cone with 45° sloping sides (Fig. 3.8b).

is about 4 ppm (Ni79). However, during mining, some low-grade ore mixes with the overburden and may increase the concentration of the pile to as high as 20 ppm U₃O₈ (Ni79). This is equivalent to 12.6 disintegrations per minute (dpm) per gram of overburden. The progeny of the uranium will contribute additional radioactivity. Although there are local disequilibria between U-238 and its principal daughters, Th-230 and Ra-226, in ore-bearing rock, secular equilibrium will be assumed (Wo79). Small quantities of Th-232 and progeny will provide additional radioactivity. There is no apparent relationship between the Th-232 and U-238 decay chains. Th-232 concentrations in ores and host rock range from less than a pCi/g to a few pCi/g regardless of the U-238 concentration (Wo79).

Table 3.12 Particle size distributions of mill tailings
and mine overburden

Mill Tailings ^(a)			Overburden ^(b)	
Particle Size, μm	Weight Percent	Conc. ^(c) Avg. Conc.	Particle Size, μm	Weight Percent
250	60.3	0.15	>2000	75
125-250	7.5	0.03		
53-125	4.2	0.03	50-2000	13
44-53	3.8	0.03		
20-44	7.8	0.75	2-50	8
7-20	7.2	1.5		
1.4-7	9.1	4.6	< 2	4
< 1.4	0.0			

(a) Source: Sc79.

(b) Source: Ro78.

(c) The concentration of radionuclides in that fraction divided by the average concentration.

Table 3.13 Natural radionuclide concentrations in various
common rock types

Rock Type	U-238		Th-232		K-40	
	ppm	pCi/g	ppm	pCi/g	ppm	pCi/g
<u>Igneous</u>						
Basic	0.9	0.3	2.7	0.3	1.2	8.4
Granite	4.7	1.6	20	2.2	5.0	35
<u>Sedimentary</u>						
Shale	3.7	1.2	12	1.3	3.2	22
Sandstone	0.45	0.15	1.7	0.2	1.1	7.7
Limestone	2.2	0.7	1.7	0.2	0.32	2.2

Source: Oa72.

Table 3.14 shows the results of an extended airborne particle sampling program near a surface mine in New Mexico (Ea79). Although the on-site source of the radioactivity measured on these filters is undetermined, ore and sub-ore piles, waste rock piles, and mining activity all probably contribute. The higher activities reflect a greater contribution from ore dusts. From these air measurements, the above assumed average uranium concentration in overburden, 12.6 dpm/g (\approx 6 pCi/g), appears reasonable. These data also indicate that the progeny of U-238 through Ra-226 are in near secular equilibrium. The Th-232 concentration is about 1 pCi/g and, as indicated above, independent of the uranium concentration. Considering all available data, the radioactive source terms for overburden piles will be as follows: (1) U-238 and progeny = 6 pCi/g (0.0020 percent U_3O_8); (2) activity ratio (dust:overburden) = 2.5 (Section 3.3.1.2); and (3) Th-232 and progeny = 1 pCi/g. Figures 3.1 and 3.2 show the uranium and thorium decay series.

Table 3.14 Annual average airborne radionuclide concentrations in the vicinity of an open pit uranium mine, pCi/g

Location	U-238	Th-230	Ra-226	Th-232	U-238/Th-232
Jackpile Housing	76	80	70	1.2	63
Paguate	13	12	13	1.3	10
Bibo	9	7	5	1.3	7
Mesita	3	2	3	0.7	4
Old Laguna ^(a)	5	2	3	0.4	13

^(a) Background location.

Source: Ea79.

Little information is available on stable element concentrations in overburden rock. Table 3.15 summarizes the analyses of a few grab samples of soil and rock from a uranium mine in New Mexico and one in Wyoming (Wo79). Except for possibly Se, V, and As, there are no significant concentrations of stable elements attributable to uranium mining. Considering the typically high natural Se and V contents of many minerals common to these areas and the limited number of analyses, the inference of pollution is indefinite. A relationship between uranium and the stable element concentrations does not appear to exist. Thus, the stable element concentrations in overburden from the model surface mine will be the average concentrations of samples 6, 7, and 8 in Table 3.15. Table 3.16 lists the average concentrations.

3.3.1.2 Ore Stockpiles

Ore is often stockpiled at the mine as well as the mill. Although ore stockpiles are much smaller than the overburden waste piles, the concentrations of most radioactive contaminants are much greater in ore-bearing rock than in overburden. In addition, ore is stockpiled at the mine for shorter periods of time than waste rock. Ore stockpile residence times vary from mine to mine and range from a few days to a few months. The recent study of 8 large surface mines cited 41 days as an average ore stockpile residence time (Ni79). We will use this value to estimate the average area of ore stockpiles.

The average of the 63 operating surface mines produced 1.2×10^5 MT of ore during 1978. Assuming 330 working days per year and a 41-day ore stockpile residence time, a 1.5×10^4 MT ore stockpile would exist at the average mine. In comparison, the recent Battelle study reported that the average of eight large surface mines produced 1550 MT of ore per day, which would yield a 6.3×10^4 MT ore storage pile, assuming the same residence time (Ni79). The ore piles vary in height at different mines and different times. One study reports a maximum pile height of 9.2 m (30 ft) (Ni79), and at another site the maximum and equilibrium ore pile heights are estimated to be 6.7 m and 3.1 m, respectively (NRC78a). Using these parameters and a bulking factor of 1.25 (Burris, E., Navajo Engineering and Construction Authority, Shiprock, NM, 2/80, personal communication), the pile surface and pad areas were computed for the two production rates and two pile heights, 9.2 m and

Table 3.15 Uranium and stable element concentrations measured in rock and soil samples from two uranium mines

Sample	Concentration, $\mu\text{g/g}$														
	As	Ba	Cu	Cr	Fe ^(a)	Hg	K ^(a)	Mn	Mo	Pb	Se	Sr	V	Zn	U
<u>Wyoming</u>															
1. Top Soil Piles	3.2	700	13	46	1.3	<4	2.2	190	2.9	23	<1	89	60	37	6
2. Sub-ore	<1.8	6800	9	<36	1.2	10	2.3	140	<2.2	22	2.1	128	<100	25	61
3. Ore	5.4	800	9	<27	1.1	<7	2.3	180	<2.9	16	28	94	200	25	370
<u>New Mexico</u>															
4. Background Soil	4.1	450	12	<23	0.9	<4	1.8	200	5.5	12	<1	72	<50	22	<5
5. Background Soil	2.3	440	9	<20	0.8	<4	1.6	190	4.9	13	<1	50	<50	19	<5
6. Waste Pile	7.8	540	11	<28	0.8	<5	1.4	260	2.5	10	<1	99	<70	23	8
7. Waste Pile	14	280	21	<43	0.7	<8	0.5	750	<2.8	31	3.1	178	180	23	189
8. Sub-ore + waste	4.1	45	22	<51	0.3	<6	0.1	446	<1.8	25	<1.4	179	<55	13	57
9. Ore	6.0	64	27	<48	0.4	<6	0.2	673	<1.8	31	1.5	323	<55	14	---

Source: Wo79.

(a) Units are percent.

Table 3.16 Concentration of radionuclides (pCi/g) and stable elements ($\mu\text{g/g}$) in overburden rock from the model surface mines

Element	Concentration	Element	Concentration
Arsenic	9	Selenium	2
Barium	290	Strontium	150
Copper	18	Vanadium	100
Chromium	<51	Zinc	20
Iron ^(a)	0.6	U-238	6
Mercury	<8	Th-230	6
Potassium ^(a)	0.7	Ra-226	6
Manganese	485	Pb-210	6
Molybdenum	2.5	Po-210	6
Lead	22	Th-232	1

^(a) Units are percent.

3.1 m, assuming the same geometric configurations as for the overburden piles (Fig. 3.8). Table 3.17 gives the results. The computed surface areas of an average ore stockpile vary with volume of ore stored and pile height, but they are relatively independent of the pile shape.

Uranium deposits exist in sedimentary, metamorphic, and igneous formations. Sedimentary formations, primarily sandstone, siltstone, mudstone, and limestone generally host stratiform ore deposits often accompanied by carbonaceous material. Vein-type deposits usually occur in fractures of igneous and metamorphic formations. In the Rocky Mountain mining regions, about 98 percent of the recovered U_3O_8 comes from sandstone and related-type rock (St78). Sedimentary formations, principally sandstone, have been the predominant host for uranium in South Texas (Ka75).

Table 3.17 Estimated average areas of ore pile surface and pad

Pile Configuration ^(a)	Pile Height, m	Surface Area of Pile, m ²	Ore Pad Area, m ²
<u>Average Large Mine^(b)</u>			
Rectangular	9.2	6,300	5,700
Truncated Cone	9.2	6,200	5,300
Rectangular	3.1	14,000	13,500
Truncated Cone	3.1	13,700	13,200
<u>Average Mine^(c)</u>			
Rectangular	9.2	1,860	1,820
Truncated Cone	9.2	2,000	1,580
Rectangular	3.1	3,660	3,420
Truncated Cone	3.1	3,590	3,340

(a) See Figure 3.8.

(b) Volume of ore = 6.3×10^4 MT (41 day production) \times 1.25 (bulking factor)
 $+ 2.0 \text{ MT/m}^3 = 3.9 \times 10^4 \text{ m}^3$.

(c) Volume of ore = 1.5×10^4 MT (41 day production) \times 1.25 (bulking factor)
 $+ 2.0 \text{ MT/m}^3 = 9.4 \times 10^3 \text{ m}^3$.

The DOE does not expect the mineralogical characteristics of uranium ore to change appreciably in the future, since the known reserves are mainly in sandstone or a related host (DOE79). This fact is apparent from the data in Table 3.18, which gives the distribution of ore reserves in the United States by type of host rock. More than 97 percent of the uranium reserves are in sedimentary formations, primarily sandstone. Hence, it is reasonable to assume that ore stockpiles in the future will continue to consist mainly of a friable (easily crumbled) sandstone rock.

No data are presently available on the particle size distribution of material in ore stockpiles. Thus, the particle size distribution of ore will be assumed to be similar to that of overburden rock.

The average grade of ore mined in 1978 was about 0.14 percent U_3O_8 , but this will decline in future years (DOE79). The average grades of ore associated with the \$30 and \$50 reserves are 0.10 percent and 0.07 percent U_3O_8 , respectively (DOE79). Assuming the average grade of ore mined in the next decade to be about 0.10 percent U_3O_8 , the average uranium concentration in ore stockpiles will be 285 pCi/g (632 dpm/g). Although secular equilibrium in the uranium decay chain may not totally exist in some cases due to leaching by groundwater with subsequent redeposition, it appears reasonable to assume that radioactive equilibrium exists in a general assessment.

As discussed earlier, ambient Th-232 concentrations in the vicinity of a uranium mine range between 1 to 2 pCi/g. However, a concentration of 0.01 percent thorium is typical for ore from some surface mines (Mi76). This concentration is equivalent to 11 pCi Th-232/g of ore.

Uranium occurs in many ores as a secondary deposition. In a reducing environment, the soluble uranyl ion converts to insoluble uranium oxide and deposits preferentially on the smaller particles. (The total surface area of a given mass of smaller particles is greater than for larger particles.) Therefore, dusts that consist primarily of small particles have a greater specific concentration than ore as a whole (Table 3.12). The common procedure for computing uranium concentration in dust is to multiply the average concentration in the ore by 2.5 (NRC77a, NRC78a).

Table 3.18 Distribution of ore reserves by the type of host

Host Type	MT of Ore (10^6)	MT of U_3O_8	Percent Total Tons, U_3O_8
Sedimentary ^(a)	1,143.2	810,000	97.1
Lignite Materials	2.2	3,000	0.4
Limestone	1.3	1,200	0.1
Igneous and Metamorphic	<u>32.7</u>	<u>20,400</u>	<u>2.4</u>
Totals	1,179.4	834,600	100.0

^(a) Principally sandstone, but includes conglomerates, shale, mudstone, etc.

Note.--The reserves are \$50 or less per pound U_3O_8 , effective January 1, 1979 (DOE79).

Table 3.19 Average stable element concentrations in sandstone ores of New Mexico

Metal	Concentration, $\mu\text{g/g}$ ^(a)	Metal	Concentration, $\mu\text{g/g}$ ^(a)
Arsenic	86 (10-890)	Manganese	960 (70-3,000)
Barium	920 (150-1500)	Molybdenum	115 (3-700)
Cadmium	ND ^(b)	Nickel	20 (7-70)
Cobalt	16 (3-150)	Lead	78 (3-300)
Copper	61 (15-300)	Ruthenium	ND
Chromium	20 (7-70)	Selenium	110 (1-625)
Iron	15,700 (3,000-70,000)	Strontium	130 (1.5-300)
Mercury	ND	Vanadium	1410 (70-7,000)
Potassium	25,000 (7,000-30,000)	Zinc	29 (10-70)
Magnesium	3,500 (700-15,000)		

^(a) Range of concentrations given in parentheses.

^(b) ND - not detected

Note.--Ore samples are Dakota and Morrison sandstone from 25 uranium mines (Hi69).

In accord with the above discussion, we assume the following estimated average radionuclide source terms for ore stockpiles: (1) U-238 and progeny = 285 pCi/g ore (0.10 percent U_3O_8); (2) Activity ratio (dust:ore) = 2.5; and (3) Th-232 and progeny = 10 pCi/g ore.

Stable elements -- molybdenum, selenium, arsenic, manganese, vanadium, copper, zinc, and lead -- often associated with uranium ore at elevated concentrations may cause deleterious environmental and health effects. Mercury and cadmium are present only on rare occasions (Th78). However, as discussed above, there is no apparent relationship between concentration of stable elements and ore grade (Wo79). Table 3.19 lists measured (Hi69) concentrations of stable elements in 25 sandstone ores from New Mexico and average concentrations computed from these data. We assume the average concentration for the ore from the model surface mine.

3.3.1.3 Sub-ore Piles

All mines recover some rock containing uranium ore that at the time of mining is uneconomic to mill. The grade of this "sub-ore" varies with the "cutoff" level assigned by the mill. Some mines process sub-ore by heap leaching, which changes the chemical properties and constituents of the pile (Section 1.3.5.1). However, most mines store the sub-ore in separate piles and recover it when it becomes economically feasible.

The sizes of sub-ore dump piles vary with the quantity of ore mined and its grade. One study suggests that the sub-ore accumulation rate equals the ore production rate (Ni79), a ratio similar to that reported for the Sweetwater uranium mining operation (NRC77a). Using this assumption with the ore production rates given above for the average large mine and average mine, 5.1×10^5 MT/yr and 1.2×10^5 MT/yr, respectively, the average sizes of sub-ore piles generated at a constant rate during the 17-year active life of a mine were based on an 8.5 year accumulation and a bulking factor of 1.25. Figure 3.8 shows the shapes of the piles assumed, and Table 3.20 gives the results for piles 30 m high. The surface areas of the two pile configurations differ very little.

The mineralogical characteristics of ore and sub-ore are very similar. Thus, the distribution in Table 3.18 will apply to sub-ore. This study considers the particle size distribution of sub-ore the same as for overburden and ore.

In the early mining years, the ore cutoff grade was usually about 0.15 percent U_3O_8 . However, this has continually decreased until today the cutoff ore grade is about 0.03 percent U_3O_8 (Ni79, NRC77a). Hence the ore content of these piles will be less than 0.03 percent U_3O_8 , and the average content has been estimated to be one-half the cutoff grade, or 0.015 percent U_3O_8 (Ni79), which is equivalent to 43 pCi U-238/g (95 dpm/g). Also, the uranium in the sub-ore, as in ore, is assumed to be in secular equilibrium with its progeny. Because the occurrence of uranium in sub-ore is the same as in ore and the mineralogies are similar, the uranium in sub-ore should be concentrated on small particles by the same factor as in ore, 2.5.

The Th-232 concentration in sub-ore is between the ambient level and that in the associated ore, 1 pCi/g to 11 pCi/g. For lack of measured Th-232 concentrations, we assume that less than 2 pCi/g of Th-232 will be present (Table 3.14). The radiological significance of an error in this assumption will be small.

From the above discussion, we assume the following estimated average radionuclide source terms for sub-ore piles: U-238 and progeny = 40 pCi/g (0.015 percent U_3O_8); activity ratio (dust:sub-ore) = 2.5; and Th-232 and progeny = 2 pCi/g. Figures 3.1 and 3.2 show the uranium and thorium progeny.

Table 3.20 Estimated average surface areas of sub-ore piles during the 17-year active mining period

Pile Configuration ^(a)	Surface Area of Pile, m ²	Terrain Covered, Hectares
Average Large Mine ^(b)		
Rectangular	1.2×10^5	11
Truncated Cone	1.2×10^5	11
Average Mine ^(c)		
Rectangular	3.5×10^4	3.2
Truncated Cone	3.6×10^4	3.0

(a) See Fig. 3.8.

(b) Volume of sub-ore = $8.5 \text{ yr} \times 5.1 \times 10^5 \text{ MT/yr} \times 1.25 \div 2.0 \text{ MT/m}^3 = 2.7 \times 10^6 \text{ m}^3$.

(c) Volume of sub-ore = $8.5 \text{ yr} \times 1.2 \times 10^5 \text{ MT/yr} \times 1.25 \div 2.0 \text{ MT/m}^3 = 6.4 \times 10^5 \text{ m}^3$.

Stable elements observed in ore will also be present in sub-ore. Because stable element concentrations specific to sub-ore are unavailable and are unrelated to ore grade, concentrations in the sub-ore from the model surface mine will be assumed equal to those in the ore (Table 3.19).

3.3.1.4 Reclamation of Overburden Piles

Reclamation is usually done only for overburden piles. Ore stockpiles are continually being disturbed and their residence time is short. Also, sub-ore piles generally are not stabilized in anticipation of recovering the uranium at a later time. Hence, only overburden and waste rock piles are considered for stabilization and reclamation. Section 1.3.2 gives a brief description of these practices.

Backfilling mined out areas of the pit is necessary for an adequate reclamation program. Because of the swelling of earthen material once mined, sufficient material should be available to completely fill the pit when mining is completed. However, even though backfilling is generally being performed at most recently active mine sites, sufficient overburden is often not replaced to eliminate the pit.

Improperly stabilized spoil piles may become sources of contaminants to the environment. The wind can suspend and transport small-sized particles containing elevated levels of contaminants. Radon-222, produced by the radioactive decay of Ra-226 contained in the rocks, can emanate from the pile surfaces. Precipitation runoff from the piles can carry particulate matter and dissolved contaminants into the natural surface drainage system if rainfall exceeds the infiltration and holding capacity of the pile. The general procedure for reducing wind and water erosion is to grade the piles to conform to the natural terrain, cover the area with a layer of topsoil, and seed it with a native grass.

These spoils consist of unweathered and unconsolidated rock, coarse gravels, and sands and allied materials isolated from the natural processes that occur on surface soils. Consequently, spoils have poor textural properties and low water-holding capacities. Having no established flora to aerate the surface and make nutrients available, spoils are barren of nutrients required for plant growth. Hence, to sustain vegetation on these piles may be difficult because of poor soil quality and the arid conditions

in the principal mining regions. Therefore, all plant growth depends on the topsoil cover, which is generally less than 30 cm thick (Re76). This is often inadequate to store sufficient water and nutrients to sustain plant growth during extended dry periods. Soil irrigation and fertilization may be required for several years until plants can sustain themselves.

Proper grading of the spoil piles, with water management and conservation, can help reclamation. The piles should have less than a 3:1 slope to reduce surface water runoff and erosion (St78). Forming catchment basins and terraces to hold water on the spoils and reduce water erosion will also increase the amount of runoff available to the plants. It also has been determined that vegetation on north-facing slopes requires about half the applied water of that on south-facing slopes (Re76). Water requirements of vegetation on horizontal surfaces and east and west slopes are about intermediate between those of the north and south slopes. Hence, spoil piles with long, north slopes will conserve water and reduce the irrigation required. Locating piles on leeward slopes and away from natural drainage will also reduce wind and water erosion.

The reestablishment of native grasses and shrubs is essential for controlling wind and water erosion and providing wildlife habitat. Wyoming requires a pre-mining vegetation inventory for use in evaluating post-mining reclamation (Wy76). Similar statutes governing mine reclamation are in effect in other states (Section 1.4). The Soil Conservation Service has recommended seed mixtures that are best suited to climatic and soil conditions in different areas of the West (St78). Newly seeded areas are usually protected from grazing by fencing for at least two growing seasons to allow the plants to become established.

Abandoned pits fill with water and form small lakes that livestock and wildlife can use for drinking water, if the water is uncontaminated. But, unless properly managed, final pits may be hazards to people and wildlife. Therefore, steep walls should be graded to give safe access into the pit, and after grading, the pit banks should be seeded to minimize erosion and prevent the sides from sloughing off.

3.3.2 Mine Water Discharge

3.3.2.1 Data Sources

The principal sources of information used to model the mining region in

Wyoming are the site-specific EIS's and ER's for active and proposed mining/milling operations and the NPDES permit data on discharge volume and quality. Several reports by state and federal agencies supplemented the foregoing, particularly with respect to estimating ambient water quality and flood volumes for various return periods and annual or monthly flows in principal streams of the region. Foremost among these is work by the State (Ha78), the U.S. Geological Survey (Cr78, Ho73), and the Soil Conservation Service (DOA75).

Self-monitoring data collected by industry and reported to EPA were also checked to ascertain compliance with NPDES permit conditions. Unfortunately, the permits do not specify limits on the volume of discharge; hence, the total mass or flux per unit of time may or may not agree with the values originally estimated by the discharges in the EIS, ER, or license application.

3.3.2.2 Quantity and Quality of Discharge

The purpose of this section is to identify water quality associated with surface uranium mining in the Wyoming Basin. This area was selected for detailed source term characterization and pathways analysis because of past and ongoing uranium production, primarily by surface mining. A subsequent section (3.4.2) similarly addresses underground mining. The analysis to follow is incomplete and preliminary, owing to the limited existing data, the lack of opportunity for significant new investigations in the time of this study, and the decision to pursue the objectives on a "model area/model mine" approach. So many variables of ore occurrence, mining practices, climate, geology, and hydrology exist that a detailed investigation is unrealistic.

Table 3.21 summarizes water quality data for seven surface and three underground mines in Wyoming. Uranium averages 0.62 mg/l and ranges from 0.02 to 1.3 mg/l. Dissolved radium-226 is typically less than 4 pCi/l, although one mine reportedly discharged 10.66 pCi/l. Suspended solids average 24.9 mg/l. There is considerable variation from one facility to another; the observed range is 2.7 to 87.2 mg/l. Zinc is the only stable element consistently monitored, probably because the NPDES permit addresses it. Concentrations average 0.04 mg/l and are well below the 0.5 mg/l limit in the permits. Barium and arsenic are less frequently monitored but appear to be in the range of 0.05 mg/l for barium to 0.005 mg/l for arsenic. Both of these values are well below the discharge limits.

Table 3.21 Summary of average discharge and water quality data for uranium mines in Wyoming and a comparison with NPDES limits

Project	Mine Type	Discharge m ³ /min	Radioactivity		TSS	Major and trace constituents, mg/ℓ					
			Total U mg/ℓ	Ra-226 pCi/ℓ		SO ₄	Zn	Fe	Ba	Cd	As
1	U	0.85	0.95	3.92	87.2		0.08	1.25			
2	U	6.57	0.41	2.28	2.7	234	0.02	0.02	0.05		
3	U	0.70	0.02	7.41	8.8		0.01				
4	S	1.89	1.30	10.66	5.0		0.01				
5	S	3.60	0.63	3.94	11.1		0.14				
6	S	5.68	0.02	2.85	10		0.05				
7	S	3.52	0.98	0.67	19.4	875	0.05			0.004	0.005
8	S	1.21	0.14	3.03	17.3		0.02				
9	S	0.10	1.14	3.6	62.5		0.16				
10	S	4.55									
<u>All Mine Types (1 through 10):</u>											
Average:		2.87	0.62	4.26	24.9	555	0.06	0.64	0.05	0.004	0.005
Standard Deviation:		2.25	0.50	3.00	29.5	453	0.06	0.87	-	-	-
<u>Underground Mines (1 through 3):</u>											
Average		2.71	0.46	4.54	32.9	234	0.04	0.64	0.05		
Standard Deviation:		3.35	0.47	2.62	47.1	-	0.04	0.87	-		
<u>Surface Mines (4 through 10):</u>											
Average		2.94	0.70	4.1	20.88	875	0.071			0.004	0.005
Standard Deviation:		1.96	0.53	3.4	21.04	-	0.063			-	-
Summary of NPDES Permit Limits											
Daily Average/Daily Maximum			2/4	3/10 ^(a) 10/30 Total Radium	20/30		0.5/10	-/2	-/1	0.05/0.1	0.5/1

(a) Total Ra-226 limit is not monitored.

Source: NPDES permits from Region VIII (R. Walline, written communication), site-specific reports (EIS, ER), and self-monitoring data.

Mean values from six surface mining projects in Wyoming were the basis for estimating the effects of mine discharge on water quality. Values from mines in the South Powder River Basin model area compare very well with the Wyoming mines, thus supporting adoption of a model mine in the Basin. There were no strong differences in water quality between surface and underground mines. Table 3.21 shows that discharge is highly variable, ranging from 0.1 to 6.57 m³/min, with an average of 2.87 m³/min. In surface mining projects, the average is 2.94 m³/min, with a standard deviation of 1.96, indicating considerable discharge variation among facilities. This study assumes an overall average flow of 3 m³/min from each surface mine in the calculations of chemical loading of local and regional streams (see Section 3.3.3 and Appendix H).

Table 3.22 shows water quality and flow rates associated with open pit mines in other areas and in various stages of operation. Ongoing development of an open pit mine in Colorado involves 28 m³/min discharge and is therefore well above the average. Radium, uranium, and suspended solids are relatively low. Producing open pit mines in New Mexico are usually dry or nearly so and are dewatered at rates of 0.6 m³/min or less. The water is used for dust control. Radium concentrations can be very high (New Mexico Projects) due to long residence time of groundwater in the ore body and the concentrating effects of evaporation. Similarly, groundwater associated with ore bodies in Texas and Wyoming may contain several hundred picocuries per liter.

Mine dewatering has the greatest potential for adverse environmental and public health impacts. Although contaminant concentrations in the effluent conform to NPDES requirements, there is long-term contaminant loading to the ambient environment. Contaminants concentrate on stream sediments because of sorption and evaporation and become available for transport by flood water. Regional or at least local dewatering of ore bodies may deplete high quality groundwater. Theoretically, dewatering may induce horizontal or vertical influx of poorer quality groundwater into productive or potentially productive aquifers, but the extent of this phenomenon is poorly documented. We strongly recommend further study because the work done to date is largely oriented toward determining engineering feasibility versus the overall environmental impact.

Table 3.22 Water quality associated with surface and underground mines in various stages of construction and operation

Project	Discharge m ³ /min	Total U mg/l	Dissolved Ra-226 pCi/ l	Pb-210 pCi/ l	Milligrams per liter				
					TSS	SO ₄	As	Mo	Se
<u>Colorado</u>									
Open pit mine:									
Development stage	28	1.044	4.10		16.2				
<u>New Mexico</u>									
Producing open pit mine, seepage to pit	0.13	2.5	180	17	168	2151	0.005	0.018	0.019
Open pit mine, ponded inflow water	0.58	2.6	220	26	23	842	0.005	0.545	0.043
<u>Texas</u>									
Active open pit mine holding pond.			50 to 100			380	< 0.01	< 0.01	< 0.01

Overland flow is not dismissed herein as a significant pathway, although its impact is of lesser importance according to data from April 1979 field studies in New Mexico and Wyoming (see Section 3.2.3.2 and Appendix G). A recent U.S. Geological Survey study for the Bureau of Indian Affairs (Ku79) addresses projected effects of runoff over long time periods if wastes and sub-ore are not stabilized or covered. The study concludes, with essentially no real data, that stream flows are too small in the sub-basin to transport wastes. In the larger basins, such as the Rio Puerco, sediment loads are so great that addition of tailings and, presumably, mine wastes would be insignificant. It is our opinion that additional field study is needed. Overland flow in a long time period could move radionuclides in the wastes into the main stream channels. Since this source will be available for many years after mine closure, if wastes are not stabilized, it may become a major one.

Seepage of contaminated water from mine holding ponds, which are operated to reduce suspended solids concentrations in mine discharge water, is believed to be insignificant. Since the ponds have relatively small areas, their seepage losses are small compared to losses by infiltration of releases to the watercourses. In some mining areas, such as the Powder River Basin, shallow groundwater quality is naturally poor. Maximum attenuation of contaminants is expected in the shallow, poorly permeable bedrock strata of the Wasatch and Fort Union Formations.

3.3.3 Hydraulic and Water Quality Effects of Surface Mine Discharge

3.3.3.1 Runoff and Flooding in the Model Surface Mine Area

3.3.3.1.1 Study Approach

Precipitation and runoff estimation for the model surface mine scenario in Wyoming considers three hydrographic units: sub-basin, basin, and regional basin. Respective surface areas are 11.4, 5,400, and 13,650 square kilometers (km^2). The mine is located in the sub-basin. The sub-basin, the basin, and the regional basin are all drained by ephemeral streams. The latter is drained by a major regional river that has wide seasonal variations in flow and is dry or nearly so about 180 days each year. The sub-basin has similar flow variability. Figure 3.9 depicts the mine in relation to the sub-basin, basin, and regional basin.

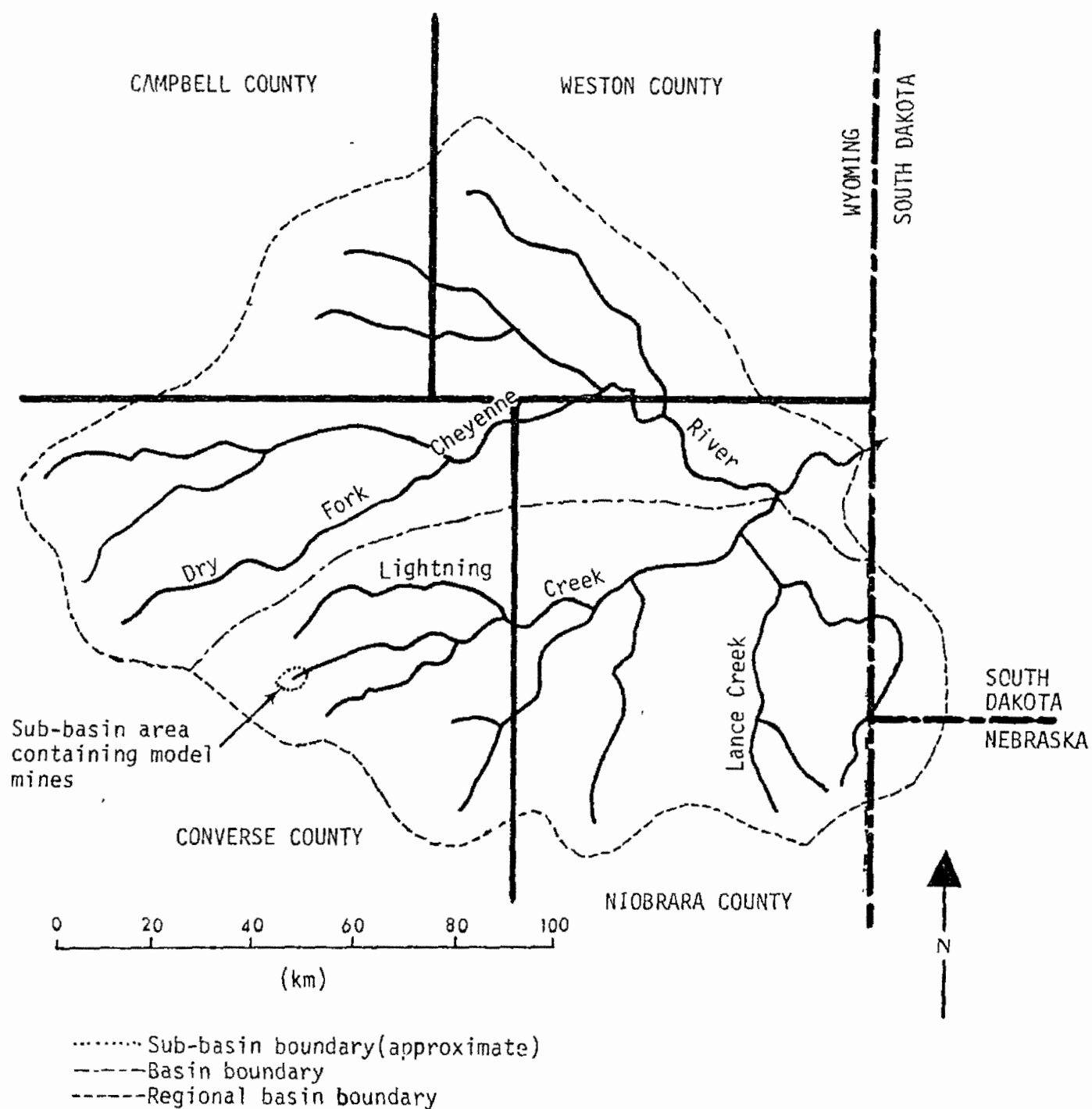


Figure 3 9 Sketch of sub-basin, basin, and regional basin showing orientation of principal drainage courses, areas of drainage, and location of mines

The first general approach defined quality and volume of mine water discharge. Hypothetical hydrographic basins were then delineated and flood flows calculated for return periods ranging from 2 to 100 years. The individual and collective effects of discharge from three mines were then evaluated in terms of perennial flow, flood flow, and chemical transport. Of key importance was an estimation of the extent of perennial streams created by mine discharge and the influence of contaminants on water quality in the river draining the regional basin.

3.3.3.1.2 Description of Area

We selected an area of active mining and milling in the South Powder River Basin of Wyoming for analysis. The area has four active or imminently active uranium mills and a number of open pit mines. Available data on the geology, hydrology, and water quality of the area are sparse, but because of the mining and milling activity are relatively well known for a remote region like northeastern Wyoming. The study team chose one mining and milling project in the area for field investigation in April 1979; hence, additional data became available and are used herein as appropriate.

Terrain in the area has low rolling hills and an average elevation of 1414 m (MSL datum). Since the climate is not very different from that of nearby Casper, Wyoming, meteorological data from that station are fairly representative of the region. There are no relatively large seasonal and annual variations in precipitation intensity, frequency, and duration. Mean annual precipitation over a 30-year record period is 28.5 cm and occurs mainly as scattered thunderstorms in late spring and early summer. These thunderstorms supply 25 to 50 percent of the total annual precipitation and are usually of high intensity, short duration, and can be quite local. Potential pan evaporation averages 110 cm per year and greatly exceeds precipitation.

Streams in the study area are ephemeral and only exhibit measurable surface flow during snowmelt and heavy thunderstorm activity. Average total monthly flow for the period 1948 through 1970 for Lance Creek and the Cheyenne River at Spencer, Wyoming reveal distinct high- and low-flow periods in the year (Fig. 3.10). We believe that the streams represent the basin and regional basin hydrographic units used herein. Large watersheds usually exhibit measurable surface flow for about 180 days per year. Small watersheds, 30 to 40 square kilometers, may not flow at all for several consec-

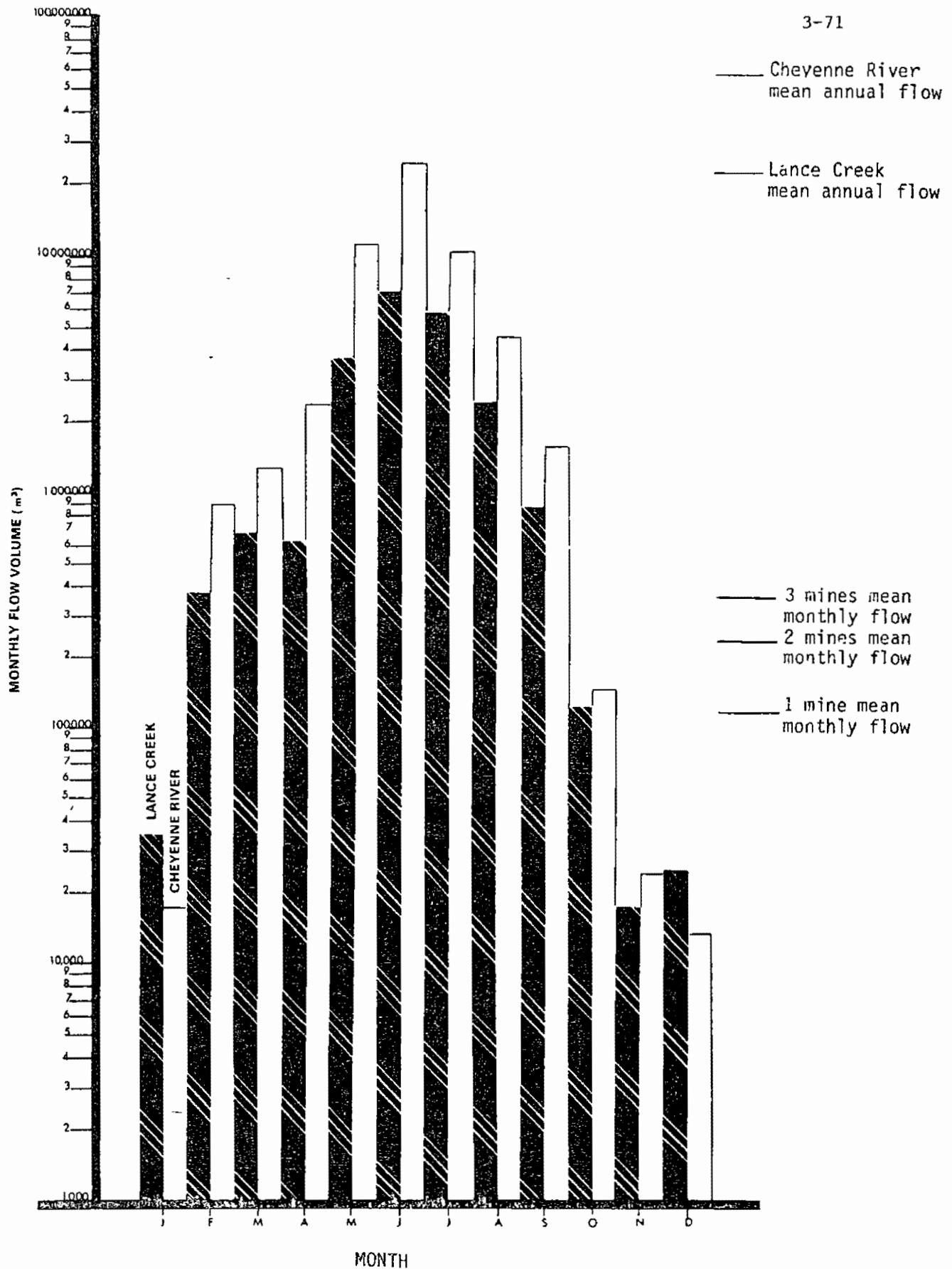


Figure 3 10 Average monthly flows for the Cheyenne River and Lance Creek near Spencer, Wyoming, for the period 1948-1970 (DOI59, DOI64, DOI69, DOI73)

utive years. Mean annual runoff is 0.8 to 1.3 cm or 0.0023 to $0.004 \text{ m}^3/\text{sec}$ per km^2 .

Peak flows in the regional basin and basin area are a result of snowmelt in at least 50 percent of the cases. This is commonly due to temporary but rapid melting from January to March. High flows can also result from widespread summer storms, but these are the exception. For small basins on the order of forty square kilometers or less, peak flows occur because of thunderstorms in the summer months. Thus peak flows in small basins versus the basin or regional basin commonly occur for different reasons and at different times in the year. A period of peak runoff from the sub-basin might coincide with a low flow or zero discharge condition in the basin or regional basin.

In the area of the Morton Ranch project (DOA75) there are 14 sub-basins, the average of which is about 11.4 km^2 . Channel slopes are 11.4 to 31.4 m/km (average 21.2 m/km), and basin slopes are about 88 m/km . These are tributary to larger streams with channel slopes of 2.17 to 17.0 m/km (average 6.63 m/km), and which drain basins with an area of $5,400 \text{ km}^2$ and a mean annual flow of $0.80 \text{ m}^3/\text{sec}$. These, in turn, are tributary to a regional basin with an area of $13,650 \text{ km}^2$ and mean annual flow of $1.47 \text{ m}^3/\text{sec}$. All three hydrographic units are drained by ephemeral streams. The main stem of the regional system is dry an average of 180 days per year. The basin drains into the regional basin, assumed here to be the Cheyenne River Basin, which drains an area of $13,650 \text{ km}^2$ (Da75, Lo76, Ra77).

Surface water in the model area is used mainly for stock watering and irrigation. The amount of irrigated area in the basin is 1400 hectares, compared to 2800 in the regional basin. Because of extreme variability in surface flow volume and water quality, almost all municipal water comes from wells completed in bedrock. Stock water is from both wells and impoundments, whereas single-family domestic supplies are primarily from wells.

3.3.3.1.3 Method of Study

Because of dilution considerations, flow volume rather than peak discharge rate is of prime concern. For the basin and regional basin areas, only peak flow rate can be readily estimated on a probability basis for annual and longer time periods of perhaps 2, 5, 10, etc. years. Peak flow in the larger hydrographic areas commonly does not coincide with that in the

smaller basins. Also, there is poor correlation between peak flow rate (Q) and total flow volume (V) for streams draining large basins. Total flow volume in the larger basins can be estimated from partial duration flow data. That is, we can estimate the percentage of the time, during the year, flow will be of a given magnitude.

Relationships among runoff volume, rainfall, and surface area in small basins (encompassing less than 30 square kilometers) in the Powder River Basin have been developed by the U.S. Geological Survey (Cr78) and the Soil Conservation Service (DOA75). Peak discharge and total annual flow in the basin and regional basin units were measured by the U.S. Geological Survey for Lance Creek at Spencer, Wyoming and for the Cheyenne River near Spencer.

We analyzed the effects of perennial or chronic mine discharge on changing existing ephemeral streams in the sub-basin, basin, and regional basin to perennial streams using a crude seepage and evaporation model. The basic equations and approach, explained in Appendix H, are similar to those used in the Generic Environmental Impact Statement on Uranium Milling (NRC79b). Adjustments were made for mine discharge rates and infiltration and evaporation losses. The main output of the model is an estimate of which stream segments might become perennial and what the net discharge would be from a number of mines operating in the same sub-basin. Water quality impacts can only be very roughly assessed. For the time being, we assume that infiltration and evaporation decrease flow but do not effect the chemical mass in the system. That is, we assume contaminants in mine drainage are deposited on or in the stream/wash substrate and remain available for transport by flood water.

The sub-basin is as shown in Fig. H.1 (Appendix H) and contains three active uranium mines, each of which discharges $4,320 \text{ m}^3/\text{day}$. Quaternary alluvium constituting the channel is assumed to have a porosity of 40 percent. The sub-basin contains seven streams or wash segments, three receiving mine water directly. Water from the mines dissipates by infiltration, evaporation, and as surface flow that may leave the sub-basin entirely. Appendix H shows the basic equations and assumptions and gives a complete summary of "losses" due to seepage and evaporation as well as any net outflow from the sub-basin.

Precipitation-runoff in the Wyoming study area correlates rather closely to basin size. Basins of about $10,000 \text{ km}^2$ area have an annual unit-area runoff of 0.43 cm/yr ; whereas an area of perhaps 25 km^2 might have a runoff of only 5 cm/yr . Decreased runoff (on a unit area basis) associates with larger basins and reflects water storage, channel losses, and evapotranspiration that occur mainly in the tributaries. Impoundments are rarely on the main stem of streams, where washouts are a problem, but rather on tributaries. The average impoundment is located about every 130 square kilometers, is rather small, and is used for stock water. Very infrequently, small flood-irrigation projects may use impounded water for grasslands. Seventy-five percent of the annual runoff occurs during the summer thunderstorm activity in May, June, and July. Snowmelt occurs rather slowly and is captured in the headwater areas, whereas rainfall events are rather intense and localized, causing excess flows that reach the main stem, Lance Creek and Cheyenne River. Sediment loads are high in both the tributary and main stem streams.

Contaminant concentrations in overland and channel flow during peak runoff events in the sub-basin are expected to follow the pattern shown in Fig. 3.11, the data for which are from the U.S. Geological Survey (H. Lowham, in preparation) for a small basin, Salt Wells Creek, in the Green River Basin of southwestern Wyoming.

Note in the inset of Fig. 3.11 that the washoff peak, that portion of the runoff enriched in dissolved and suspended materials, precedes the runoff peak. Runoff in small basins is typically associated with brief but intense thunderstorms that flush the land surface. Total suspended solids (TSS) concentrations are disproportionately high in the peak flow events. Discharges of $170 \text{ m}^3/\text{min}$ carry $100,000 \text{ mg/l}$ TSS; whereas flows of 1 cfs might carry only 500 mg/l . The leading edge of the high flow has the greatest concentration of suspended solids and dissolved chemical load. Figure 3.12 depicts discharge and specific conductance values as a function of time for the same small basin in Wyoming. Specific conductance (SC) is a rough measure of the total dissolved solids (DS) content, following the approximate relationship: $\text{DS} = 0.71 \text{ SC}$. Note that the first rise in the flow hydrograph occurs about three hours after the peak for specific conductance, indicating the presence of a contaminated "front" laden with salts and other suspended and soluble materials. The second peak on the flow hydrograph similarly precedes and is associated with degraded water quality due to this flushing

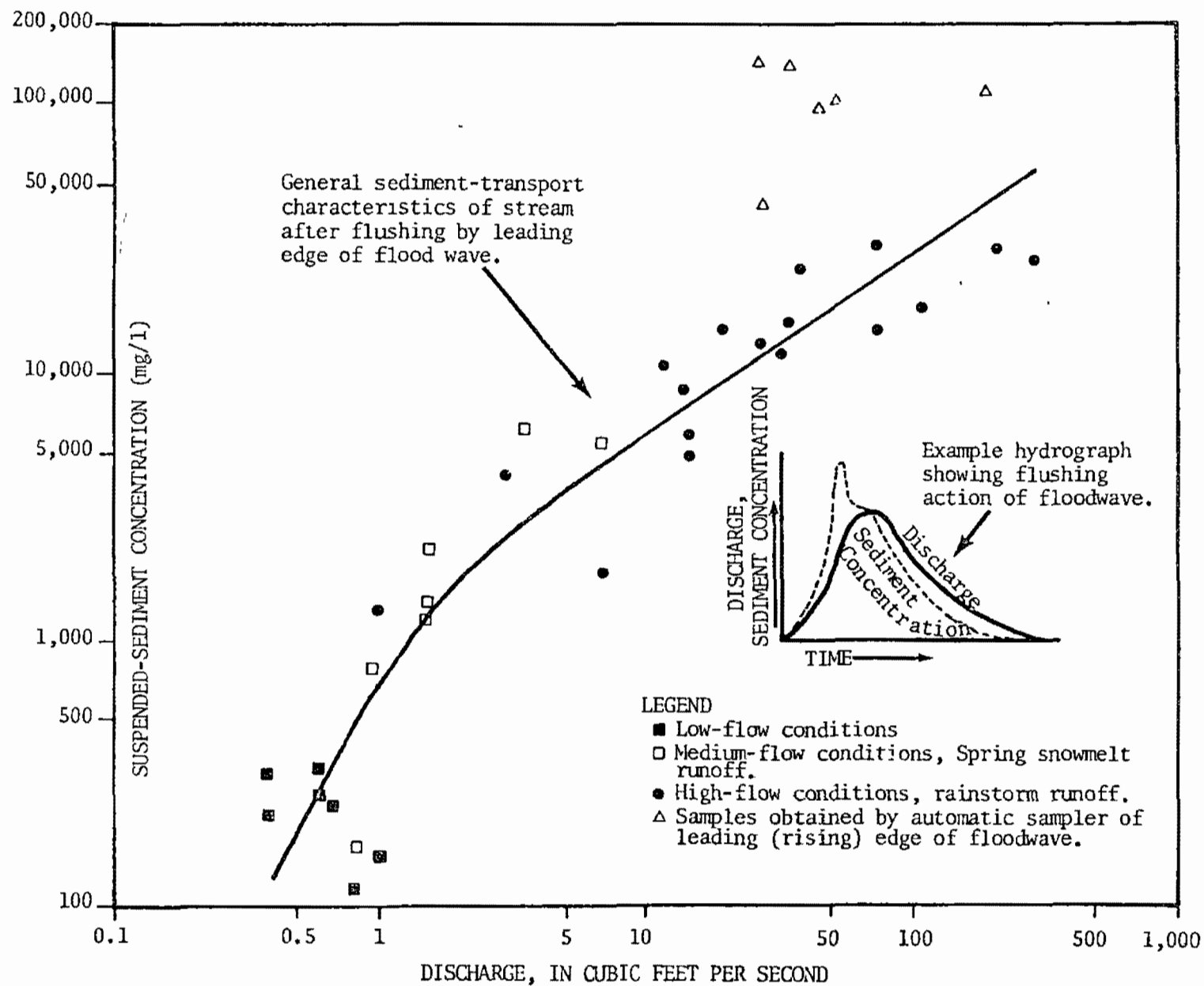


Figure 3 11 Suspended sediment concentration to discharge. Salt Wells Creek and tributaries, Wyoming (From U S Geological Survey data, H Lowham, in preparation)

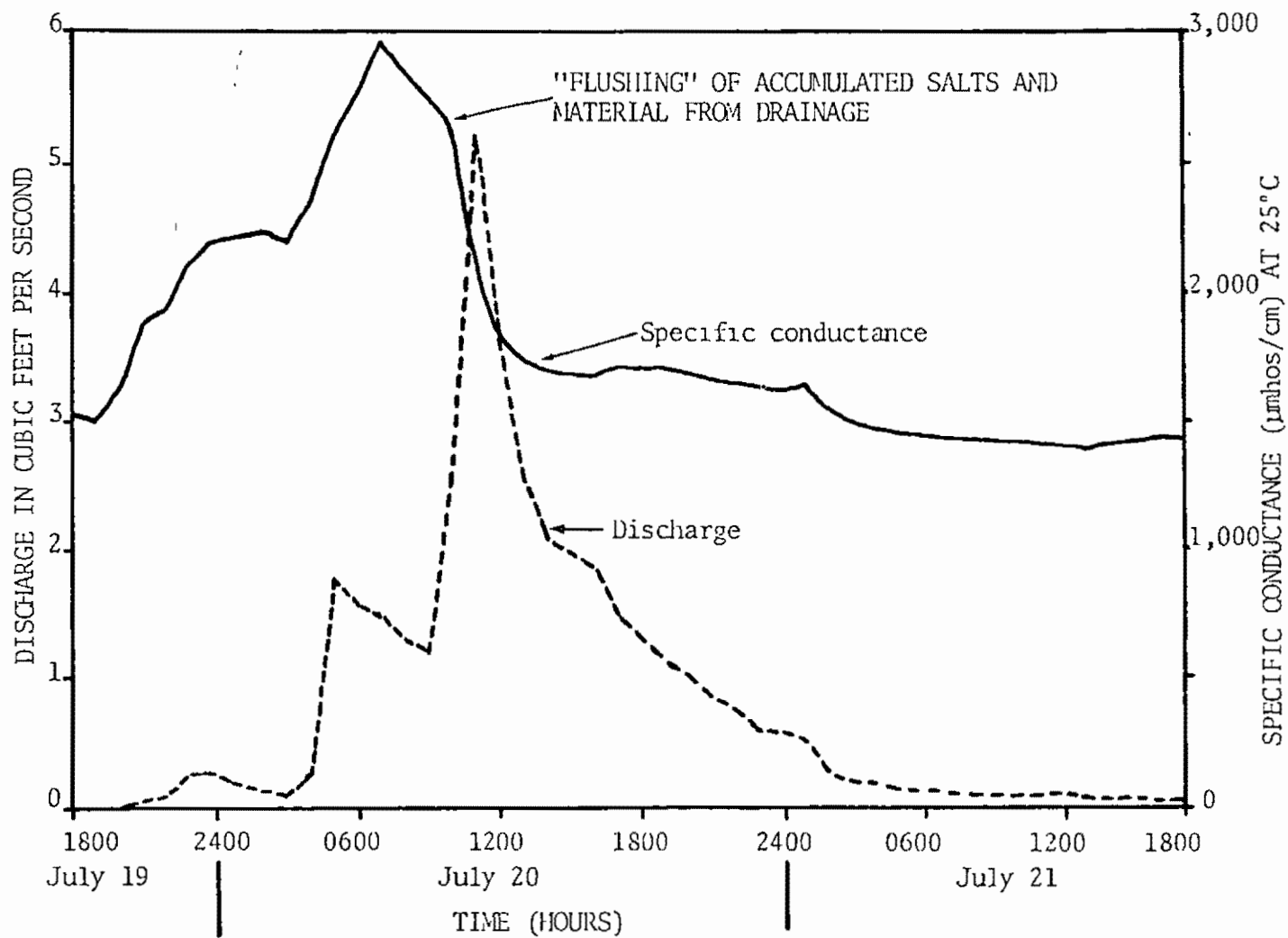


Figure 3.12 Relation of discharge and specific conductance to time at Salt Wells creek, Green River Basin, Wyoming (From U S Geological Survey data, H Lowham, in preparation)

action. About 33 hours after precipitation begins, runoff water quality and flow very nearly approximate antecedent conditions. This indicates rather thorough flushing, most of which occurred in an 18-hour period.

Assuming similarity between the surface mining area and the situation described above, we believe intense flows of rather short duration flush most of the contaminants from the land surface and stream channels. Although sub-basin floods are expressed in terms of return period for Wyoming and in terms of partial duration (1-day, 7-days) and varying return periods for New Mexico, we believe the basic approaches (total flow vs. partial duration) to be rather similar because of the "flashy" nature of runoff in both study areas. In the New Mexico case, the mean discharge rate and the flow volume for the 7-day event are very often less than that for the 1-day event for the same return period. This also confirms the intense, short-term nature of runoff processes in the Wyoming and New Mexico model areas.

3.3.3.1.4 Discussion of Results

This section addresses the interaction between mine drainage and flood waters. Flood magnitude is addressed first, followed by calculation of water quality effects due to mine water. The U.S. Geological Survey technique (Cr78) for estimating floods in small basins in northeastern Wyoming was used to estimate peak discharge and total flow volume in the sub-basin. Multiple regression analysis reveals that the variables of area, slope, and relief provide roughly 90 percent correlation between rainfall and runoff (Cr78). Considering the numerous assumptions made throughout the analysis, only the area variable is used herein. It accounts for 70 percent of the flow. Table 3.23 shows the peak discharge rate and total flow volume from the sub-basin for floods with recurrence intervals (r) of 2 to 100 years. The basic equation for calculating discharge rate or flow volume is--

$$Q_r \text{ or } V_r = a A^{b_1} \quad (3.1)$$

where a = regression constant

b_1 = drainage area coefficient for area: peak discharge area: volume relationships

A = basin area

= 11.4 km²

Q_r, V_r = discharge rate and flow volume for flooding events with return periods of 2, 5, 10, 25, 50, 100 years.

The flow volume for the two-year flood is $32,921 \text{ m}^3$, and the instantaneous peak flow rate is $387 \text{ m}^3/\text{min}$. For comparison, we assume that the model mine discharges $3.00 \text{ m}^3/\text{min}$ or $1.6 \times 10^6 \text{ m}^3$ per year. Assuming the annual flood volume is $30,000 \text{ m}^3$, it is apparent that annual dilution is essentially nil and can be expected to be zero for perhaps 8 to 10 months of the year when there is no natural runoff. As Table 3.24 shows, flow volumes calculated using the Soil Conservation Service (DOA75) methodology average 1.5 times greater than those derived using the USGS approach. The latter are used herein in the interests of conservatism, i.e., there is less volume for dilution of contaminants.

Table 3.23 Peak discharge and total volume for floods of 2, 5, 10, 25, 50 and 100 year recurrence intervals

Recurrence Interval, r, in years	Regression		Volume (m^3)	Regression		Peak Discharge ($\text{m}^3/\text{min.}$)
	Constant, a	b_1		Constant, a	b_1	
2	9.62	0.689	32921	96.21	0.582	387
5	18.08	0.713	64116	199.6	0.612	840
10	24.87	0.727	90009	292.8	0.632	747
25	34.71	0.739	127862	441.1	0.660	1269
50	42.82	0.748	159920	575.4	0.679	2674
100	51.58	0.756	194937	731.1	0.699	3500

Peak flood discharges in the basin and regional basin range from 4,053 to $31,401 \text{ m}^3/\text{min}$ for recurrence intervals of 25 years or less. For 50- and 100- year flooding events, peak discharge approximates 19,370 to $44,860 \text{ m}^3/\text{min}$. In the regional basin, discharge is $1.7 \text{ m}^3/\text{min}$ or less approximately eight months of the year and equals or exceeds $17 \text{ m}^3/\text{min}$ for about three months of the year, typically in the winter and early spring.

Maximum discharge from the basin and sub-basin is expected in the late spring and early summer months because of thunderstorms. At this time, flow in the river draining the regional basin is also at or near maximum, thus there is high probability for considerable dilution of runoff contaminated by mine drainage.

Total flow volumes for the basin and regional basin were estimated from U.S. Geological Survey records for the period 1948 to 1970. Figure 3.10 shows average monthly flows in cubic meters for the Cheyenne River and Lance Creek near Spencer, Wyoming. Immediately apparent is the close similarity in overall runoff pattern for the year.

Table 3.24 Summary of calculated total flow in the Wyoming model area sub-basin using the USGS and SCS methods

Recurrence Interval, r, in years	Sub-basin Total flow (m^3) ^(a)	Sub-basin Total flow (m^3) ^(b)
2	32,921	14,467
5	64,116	NC ^(c)
10	90,009	98,419
25	127,862	170,815
50	159,920	231,618
100	194,937	295,257

^(a)Source: Cr78.

^(b)Source: DOA75.

^(c)NC = Not calculated.

Minimum flows occur in November, December, and January, and peak runoff in both basins occurs in May, June, and July. Long-term average annual flow in the basin is $2.18 \times 10^7 m^3$ and $5.64 \times 10^7 m^3$ in the Cheyenne River. These are almost exactly proportional to the respective basin areas of $5,360 km^2$ and $13,650 km^2$, indicating similar climatic and runoff conditions.

Assuming there are 3 mines operating for a 17-year period and that each mine discharges on the average $3.00 \text{ m}^3/\text{min}$ continuously, total annual flow volume from the mines is $4.7 \times 10^6 \text{ m}^3$. Cumulative discharge from the sub-basin is $7.04 \text{ m}^3/\text{min}$ or $3.7 \times 10^6 \text{ m}^3/\text{yr}$, which causes development of a perennial stream 12.8 km long within the basin. Insofar as the basin channel length is 141 km, the perennial stream ceases to flow well within the basin.

Appendix H explains the methodology and intermediate steps involved in deriving these foregoing values. Mine drainage water is not expected to flow the full length of Lance Creek or reach the Cheyenne River. However, on the basis of total monthly flow, the volume of mine drainage from one mine exceeds the flow in Lance Creek and the Cheyenne River for three months of the year, whereas flow from three mines exceeds basin flow for five months and regional basin flow for four months each year (Fig. 3.10).

The aqueous pathway for mine drainage is considered in terms of chronic, perennial transport in the mine water, per se, and transport by flood waters that periodically scour the channels where most of the sorbed contaminants would be located. Considering the random nature of flooding and the resulting uncertainty as to when the next 2-, 5-, or 10-year, etc. flood may occur, it is assumed that most contaminants accumulate on an annual basis and are redissolved by floods of varying return periods (2 to 10 years) and volumes. Many combinations of buildup and flooding are possible, such as buildup for 5 years or 10 years with perhaps several 2-year storms and one 5-year storm. Insofar as numerous assumptions are made in calculating volume and quality of mine discharge, basin runoff, and fate of the contaminants in the aqueous system, use of annual accretion and varying flood volumes in the sub-basin is considered adequate for estimating flood water quality.

Dilution of contaminated flows originating in the sub-basin and extending into the basin were conservatively calculated by assuming that the total flow during the low period equaled the mean annual flow. Thus, high flows and associated increased dilution are ignored, tending to make the analysis conservative. Contaminated flows from the sub-basin are diluted into these adjusted mean annual flows. Definition of the source term on an annual basis is most compatible with the radiation dose and health effects calculations in Section 6. Use of the low flow segment of the total annual

flow regime is decidedly conservative since total flow during the five months of low flow conditions amounts to $111,610 \text{ m}^3$ and $218,336 \text{ m}^3$ for the basin and regional basin, respectively. Average annual flow for the period of record (22 years) is considerably higher, amounting to $2.184 \times 10^7 \text{ m}^3$ for the basin and $5.64 \times 10^7 \text{ m}^3$ for the regional basin.

Runoff in the basin and regional basin is expected to markedly dilute contaminated flood flows originating in the basin. Such floods would scour contaminants from about 23 kilometers of channel affected by contaminants from the three active mines. Peak runoff events in the sub-basin are most likely in the late spring-early summer season when runoff in the basin and regional basin is the maximum or near maximum, on the average. However, peak runoff from the sub-basin could also occur when the basin and regional basin are at low flow or zero discharge. Such contrasts are present between the basin and regional basin flow regimes. From September through December, Lance Creek can be expected to have no discharge from 45 to 65 percent of the time, whereas the Cheyenne River will be dry, on the average, from 65 to 85 percent of the time (Fig. 3.13). Thus there is a distinct chance that contaminants transported in Lance Creek would not be immediately diluted upon reaching the Cheyenne River.

Before discussing the calculated concentrations of contaminants in the basin and regional basin streams, several other conditions need to be mentioned. In water-short regions like Wyoming, extensive use is made of impoundments to capture and store runoff. On Lance Creek, the model for the basin, the volume of existing impoundments is $15.78 \times 10^6 \text{ m}^3$ or 72 percent of the annual average runoff. In the regional basin, modeled after the Cheyenne River, there are $4.2 \times 10^7 \text{ m}^3$ of storage volume, which is 74 percent of the average flow of $56.4 \times 10^6 \text{ m}^3$. Thus, it is very likely that discharge from the sub-basin or basin will not exit the basin, particularly in the periods of low flow. Contaminant concentrations, particularly those affected by sorption and precipitation reactions, are likely to be reduced as a result of sedimentation and long residence time in the impoundments, although there is some potential for overtopping, disturbance by cattle, and so on. Significant adverse impacts are not likely considering precipitation and sorption reactions which are likely to remove contaminants from the food chain. Proof of this is lacking and we recommend confirmatory studies for the stable elements. Previous studies (Ha78; Wh76) emphasized radiological contaminants.

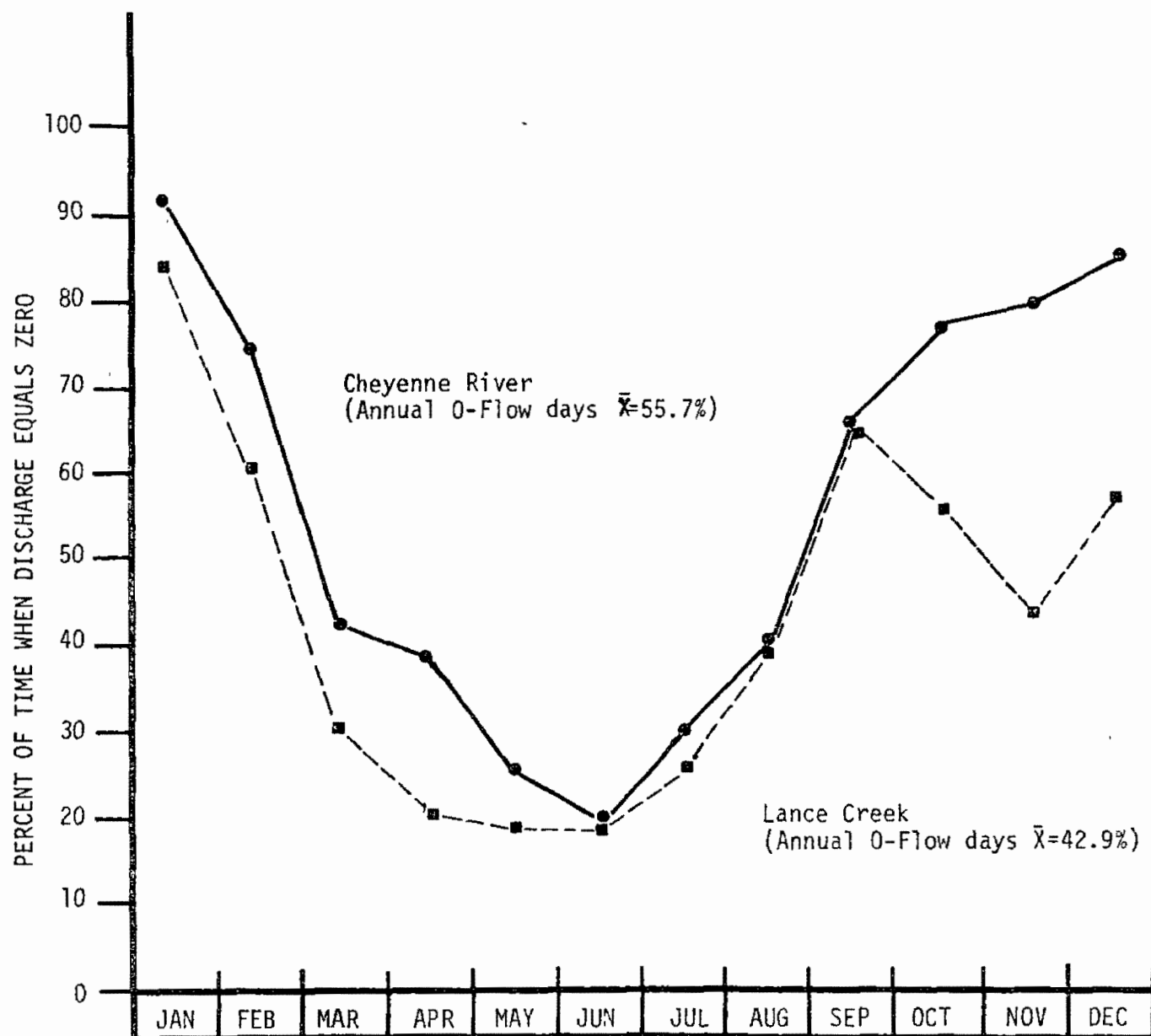


Figure 3.13 Periods of no flow in Lance Creek and the Cheyenne River near Riverton, Wyoming for the period 1948-1978
(Summarized from flow records provided by H. Lowham, U.S. Geological Survey, Cheyenne, WY)

Radium-226 is strongly sorbed onto stream sediments and (or) it is subject to precipitation. Partial re-solution in subsequent floods occurs but it is assumed that only 10 percent of the mass deposited on an annual basis goes back into solution in flood waters. The rationale for this assumption is based on laboratory studies (Sh64; Ha68), field data from New Mexico (Ka75; Ku79), and review of the literature. Pertinent field and laboratory data specific to surface water quality in the Wyoming uranium mining areas are scarce, although studies by the State (summarized by Harp, 1978) are noteworthy. Sulfate is regarded herein as rather mobile and, as such, most of it infiltrates the shallow aquifer. Therefore, only 20 percent of the mass from a given mine on an annual basis is assumed available for re-solution in flood waters. The fate of zinc, arsenic, and cadmium is insufficiently understood to predict what fraction in the mine discharge will be removed from solution versus remain available for re-solution. Studies along these lines are necessary. Similarly, not all of the contaminants potentially present in mine waters from Wyoming are necessarily shown in Tables 3.21 and 3.25, which were developed based on available data from NPDES permits, environmental reports, and environmental impact statements. In the case of suspended solids, there is no calculation of non-point source contributions from mined lands. Sediment loads from such sources could be locally significant, but mined land reclamation and natural recovery seems to effectively mitigate problems. Only suspended solids from mine drainage, per se, are considered.

Table 3.25 shows the flood flow volumes (in the sub-basin) associated with events having return periods of 2, 5, 10, 25, 50, and 100 years. Also shown are the contaminant concentrations calculated from the annual contaminant loading diluted into the foregoing floods. As expected, concentrations are high because of the low dilution volumes associated with the small sub-basin. Surface water in the sub-basin might be impounded therein for use by stock or, less possibly, irrigation, but it is more likely that the principal impoundments would be in the larger hydrographic unit, the basin. The flood flow volumes shown represent runoff from the entire sub-basin. When the second and third mines begin to discharge, the annual loading and concentration values shown would have to be doubled or tripled. The reader should remember that background concentrations already present in flood runoff would be additive to the values in Table 3.25. However, these have been assumed

Table 3.25 Annual contaminant loading from one uranium mine and resulting concentrations in floods within the sub-basin for return periods of 2 to 100 years

Contaminant and concentration in mine effluent	Chemical mass available for transport on an annual basis	Flood flow volumes (m^3) and contaminant concentrations associated with return periods of 2 to 100 years ^(c)					
		$V_2 = 32921$	$V_5 = 64116$	$V_{10} = 90009$	$V_{25} = 127862$	$V_{50} = 159920$	$V_{100} = 194937$
		C_2	C_5	C_{10}	C_{25}	C_{50}	C_{100}
Total uranium	0.070 mg/l 110 kg/yr	3.34	1.72	1.22	0.86	0.69	0.56
Radium-226	4.1 pCi/l 0.00065 Ci/yr ^(a)	19.7	10.1	7.2	5.1	4.1	3.3
Total suspended solids	20.9 mg/l 32,955 kg/yr	1001	514	366	258	206	169
Sulfate	875 mg/l 275,940 kg/yr ^(b)	8381	4304	3066	2158	1723	1416
Zinc	0.071 mg/l 112.0 kg/yr	3.40	1.75	1.24	0.876	0.700	0.575
Cadmium	0.004 mg/l 6.31 kg/yr	0.192	0.098	0.070	0.049	0.039	0.032
Arsenic	< 0.005 mg/l 7.88 kg/yr	0.239	0.123	0.088	0.062	0.049	0.040

(a) Ten percent of the annual loading is assumed available for solution. The balance is assumed sorbed onto sediments or present in insoluble precipitates.

(b) Twenty percent of the annual loading is assumed available for transport and the balance is assumed to have infiltrated to the water table or it is present as an insoluble precipitate.

(c) V_r and C_r refer to, respectively, flood volume, in cubic meters, and concentration in milligrams per liter or picocuries per liter for an r -year flood. Concentrations are in milligrams per liter except radium-226, in pCi/l.

Note.--Assumptions: Mine discharges continuously at a rate of $3.00 m^3/min$ and concentrations are the average of those shown in Table 3.21. All suspended and dissolved contaminants remain in or on the stream sediments and are mobilized by flood flow.

equal to zero in order to estimate incremental increases due to mining and to simplify the calculations.

Table 3.26 shows contaminant concentrations in the basin and regional basin streams from the discharge of one mine. For cases involving two or more mines, the concentration shown would be scaled up by a factor of two or more. Basically, the table shows the effects of taking contaminated flood waters from the sub-basin and diluting them in the low flow volume of the basin and regional basin. As expected, concentrations decrease with floods of greater volume and longer return period. Additional dilution occurs when discharge from the basin enters the regional basin. Taking the two-year runoff event in the sub-basin, for example, uranium is diluted from 3.34 mg/l (Table 3.25) to 0.76 mg/l in the basin and then to 0.44 mg/l in the regional basin. There is some question as to whether the lesser sub-basin floods, particularly those with return periods of 25 years or less, would actually flow the length of the basin and enter the regional basin. Because much of the 22.7 km reach of stream directly affected by mine discharge is located in the basin, it is conservatively assumed that the contaminants will reach the basin and eventually the regional basin. The foregoing analysis is structured as a worst-case, maximum-concentration scenario.

Concentrations of contaminants in flood waters affected by mine drainage are compared to water standards for potable and irrigation uses (Table 3.27). Radium-226 concentrations in the basin and regional basin streams (Table 3.27) range from 1.6 to 4.5 pCi/l and are below the drinking water standard (for Ra-226 + Ra-228) of 5 pCi/l. Uranium concentrations range from 0.26 to 0.76 mg/l, which is roughly equivalent to 176 to 514 pCi/l. On the basis of chemical toxicity alone, such concentrations would probably present no problem for short periods, but radioactivity is another matter. Reevaluation of the standard for uranium in potable water is presently receiving attention within the Agency (R. Sullivan and J. Giedt, USEPA, oral communication, 1980). Briefly, there is consensus that the radiotoxicity of uranium is similar to that of radium-226 and 228. For continuous ingestion at a rate of 2 liters per day, it is suggested that potable water contain no more than 10 pCi/l (0.015 mg/l) natural uranium to reduce the incidence of fatal cancers to no more than 0.7 to 3 per year per million population (Office of Drinking Water guidance to the State of Colorado, July 7, 1979). Realizing that the

Table 3.26 Concentrations in basin and regional basin streams as a result of surface mine discharge

Parameter	Concentrations (mg/l ; pCi/l in the case of radium) in basin discharge under low flow conditions due to influx of sub-basin floods with 2, 25, and 100 year return periods ^(a)			Concentrations (mg/l ; pCi/l in the case of radium) in regional basin discharge under low-flow conditions due to influx of basin discharge, also under low-flow conditions, and sub-basin floods with 2, 25, and 100 year return periods ^(b)		
	C ₂	C ₂₅	C ₁₀₀	C ₂	C ₂₅	C ₁₀₀
Total Uranium	0.76	0.46	0.36	0.44	0.32	0.26
Radium-226	4.5	2.7	2.1	2.6	1.9	1.6
Total Susp. Solids	228	138	107	131	95	79
Sulfate	1909	1152	900	1098	797	668
Zinc	0.774	0.468	0.366	0.445	0.324	0.271
Cadmium	0.044	0.026	0.020	0.025	0.018	0.015
Arsenic	0.054	0.033	0.025	0.031	0.023	0.019

^(a) Calculated as follows: Assuming a two year flood, uranium concentration in the outflow from the sub-basin equals 3.34 mg/l and flow equals 32,921 m³ (see Table 3.25). Average total flow for 5 months of low flow conditions in the basin equals 111,610 m³. The concentration in the basin outflow, after dilution of the contaminated inflow from the sub-basin for floods of varying recurrence intervals equals:

$$C_{\text{Basin}} = \frac{V_{\text{Sub-basin}} \times C_{\text{Sub-basin}}}{(V_{\text{Sub-basin}} + V_{\text{Basin}})} = \frac{(32921 \text{ m}^3) (3.34 \text{ mg/l})}{32921 \text{ m}^3 + 111610 \text{ m}^3} = 0.76 \text{ mg/l}$$

^(b) Calculations similar to "a" above, except average total flow volume for 5 months of low flow in the regional basin equals 218,336 m³. Hence,

$$C_{\text{Regional basin}} = \frac{V_{\text{Sub-basin}} \times C_{\text{Sub-basin}}}{(V_{\text{Sub-basin}} + V_{\text{Regional Basin}})}$$

Table 3.27 Comparison of potable and irrigation water standards and surface water quality affected by surface mine drainage

Parameter	Range of contaminant concentrations in flood flow affected by mine discharge ^(a)				Potable water standards ^(b)		Irrigation ^(c)
	Basin		Regional Basin		Maximum Permissible	Recommended Limiting	Recommendations for maximum concentration
	Min.	Max.	Min.	Max.	Concentration	Concentration	for continuous use on all soils (mg/l)
					(mg/l)	(mg/l)	
Total U	0.36	0.76	0.26	0.44	0.015/3.5/0.21 ^(d)		---
Ra-226 + 228	2.1	4.5	1.6	2.6	---	5 pCi/l	5 pCi/l
TSS	107	228	79	131	---	---	---
Sulfate	900	1909	668	1098	---	250	200
Zinc	0.366	0.774	0.271	0.445	---	5.0	2.0
Cadmium	0.02	0.044	0.015	0.025	0.01	---	0.010
Arsenic	0.025	0.054	0.019	0.031	0.05	0.01	0.10

^(a) Concentrations in milligrams per liter, except Ra-226 -228 which are in picocuries per liter.

^(b) Sources: U.S. Environmental Protection Agency (EPA76) and, in the case of uranium, suggested guidance from the National Academy of Sciences (NAS79) to the USEPA and from USEPA (Office of Drinking Water) to the State of Colorado (La79).

^(c) Source: NAS72.

^(d) 0.015 mg/l : Suggested maximum daily limit based on radiotoxicity for potable water consumed at a rate of 2 liters per day on a continuous basis.

3.5 mg/l : Suggested maximum 1-day limit based on chemical toxicity and intake of 2 liters in any one day.

0.21 mg/l : Suggested maximum 7-day limit based on chemical toxicity and intake of 2 liters per day for 7 days.

limit of 10 pCi/l (0.015 mg/l) may not be cost effective, the Agency is contracting to develop the economic and technical basis for a uranium (in water) standard. The National Academy of Science, at the request of the Agency, evaluated the chemical toxicity of uranium. A maximum, 1-day concentration of 3.5 mg/l (7 mg/day based on daily intake of 2 liters) is the "Suggested No Adverse Response Level" (SNARL). The corresponding concentration for a 7-day period is 0.21 mg/l.

There are numerous complicating factors surrounding the foregoing suggested radiotoxicity and chemical toxicity limits for uranium. These include economic justification, technical feasibility, gut to blood transfer factors, and overall health of the receptor, to name a few. Of importance is the fact that a stricter standard for uranium in water is likely and that present NPDES limits of 1 mg/l or previous drinking water limits on the order of 5 to 8 mg/l are or will be superseded. For these reasons, the calculated uranium concentrations in the aqueous pathway are considered relative to the more recent, suggested limits of 0.015, 3.5, and 0.21 mg/l.

Although mine effluents are not considered potable water, they infiltrate shallow aquifers that are potable in terms of the Safe Drinking Water Act. The extent to which shallow aquifers in uranium mining areas are used for potable water supply is presently small, but accurate surveys of well locations and water quality are scarce. If a limit of 0.015 mg/l for uranium in potable water is set, it appears that uranium instead of radium-226 may be the primary pollutant of concern in both surface runoff and related shallow groundwater.

Of the remaining contaminants, sulfate and possibly cadmium might exceed drinking water standards. Cadmium may also limit use of the water for irrigation. These results provide only a rough estimate of water quality effects. There are other stable toxic elements to consider, but there are insufficient data. Multiple mine sources would increase the concentration, but ion exchange, sorption, etc. would reduce them. The net effect is simply unknown. It does appear that uranium, in particular, deserves additional study in light of new interpretations concerning radiotoxicity.

3.3.3.2 Impacts of Seepage on Groundwater

The previous analysis assumed no infiltration (to groundwater) of dissolved or suspended contaminants, thereby creating a maximum or worst-case situation with respect to transport via floodwaters. In fact, contaminants will also infiltrate through the stream deposits. Anions and selected stable elements like uranium, selenium, and molybdenum are most likely to migrate downward. Insofar as the alluvial, valley fill aquifer may be used locally, particularly in the case of larger drainage basins and the regional basin, some analysis of potential impacts is offered herein.

Effects of mine drainage impoundments used to settle suspended solids are excluded from the present analysis. Such impoundments are relatively small, commonly less than 1 or 2 hectares, and tend to become self-sealing due to settling of fines. Potable water supplies at the mines are usually from deep exploration borings converted to water wells or from mine water. Problems may exist with such water being contaminated, as has been documented in the Grants Mineral Belt (EPA75), but we do not believe seepage from settling ponds to be a factor.

Infiltration of water discharged to ephemeral stream courses was not calculated separately. It was combined into a lumped term incorporating infiltration and evaporation. Both losses are, in part, a function of surface area. Infiltration takes place primarily in the basin. When three mines are operating, 22.7 km of perennial stream is created and extends into a portion of the basin. Infiltration of the mine effluent adds primarily to the amount of water in storage in the alluvium, versus acting as a source of recharge to the deeper, consolidated strata.

As with many of the intermontane basins in Wyoming, water in the South Powder River Basin is primarily groundwater recharged by sporadic runoff from limited precipitation (Ke77). Some stock ponds that collect surface runoff are supplemented by groundwater from wells or springs. Mine water discharged from one underground mine is used to irrigate approximately 65 hectares of native grass, alfalfa, oats, and barley. In general, groundwater is not used for irrigation (Ho73). Groundwater use for domestic supplies is largely confined to the Dry Fork of the Cheyenne River (Ke77). The number of wells is close to a density of one per 400 ha (Ke77). Typical wells are completed in the alluvium and yield less than 100 m^3/min .

Geological formations in the southern portion of the Powder River Basin include in descending order and increasing age; the 1) Alluvium, 2) Wasatch Formation, 3) Fort Union Formation, 4) Lance Formation, 5) Fox Hills Formation, and 6) older rocks too deep to be affected by uranium mining (NRC78c). Table 3.28 shows the well depth for each formation, anticipated well yields, and the total dissolved solids content in the vicinity of an active uranium mining and milling project in the South Powder River Basin.

Water quality in the Wasatch and Fort Union Formations ranges widely and appears to correlate with the permeability of the water-bearing sand and proximity to outcrops. No relation of water quality to depth is apparent. Analyses of water from Cenozoic rocks show dissolved solids ranging from less than 100 to more than 8000 mg/l (Ho73). Of the 258 analyses performed by the USGS, 55 showed dissolved solids less than 500 mg/l, 133 less than 1000 mg/l, and 125 more than 1000 mg/l. Sodium, sulfate, and bicarbonate are the dominant ions, and water is usually excessively hard. Iron is characteristically a problem in water from the Wasatch and Fort Union Formations (Ho73). Element distributions show considerable variability due to clay lenses in the sandy units (NRC78c). The clays act as barriers to groundwater movement and preferentially concentrate some elements. Table 3.29 shows the ambient groundwater quality in the immediate area of three active mills in the South Powder River Basin.

In the Wyoming model mine sub-basin, total inflow equals $9 \text{ m}^3/\text{min}$ or $4.73 \times 10^6 \text{ m}^3/\text{yr}$, and total annual infiltration loss equals $4.65 \times 10^6 \text{ m}^3$ (calculated in Appendix H). Restated, 98.2 percent of the discharge infiltrates and the remainder evaporates.

Infiltration of $4.65 \times 10^6 \text{ m}^3/\text{yr}$ is not likely to continue for the full duration of mining unless the bedrock strata have the same or similar permeability as the alluvium and (or) there is an extensive zone of unsaturated alluvium to provide storage. The alluvium in the Wyoming study area is concentrated along the stream axes, is relatively thin, and is underlain by less permeable bedrock strata. It is probable that a zone of saturated alluvium will gradually develop and extend downstream as mine discharge continues. Recharge from the alluvium to the underlying Wasatch or Fort Union Formations will occur but at a low rate compared to infiltration. Water quality in the alluvium is highly variable (Table 3.29); it may or may not be affected by mine drainage. Adverse impacts, if any, are likely to be a result of uranium, sulfate, and mobile elements.

Table 3.28 Northeastern Wyoming groundwater sources

Geologic Period	Aquifer	Depth Range of Wells, m	<u>Anticipated Well Yield, gpm</u>		Total Dissolved Solids, mg/l
			Common	High	
Quaternary	Alluvium	3-30	20-945	1140-2270	106-7340
Tertiary	Wasatch	12-300	4-150	380-2370	160-6620
	Fort Union	45-180	4-110	380	484-3250
Cretaceous	Lance	45-365	4-190	1900	450-3060
	Fox Hills	210-700	75-260	760-1900	1240-3290
	Mesaverde	12-915	57-150	225-265	550-1360
	Cody	30-335	4-20	380-760	6392-12,380
	Frontier	20-610	4-20	380-1135	390-2360
	Dakota	75-1830	95-380	760-3410	218-1820
	Sundance	120-210	4-20	95	894-2310
Triassic	Spearfish	6-275+	4-115	380-760	2590
Pennsylvanian	Minnelusa	75-1980	95-950	1860-7470	255-3620
Mississippian	Pahasapa	150-2320	380-9460	26,500-35,600	290-3290
Ordovician	Bighorn	0-60	3785	3785	427-3219
Cambrian	Flathead	20-1800	760		124

Source: NRC78b.

Table 3.29 Groundwater quality of wells sampled by the three major uranium producers in the South Powder River Basin, Wyoming

Parameter	Range of Concentration Reported (mg/l)		
	Kerr-McGee ^(a)	TVA ^(b)	Exxon ^(c)
pH	7.4-8.0	7.4 - 8.5	7.3-8.1
Spec. cond.			
μmhos/cm	210-1100	250-1300	290-600
Ca	28-343	10-200	26-150
Mg	8-81	2-80	1-13
Na	5-71	10-300	54-121
HCO ₃	30-380	70-110	90-412
SO ₄	28-980	8-1000	58-575
Cl	<5-57	11-25	6-16
Zn	0.006-18.0	0.03 -3	ND- 0.14 ^(d)
Fe		0.2 -20	0.01- 1.64
Ba			ND- 0.05
Radium (pCi/l)	0.41 - 5.18	0.2 -18	0.4 -12.0
Uranium (mg/l)	< 0.002- 2.3	0.002-60	0.0004 - 0.21

^(a) Shallow wells up to 61 meters depth, Tables 2.6-7 through 2.6-10 of reference Ke77.

^(b) From Figs. C1 and C3 of reference NRC78b.

^(c) Table 2.12 of reference NRC78d.

^(d) ND: Not detectable.

An actual example of this saturated front developing and moving down-gradient is present at the Kerr-McGee Nuclear Corporation's Bill Smith Mine in South Powder River Basin (Ke77). The mine discharges to a tributary of Sage Creek at a rate of about $1.7 \text{ m}^3/\text{min}$. From the period January 1974 to late 1976, a flow front 23 km long developed as a result of infiltration into the sandy alluvium. The discharge water maintains a high groundwater level in the stream bed. Unfortunately, no information is available on the geometry of the stream channel to evaluate the volume of water that has infiltrated in the three-year period or on any water quality changes that have occurred.

In summary, additional field data are needed to properly address the water quality effects of infiltration. Both theory and at least one field example indicate extensive infiltration of effluent containing at least some mobile stable and radioactive contaminants. Therefore, we recommend additional field investigations to determine, at the minimum, any hydraulic and water quality effects of mine discharge on shallow aquifers and the influence of dewatering on regional water levels and water quality, regardless of pre-existing or anticipated local water use patterns.

3.3.4 Gases and Dusts from Mining Activities

Dusts and toxic gases are generated from routine mining operations. Combustion products are produced by large diesel and gasoline-powered equipment in the mine and by trucks transporting the overburden, ore, and sub-ore from the pit to storage pile areas. Dusts are produced by blasting, breaking, loading, and unloading rock and ore and by haulage trucks moving along dirt roads. Finally, Rn-222 will emanate from exposed ore in the pit and from the ore as it is broken, loaded, and unloaded. These sources will be discussed individually.

3.3.4.1 Dusts and Fumes

Most vehicular emissions are from the combustion of hydrocarbon fuels in heavy-duty, diesel-powered mining equipment. Surface mines produce considerably more emissions than underground mines, since the overburden must be removed before the ore can be mined. The principal emissions are particulates, sulfur oxides, carbon monoxide, nitrogen oxides, and hydrocarbons. The quantity of these combustion products released to the atmosphere depends on the number, size, and types of equipment used.

The EPA estimates the following emissions from mining 1350 MT of ore per day from a surface mine (Re76).

<u>Pollutant</u>	<u>Emissions per Operating Day, kg/d</u>	
	<u>Mining Operations</u>	<u>Overburden Removal</u>
Particulates	17.0	18.9
Sulfur oxides	35.4	39.3
Carbon monoxide	294.2	327.4
Nitrogen oxides	484.6	538.4
Hydrocarbons	48.4	53.8

Assuming a 330 operating-day-year (Ni79), we adjusted these emission rates to ore production for the average surface mine (1.2×10^5 MT/yr) and the average large surface mine (5.1×10^5 MT/yr) as described in Sections 1.3.1 and 3.3.1. Table 3.30 shows the total airborne combustion product emissions. These estimated emission rates are somewhat higher than rates previously suggested by the U.S. Atomic Energy Commission (AEC74).

Table 3.30 Estimated air pollutant emissions from heavy-duty equipment at surface mines

<u>Pollutant</u>	<u>Emissions, MT/yr^(a)</u>	
	<u>Average Mine^(b)</u>	<u>Average Large Mine^(c)</u>
Particulates	3	14
Sulfur oxides	7	28
Carbon monoxide	55	235
Nitrogen oxides	91	387
Hydrocarbons	9	39

(a) Based on (Re76) and 330 operating days per year (Ni79).

(b) Ore production = 1.2×10^5 MT/yr.

(c) Ore production = 5.1×10^5 MT/yr.

Dust is produced from blasting, scraping, loading, transporting, and dumping ore, sub-ore, and overburden. Additional dust is produced when the ore is reloaded from the stockpile for transportation to the mill. Dust emissions vary widely, depending upon moisture content, amount of fines, number and types of equipment operating, and climatic conditions. Because ore is usually wet, the relative amounts of dust produced from mining and handling it are usually small. We selected the following emission factors from those suggested by the EPA for the above listed mining activities (Hu76, Ra78, Da79):

$$\text{Blasting} = 5 \times 10^{-4} \text{ kg dust/MT}$$

$$\text{Scraping and bulldozing} = 8.5 \times 10^{-3} \text{ kg dust/MT}$$

$$\text{Truck loading} = 2.5 \times 10^{-2} \text{ kg dust/MT}$$

$$\text{Truck dumping} = 2 \times 10^{-2} \text{ kg dust/MT}$$

We applied these emission factors to the ore, sub-ore, and overburden production rates of the average mine and average large mine and estimated average annual dust emissions for these mining activities (see Table 3.31). These are probably maximum emission rates because blasting is not always required, and some emission factors appear to have been based upon data from crushed rock operations, which would contain more fines than rock removed from surface mines. One-half the emission factor values were applied to ore and sub-ore because they are usually wet, except when reloading ore from the stockpile, in which case it is assumed to have dried during the 41-day residence period (Section 3.3.1.2).

The movement of heavy-duty haul trucks is probably the largest single source of dust emissions at surface mines. An emission factor (EF) for this source can be computed by the following equation (EPA77b).

$$EF = 2.28 \times 10^{-4} (s) \left(\frac{V}{48} \right) \frac{365-W}{365} (TF) (f) \quad (3.2)$$

where,

EF = Emission factor, MT/vehicle kilometer traveled (MT/VKmt),

S = Silt content of road surface, percent,

V = Vehicle velocity, kmph [Note: This term becomes $\left(\frac{V}{48}\right)^2$ for velocities less than 48 km/hr (EPA77b, DA79)],

W = Mean annual number of days with 0.254 mm or more rainfall,

TF = Wheel correction factor, and

f = Average fraction of emitted particles in the <30 μm diameter suspended particle size range; particles having diameters greater than 30 μm will settle rapidly near the roadway.

Values selected for these terms in the solution of Equation 3.2 are --

S = 10 percent (Da79),

V = 32 km/hr for heavy-duty vehicles and 48 km/hr for light vehicles (therefore, the velocity term is $(32/48)^2$ and $(48/48)$, respectively),

W = 90 days (EPA77b),

TF = 2.5 (Da79) (heavy-duty vehicles only), and

f = 0.60, since the weight percent of particles of less than 30 μm and greater than 30 μm in diameter is generally considered to be 60 and 40 percent, respectively (EPA77b).

Substituting these values into Equation 3.2 yields 1.15×10^{-3} MT/VKmt and 1.03×10^{-3} MT/VKmt for the emission factors of heavy-duty haul trucks and light duty vehicles, respectively.

Table 3.31 shows estimated dust emissions for the movement of heavy-duty haul trucks using the following information:

Table 3.31 Average annual dust emissions from mining activities

Mining Activity	Dust Emissions, MT/yr					
	Average Mine ^(a)			Average Large Mine ^(b)		
	Ore ^(c)	Sub-ore ^(c)	Overburden	Ore ^(c)	Sub-ore ^(c)	Overburden
Blasting	0.03	0.03	3.0	0.13	0.13	20
Scraping/bulldozing	NA ^(d)	NA	51	NA	NA	340
Truck Loading	<u>1.5</u>	<u>1.5</u>	<u>150</u>	<u>6.4</u>	<u>6.4</u>	<u>1000</u>
Total at Pit Site	<u>1.53</u>	<u>1.53</u>	<u>204</u>	<u>6.53</u>	<u>6.53</u>	<u>1360</u>
Truck Dumping	1.2	1.2	120	5.1	5.1	800
Reloading stockpiled ore ^(e)	<u>3.0</u>	<u>NA</u>	<u>NA</u>	<u>13</u>	<u>NA</u>	<u>NA</u>
Total at Pile Sites	<u>4.2</u>	<u>1.2</u>	<u>120</u>	<u>18.1</u>	<u>5.1</u>	<u>800</u>
Vehicular dust ^(f)	14	14	304	59	59	2020
Wind suspended dust from storage piles	10	3	30	44	10	94

(a) Based on annual production rates of 1.2×10^5 MT of ore and sub-ore and 6.0×10^6 MT of overburden.

(b) Based on annual production rates of 5.1×10^5 MT of ore and sub-ore and 4.0×10^7 MT of overburden.

(c) Assumed wet.

(d) NA - not applicable.

(e) Assumed dry.

(f) Dust emissions from heavy-duty vehicular traffic along ore, sub-ore and overburden haul roads.

EF = 1.15×10^{-3} MT/VKmt,

Truck capacities = 31.8 MT for ore and sub-ore and

109.1 MT for overburden (Da79),

Round-trip haul distance = 3.2 km to ore and sub-ore piles

and 4.8 km to overburden dump, and

Annual production rates = given in Section 3.3.1 and in the footnotes of Table 3.31.

Additional dust emissions will occur from the movement of light-duty vehicles along access roads. Using the emission factor derived above (1.03×10^{-3} MT/VKmt) and assuming that there are 24 km of access roads traveled 4 times a day for 330 operating days per year, about 33 MT of dust will be produced from this source annually. Emissions during haulage road maintenance is relatively small and will not be considered.

Table 3.31 also shows average annual dust emissions from wind erosion of overburden, sub-ore, and ore piles at the model surface mines. For these computations, we assumed the model overburden pile to be that of Case 2 and in the shape of a 65-m high truncated cone (Table 3.11). The same was assumed for the average mine, except the pile height was 30 m. The sub-ore piles of both mines were assumed to have a truncated cone configuration (Table 3.20). The same configuration was also assumed for the ore piles, but the pile heights were 9.2 m for the average large mine and 3.1 m for the average mine (Table 3.17).

Emission factors, computed in Appendix I, are 0.850 MT/hectare-yr for overburden and sub-ore piles and 0.086 kg/MT for the ore stockpiles. The first emission factor was multiplied by the overburden and average sub-ore pile areas; the second factor was multiplied by the annual ore production.

In computing the Table 3.31 dust emissions, we assumed no effective dust control program and that there was no vegetation on overburden and sub-ore piles. Haul roads are normally sprinkled routinely during dry periods, and stabilizing chemicals are applied primarily to ore haul roadways at some mines. Sprinkling can reduce dust emissions along haul roads by 50 percent, and up to 85 percent by applying stabilizing chemicals (EPA77b, Da79).

The dust emissions from vehicular traffic (Table 3.31) (transportation) were summed with those produced by light vehicular traffic (33 MT/yr) and considered as one source of emissions. Concentrations of contaminants in the dust are unknown. Some spillage of ore and sub-ore along haul roads will undoubtedly raise uranium levels in roadbed dust. As an estimate, uranium and daughter concentrations in the dust were considered to be twice background, 8 ppm (2.7 pCi/g), while concentrations of all other contaminants were considered to be similar to those in overburden rock (Section 3.3.1.1, Table 3.16). Table 3.32 shows the annual emissions computed with these assumptions.

Table 3.33 lists annual contaminant emissions from mining activities (scrapping, loading, dumping, etc.) according to source location, at the pit and at the piles. Contaminant emissions were computed by multiplying the total annual dust emissions at each pile (Table 3.31) by the respective contaminant concentrations in each source -- overburden (Section 3.3.1.1; Table 3.16), sub-ore (Section 3.3.1.3; Table 3.19) and ore (Section 3.3.1.2; Table 3.19). Contaminant emissions at the site of the pit were computed by multiplying the total annual dust emissions of ore, sub-ore, and overburden (Table 3.31) by their respective contaminant concentrations. The three products of the multiplication were then summed to give the values in the 4th and 8th data columns of Table 3.33. The health impact of the sources at each location will be assessed separately in Section 6.1.

Table 3.34 lists annual contaminant emissions due to wind suspension and transport of dust. These values were computed by multiplying the annual mass emissions (Table 3.31) by the contaminant concentrations in overburden, sub-ore, and ore listed in Sections 3.3.1.1, 3.3.1.3, and 3.3.1.2, respectively. The uranium and uranium daughter concentrations were also multiplied by an activity ratio (dust/source) of 2.5 (Section 3.3.1.2). Although some metals may also be present as secondary deposits, it was believed that there were insufficient data to justify multiplying their concentrations by the 2.5 ratio.

3.3.4.2 Radon-222 from the Pit, Storage Piles, and Ore Handling

Rn-222 will be released from the following sources during surface mining operations:

Table 3.32 Average annual emissions of radionuclides (μCi) and stable elements (Kg) from vehicular dust at the model surface mines

Contaminant	Average Large Surface Mine ^(a)	Average Surface Mine ^(b)
Arsenic	20	3.3
Barium	630	106
Copper	39	6.6
Chromium	< 111	< 19
Iron	13,030	2,190
Mercury	< 17	< 2.9
Potassium	15,200	2,560
Manganese	1,050	177
Molybdenum	5.4	0.9
Lead	48	8.0
Selenium	4.3	0.7
Strontium	330	55
Vanadium	220	37
Zinc	43	7.3
Uranium-238 and each daughter	5,860	990
Thorium-232 and each daughter	2,170	370

(a) Mass emissions = 2,170 MT/yr.

(b) Mass emissions = 365 MT/yr.

Table 3.33 Average annual emissions of radionuclides (μCi) and stable elements (kg) from mining activities at the model surface mines

Contaminant	Average Surface Mine ^(a)				Average Large Surface Mine ^(a)			
	Overburden Pile Site	Sub-ore Pile Site	Ore Pile Site	Pit Site	Overburden Pile Site	Sub-Ore Pile Site	Ore Pile Site	Pit Site
Arsenic	1.1	0.10	0.36	2.1	7.2	0.44	1.6	13
Barium	35	1.1	3.9	62	232	4.7	17	406
Cobalt	NR ^(b)	0.02	0.07	0.05	NR	0.08	0.29	0.21
Copper	2.2	0.07	0.26	3.9	14	0.31	1.1	25
Chromium	<6	0.02	0.08	<10	<41	0.10	0.36	<70
Iron	720	19	66	1,270	4,800	80	284	8,360
Lead	2.6	0.09	0.33	4.7	18	0.40	1.4	31
Magnesium	NR	4.2	15	11	NR	18	63	46
Manganese	58	1.2	4.0	102	388	4.9	17	672
Mercury	<1	ND ^(c)	ND	<1.6	<6.4	ND	ND	<11
Molybdenum	0.3	0.14	0.48	0.86	2.0	0.59	2.1	4.9
Nickel	NR	0.02	0.08	0.06	NR	0.10	0.36	0.26
Potassium	840	30	105	1,500	5,600	128	453	9,850
Selenium	0.2	0.13	0.46	0.74	1.6	0.56	2.0	4.2
Strontium	18	0.16	0.55	31	120	0.66	2.4	206
Vanadium	12	1.7	5.9	25	80	7.2	26	154
Zinc	2.4	0.04	0.12	4.2	16	0.15	0.52	28
Uranium-238 & each daughter	1,800	120	2,990	4,300	12,000	510	12,900	25,700
Thorium-232 & each daughter	120	2.4	42	220	800	10	180	1,440

(a) Mass emissions from Table 3.31.

(b) NR - Not reported.

(c) ND - Not detected.

Table 3.34 Average annual emissions of radionuclides (μCi) and stable elements (kg) in wind suspended dust at the model surface mines

Contaminant	Average Large Surface Mine			Average Surface Mine		
	Overburden	Sub-Ore	Ore	Overburden	Sub-Ore	Ore
	Pile	Pile	Stockpile	Pile	Pile	Stockpile
Arsenic	0.85	0.86	3.8	0.27	0.26	0.86
Barium	27	9.2	40	8.7	2.8	9.2
Cobalt	NR ^(a)	0.16	0.70	NR	0.05	0.16
Copper	1.7	0.61	2.7	0.54	0.18	0.61
Chromium	<4.8	0.20	0.88	<1.5	0.06	0.20
Iron	564	157	690	180	47	157
Mercury	<0.75	ND ^(b)	ND	<0.24	ND	ND
Potassium	660	250	1,100	210	75	250
Magnesium	NR	35	154	NR	11	35
Manganese	46	9.6	42	15	2.9	9.6
Molybdenum	0.24	1.2	5.0	0.08	0.35	1.2
Nickel	NR	0.20	0.88	NR	0.06	0.20
Lead	2.1	0.78	3.4	0.66	0.23	0.78
Selenium	0.19	1.1	4.8	0.06	0.33	1.1
Strontium	14	1.3	5.7	4.5	0.39	1.3
Vanadium	9.4	14	62	3.0	4.2	14
Zinc	1.9	0.29	1.3	0.60	0.09	0.29
Uranium-238 & each daughter	1,410	1,000	31,300	450	300	7,100
Thorium-232 & each daughter	94	20	440	30	6.0	100

(a) NR - Not reported.

(b) ND - Not detected.

1. Ore, sub-ore, and overburden during rock breakage and loading in the pit and unloading on the respective piles. (Since rock breakage, loading, transporting, and unloading usually occur in a short time period, they are considered one release.)
2. Ore during reloading from the stockpile after a 41-day residence time (Section 3.3.1.2).
3. Exposed surfaces of overburden, ore, and sub-ore in the active pit area.
4. Overburden, ore, and sub-ore pile surfaces.

The annual quantities of Rn-222 released from sources 1 and 2 above were computed using the following factors and assumptions:

1. Rn-222 is in secular equilibrium with U-238.
2. The density of ore, sub-ore, and overburden is 2.0 MT/m^3 .
3. Annual production rates of ore, sub-ore, and overburden are those given previously in this Section and in footnotes "a" and "b" of Table 3.31.
4. All Rn-222 present, 0.00565 Ci/m^3 per percent U_3O_8 , is available with an emanation coefficient of 0.27. [Although an emanation coefficient of 0.2 is commonly used (Ni79), recent emanation-coefficient measurements for 950 samples of domestic uranium ores by the Bureau of Mines indicate a value between 0.25 and 0.3 to be more appropriate (Au78, Tanner, A.B., Department of Interior, Geological Survey, Reston, VA, 11/79, personal communication). Therefore, an emanation coefficient of 0.27 was selected.]
5. The quantities of U_3O_8 present in ore, sub-ore, and overburden are 0.10, 0.015, and 0.0020 percent, respectively.

Substituting these values into the following equation yields the Rn-222 releases given in Table 3.35 for the average mine and the average large mine.

$$\text{Rn-222 (Ci/yr)} = (\text{Percent } \text{U}_3\text{O}_8) \left(\frac{0.00565 \text{ Ci}}{\text{m}^3 \times \text{percent}} \right) (0.27) \left(\frac{\text{m}^3}{2.0 \text{ MT}} \right) \quad (3.3)$$

$$\times (\text{Production Rate, } \frac{\text{MT}}{\text{yr}})$$

The quantities of Rn-222 that emanate from exposed overburden, ore, and sub-ore surfaces in the pit were estimated by the following method. Exposed

surface areas of ore and sub-ore are assumed equal since equal quantities of each are mined. The computation assumes an ore plus sub-ore zone 12 m thick (h_1) in the shape of a truncated cone with 45 degree sloping sides (Fig. 3.14). The radii of the zone, r_1 and r_2 , can be computed using the following equation from the relationship $r_2 = r_1 + 12$ and the volumes of ore plus sub-ore mined in a 2.4 year period -- $1.22 \times 10^6 \text{ m}^3$ and $2.8 \times 10^5 \text{ m}^3$ at the average large mine and average mine, respectively (the bulking factor is not considered in computing the pit volume).

$$V (\text{ore} + \text{sub-ore zone}) = 1/3 \pi h_1 (r_1^2 + r_1 r_2 + r_2^2) \quad (3.4)$$

The computed radii, r_1 and r_2 , were 174 m and 186 m at the average large mine and 80 m and 92 m at the average mine. The surface areas (S_A) of exposed ore and sub-ore in the pit are then one-half that given by the equation,

$$S_A = 1/2 \pi (d_1 + d_2)(\text{slant height}) + \pi r_1^2, \quad (3.5)$$

where d_1 and d_2 are the diameters related to r_1 and r_2 . Exposed surface areas of ore and sub-ore were computed to be equal and $57,170 \text{ m}^2$ at the average large mine and $14,650 \text{ m}^2$ at the average mine.

The shape of the overburden zone was assumed to be the same as the ore and sub-ore zone (Fig. 3.14). The thickness, h_2 , and radius, r_3 , of this zone can be computed using the following equation with the relationship, $r_3 = r_2 + h_2$, and knowing the volume-- $4.8 \times 10^7 \text{ m}^3$ and $7.2 \times 10^6 \text{ m}^3$ --at the average large mine and average mine, respectively.

$$V (\text{overburden}) = 1/3 \pi h_2 (r_2^2 + r_2 r_3 + r_3^2) \quad (3.6)$$

Since r_2 was computed above to be 186 m at the average large mine and 92 m at the average mine, Equation 3.6 becomes

$$4.8 \times 10^7 = 1.087 \times 10^5 h_2 + 584 h_2^2 + 1.047 h_2^3 \quad (3.7)$$

for the average large mine, and

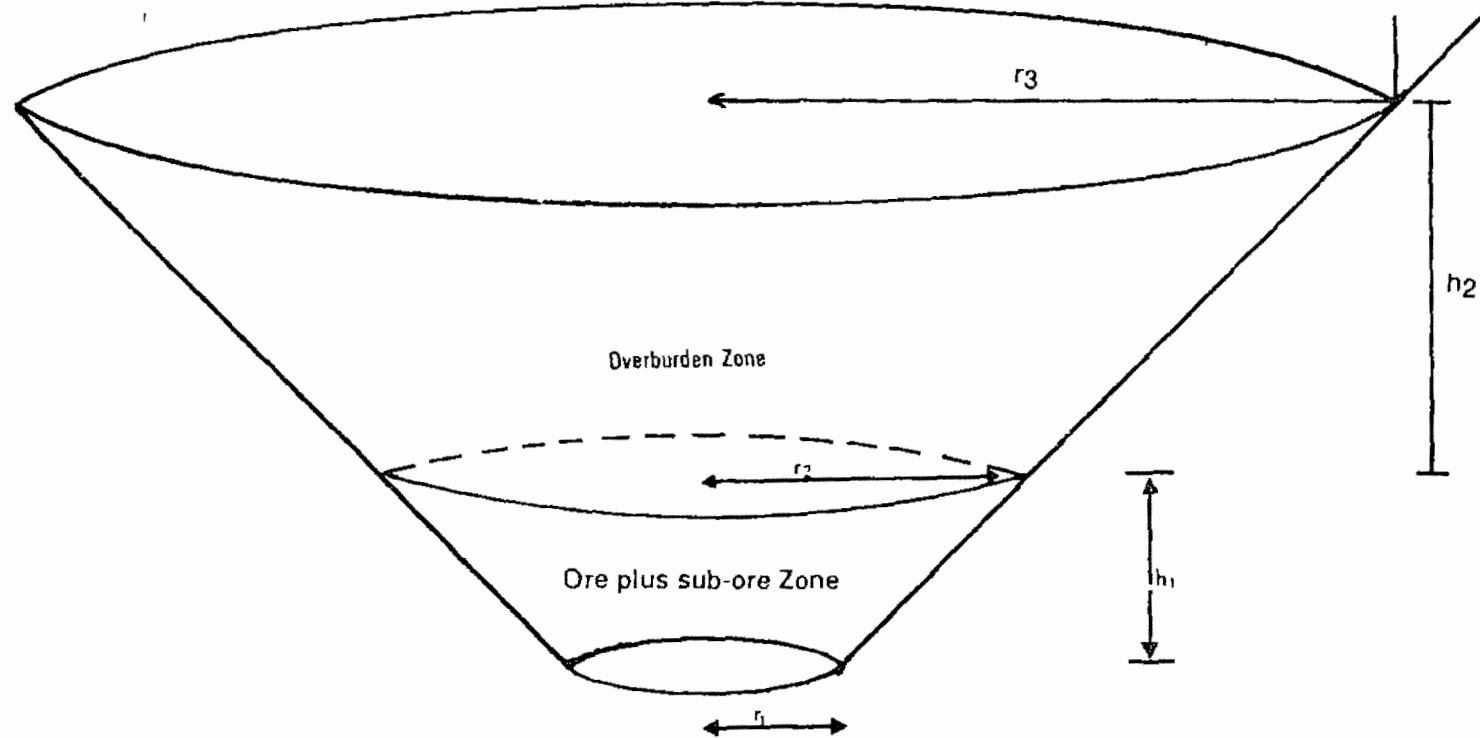


Figure 3.14 Configuration of open pit model mines.

$$7.2 \times 10^6 = 2.659 \times 10^4 h_2 + 289 h_2^2 + 1.047 h_2^3 \quad (3.8)$$

for the average mine.

Solving these equations yields the following parameters:

	h_2	r_3	r_2
average large mine	188 m	374 m	186 m
average mine	105 m	197 m	92 m

The surface area (S_A) of the exposed overburden is then given by the following equation.

$$S_A = 1/2 \pi (d_2 + d_3) (\text{slant height}), \quad (3.9)$$

where d_1 and d_2 are the diameters related to r_2 and r_3 . Areas computed were $4.68 \times 10^5 \text{ m}^2$ and $1.34 \times 10^5 \text{ m}^2$ for the average large mine and average mine, respectively.

Multiplying the exposed ore, sub-ore, and overburden areas by their U_3O_8 contents (0.10%, 0.015% and 0.002%, respectively) and by a Rn-222 exhalation rate of 0.092 Ci/m^2 per year per percent U_3O_8^* and summing gives the annual Rn-222 releases shown in Table 3.35.

The emanation of Rn-222 from overburden, sub-ore, and ore storage piles is based on an exhalation rate of 0.092 Ci/m^2 per yr per percent U_3O_8 (Ni79), and ore grades of 0.002 percent, 0.015 percent, and 0.10 percent, respectively. The surface areas used were those computed previously for the case 2 model mines and listed in Tables 3.11, 3.17 and 3.20. The areas for the average large mine and average mine are $1.1 \times 10^6 \text{ m}^2$ and $2.2 \times 10^5 \text{ m}^2$ for overburden piles, $1.2 \times 10^5 \text{ m}^2$ and $3.6 \times 10^4 \text{ m}^2$ for sub-ore piles, and $6.2 \times 10^3 \text{ m}^2$ and $3.6 \times 10^3 \text{ m}^2$ for the ore piles, respectively. Applying these parameters, the annual Rn-222 emissions from the overburden, sub-ore, and ore piles at the average mine and average large mine were computed. Table 3.35 presents the results.

The total annual Rn-222 released during surface mining operations is the sum of the releases from the sources considered: 331 Ci from the average mine and 1261 Ci from the average large mine. Considering ore production and

*The average value of measured exhalation rates at surface uranium mines (Ni79).

Table 3.35 Radon-222 releases during surface mining, Ci/yr

Source	Average Mine	Average Large Mine
Ore loading and unloading	9	39
Reloading ore from stockpile	9	39
Sub-ore loading and unloading	1	6
Overburden loading and unloading	9	61
Exposed surface of overburden, ore, and sub-ore in the pit	180	691
Ore stockpile exhalation	33	57
Sub-ore pile exhalation	50	166
Overburden pile exhalation	<u>40</u>	<u>202</u>
Total	331	1261

grade differences, these values agree reasonably well with those computed by other procedures (Tr79).

3.4 Underground Mining

3.4.1 Solid Wastes

During underground mining, like surface mining, materials are removed, separated according to ore content, and stored on the surface for various periods of time (Section 1.3.3). These separate piles consist of waste rock produced from shaft sinking operations and from cutting inclines, declines, and haulage drifts through barren rock, sub-ore, and ore. The waste rock is similar to overburden removed at surface mines, except much smaller quantities are involved and none are returned to the mine. However, as mining progresses, waste rock is sometimes used to backfill mined out areas of the mine and retained beneath the surface. The ore and sub-ore will also be similar in nature to those described previously for surface mines, as is their potential to be sources of contamination to the environment (Fig. 3.15).

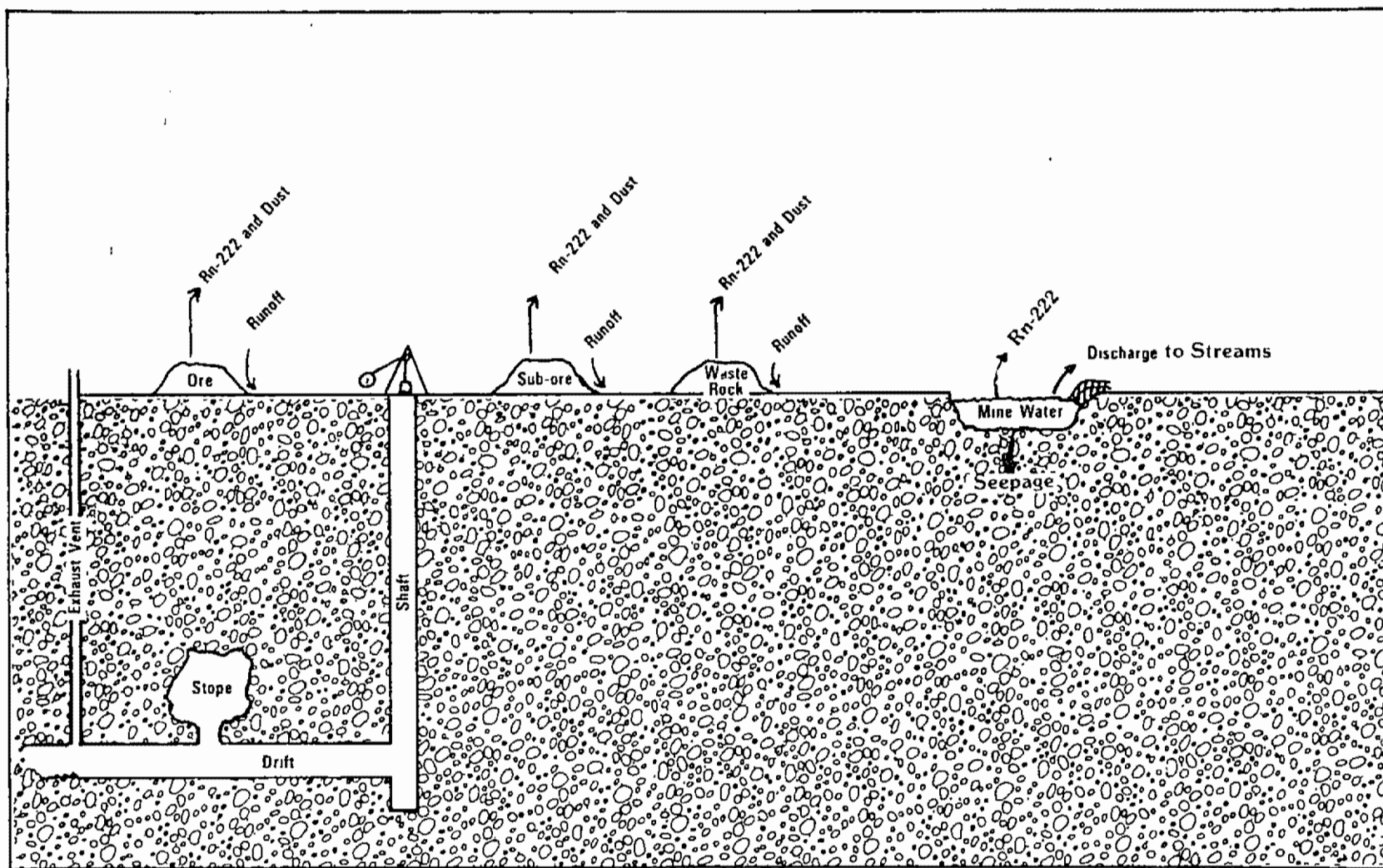


Figure 3 15 Potential sources of environmental contamination from active underground uranium mines

3.4.1.1 Waste Rock Piles

Much smaller quantities of waste rock accumulate at underground mines than overburden at surface mines. The weight ratio of waste rock to ore depends mainly upon the size, depth, and age of the mine. During the initial mining stages, all material removed is waste rock. As entry into the ore body occurs and ore mining begins, the quantity of waste rock removed per metric ton of ore decreases sizably. Once in the ore body, as little waste rock as possible is mined. The ratio of ore to waste rock removed from underground mines varies considerably. At seven presently active underground mines, the ore to waste rock ratio varies from 1.5:1 to 16:1, with an average ratio of 9.1:1 (Jackson, P.O., Battelle Pacific Northwest Laboratory, Richland, WA, 12/79, personal communication). As future mines become larger and deeper, the overall ore to waste rock ratio will probably decrease.

Since the annual average ore capacity of underground mines was 1.8×10^4 MT in 1978 (Section 1.3.1), the average of the 305 underground mines would have produced 2.0×10^3 MT of waste rock during that year, assuming the average 9.1:1 ore to waste rock ratio. This will be considered the production rate of the "average underground mine." Like surface mines, relatively few of the 305 active underground mines account for a significant portion of the total ore produced by the underground method. Also, future underground mines are expected to have larger capacities than many of the current mines (Th79). Therefore, a second underground mine will be considered, which is defined as the "average large mine." Its annual ore production rate is assumed to be 2×10^5 MT, the average ore capacity of five large underground operations (Ja79b, TVA79, TVA76, TVA78a, TVA78b). The quantity of waste rock removed annually will be 2.2×10^4 MT, assuming the ore to waste rock ratio to be the same as for the average mine. Assuming the density of waste rock to be about 2.0 MT/m^3 and a bulking factor of 1.25 (Burris, E., Navajo Engineering Construction Authority, Shiprock, N.M., 2/80, personal communication), the average mine and average large mine will produce waste rock at an annual rate of $1.3 \times 10^3 \text{ m}^3$ and $1.4 \times 10^4 \text{ m}^3$, respectively. Since waste rock is not presently used to backfill mined-out areas, this rate of accumulation will continue for the life of the mine, which is assumed to be the same as that for an open pit mine, 17 years.

Table 3.36 lists estimated average surface areas of the waste rock piles during the lifetimes of the two mines defined above. The following para-

Parameter	Average Mine	Average Large Mine
Waste rock production rate, MT/yr	2.0×10^3	2.2×10^4
Rock density, MT/m ³	2.0	2.0
Bulking factor	1.25	1.25
Waste rock volume, m ³ /yr	1.3×10^3	1.4×10^4
Active mine life, yr	17	17
Pile height, m	6	12

These estimated areas assume no backfilling and that the piles are on level terrain. Because waste rock is sometimes used to backfill and is often dumped into a gorge or ravine, these surface areas represent maximum conditions.

The mineralogy, physical characteristics, and composition of waste rock from underground mines are assumed to be identical to the overburden removed from open pit mines (Section 3.3.1.1). Also, reclamation procedures for waste rock piles at underground mines should be similar to those described in Section 3.3.1.4 for overburden dumps.

3.4.1.2 Ore Stockpiles

Because ore is often stockpiled at the mine and/or at the mill, it becomes a potential source of contamination to the mine environment during the storage period. These piles will be smaller than the waste rock piles, but the concentration of most contaminants in the ore-bearing rock will be much greater.

Ore stockpile residence times can vary considerably with time and ore management. Residence times commonly range from a few days to a few months. The same residence time will be assumed for underground mines as was selected above for surface mines, 41 days. Assuming a 330 operating-day-year and a 1.25 bulking factor, the ore stockpiles of the average mine and average large mine will contain 1,400 m³ and 15,500 m³ of ore, respectively. The surface areas of the ore stockpiles were computed using these volumes and assuming 3.1 m high rectangular piles (NRC78a). Table 3.37 lists the estimated surface areas.

Table 3.36 Estimated average surface areas of waste rock piles at underground mines

Mine Size	Average Accumulation, ^(a) m ³	Surface Area of Pile, m ²	Surface Area of Pad, m ²
Average mine ^(b)	1.1×10^4	2,700	2,460
Average large mine ^(c)	1.2×10^5	14,100	12,800

^(a) Assumes average volume of waste rock accumulated during 17-yr. mine life with no backfilling (1/2 total volume accumulation).

^(b) Annual waste rock production = 2.0×10^3 MT.

^(c) Annual waste rock production = 2.2×10^4 MT.

Note.--Waste rock piles are rectangular with length twice the width and sides sloping at 45° (Fig. 3.8 a).

Table 3.37 Estimated surface areas of ore stockpiles at underground mines

Mine Size	Steady State Accumulation, ^(a) m ³	Surface Area of Pile, m ²	Surface Area of Pad, m ²
Average mine ^(b)	1,400	680	620
Average large mine ^(c)	15,500	5,800	5,480

^(a) Assume 41-day residence time.

^(b) Annual ore production = 1.8×10^4 MT.

^(c) Annual ore production = 2×10^5 MT.

Note.--Ore stockpiles are rectangular with length twice the width and sides sloping at 45° (Fig. 3.8 a). Pile height is assumed to be 3.1 m (NRC78a).

The mineralogy, physical characteristics, and composition of ore from underground mines are assumed to be identical to the ore removed from surface mines (Section 3.3.1.2). The U_3O_8 grade of ore may average somewhat higher from underground mines than from surface mines. However, a grade of 0.1 percent U_3O_8 probably approximates reasonably well the ore reserves minable by the underground method (DOE79). Uranium and its decay products in airborne dust from these ore piles will be concentrated by a factor of 2.5 (Section 3.3.1.2).

3.4.1.3 Sub-Ore Piles

The quantity of sub-ore mined at an underground mine, as at a surface mine, is considered to be about equal to the quantity of ore mined, 1.8×10^4 MT at the average mine and 2×10^5 MT at the average large mine. Assuming sub-ore to have a density of 2.0 MT/m^3 and after removal a bulking factor of 1.25, the average volume of sub-ore to be on the surface during the 17-yr operational life of the average mine and average large mine will be $9.6 \times 10^4 \text{ m}^3$ and $1.1 \times 10^6 \text{ m}^3$, respectively (i.e., one-half the total of 17-yr accumulation).

Although sub-ore is often placed on top of piles of previously mined waste rock (Perkins, B.L., New Mexico Energy and Minerals Department, Santa Fe, NM, 12/79, personal communication), we assumed separate rectangular piles in computing the surface areas of the piles at the model mines. Table 3.38 lists the estimated surface and pad areas of the sub-ore piles. These computations were based on pile heights of 6 m at the average mine and 12 m at the average large mine.

At underground mines, the cutoff grade ranges from 0.02 to 0.05 percent U_3O_8 , yielding an average sub-ore grade of 0.035 percent U_3O_8 (99 pCi/g) (Perkins, B.L., New Mexico Energy and Minerals Department, Santa Fe, N.M., 12/79, personal communication). The mineralogy, physical characteristics, and other constituents of sub-ore from underground mines are assumed identical to the sub-ore removed from surface mines (Section 3.3.1.3).

Table 3.38 Estimated average surface areas of sub-ore
piles at underground mines

Mine Size	Average Accumulation, (a) m ³	Surface Area of Pile, m ²	Surface Area of Pad, m ²
Average mine ^(b)	9.6×10^4	18,800	17,700
Average large mine ^(c)	1.1×10^6	104,900	99,400

(a) One-half that which will accumulate during the 17-yr mine life.

(b) Annual sub-ore production = 1.8×10^4 MT.

(c) Annual sub-ore production = 2.0×10^5 MT.

Note.--Sub-ore piles are rectangular with length twice the width and sides sloping at 45° (Fig. 3.8a).

3.4.2 Mine Water Discharge

3.4.2.1 Data Sources

Information concerning the amount and quality of water discharged from underground uranium mines in New Mexico is from field surveys conducted in 1975 (EPA75, P. Frenzel, USGS, written communication, 1979) and Wogman (Wo79), from site-specific environmental impact statements and reports, from NPDES permits, and from a State study (Pe79).

Many mining companies maintain that permits are not required because the formerly ephemeral streams into which discharge occurs are, in effect, a result of the discharges and do not meet the definition of navigable bodies of water. Nevertheless, the companies have applied for permits, together with a request to the courts for a ruling concerning their necessity.

The New Mexico district office of the U.S. Geological Survey (L. Beal, USGS, written communication, 1979) provided discharge rate and volume for the regional drainage systems, namely the Rio San Jose, Rio Puerco (east), and the Rio Grande. We followed procedures developed by the USGS (Bo70) to calculate runoff from ungaged basins.

3.4.2.2 Quality and Quantity of Discharge

To estimate average or typical conditions for mine water discharge, 11 projects in Colorado, New Mexico, and Utah were selected. Table 3.39 shows the summarized flow and water quality data. The center of current domestic underground mining is in the Colorado Plateau and the San Juan Basin. In this area, there is an increasing trend toward underground mining. In Wyoming, both underground and surface mining activity are significant. In Texas, surface mining and, to a lesser extent, in situ leaching are the principal methods used. Climatic and geologic characteristics and land and water use patterns in the Colorado-Utah-New Mexico uranium area are broadly similar; and the Grants Mineral Belt in general and the Ambrosia Lake District in particular are representative of this area. There are many complicating variables such as the geologic and geochemical characteristics of the ore body and host rock. Water-yield and quality associated with mines also vary within the region, as do the size and relative location of the populace. The Grants Mineral Belt scenario is conservative. The mines discharge relatively large amounts of water to streams that are used for irrigation and stock watering and that flow by or through local centers of population.

Table 3.39 shows discharge from selected underground uranium mines in the Colorado Plateau areas of Colorado, New Mexico, and Utah. On the average, discharge is $2.78 \text{ m}^3/\text{min}$, with a standard deviation of $4.34 \text{ m}^3/\text{min}$. The selected underground mines discharge an amount of water similar to that from the Wyoming surface mines. In the Grants Mineral Belt area, average flow from 28 underground mines is $2.4 \text{ m}^3/\text{min}$ (J. Dudley, New Mexico Environmental Improvement Division, written communication). Of the 27 active underground mines being dewatered, 17 discharge to the environment at an average rate of $3.2 \text{ m}^3/\text{min}$. The remainder are in a closed circuit. That is, their discharge is used as mill feed water. The range for 17 mines is 0.2 to $19 \text{ m}^3/\text{min}$. Average discharges from New Mexico underground mines are significantly greater than those from mines in Colorado and Utah, which average $0.68 \text{ m}^3/\text{min}$. Most of the ore production in New Mexico has been from mines 200 to 300 meters deep. In recent years, mines have become progressively deeper and involve more dewatering. For example, the Gulf Mount Taylor mine, which is not yet producing ore, discharges $15 \text{ m}^3/\text{min}$ and will produce ore from a depth

of 1,200 meters. Most of the water is now diverted to a nearby ranch for irrigation and stock watering. When the mill goes on line, most of the mine water will be used there.

Of the 16 active mines in the Ambrosia Lake district, 13 discharge to offsite areas at an average rate of approximately $1.6 \text{ m}^3/\text{min}$. For modeling and to be conservative, we assumed that 14 active mines are present in the model mine area and that the average discharge rate per mine is $2.0 \text{ m}^3/\text{min}$. This is somewhat less than the average condition for the Grants Mineral Belt ($3.2 \text{ m}^3/\text{min}$) as a whole in terms of discharge rate, but the high density of mines assumed present in the model area partly compensates for the difference.

For the New Mexico project shown in Table 3.39, numbers 4, 5, 6, and 7 have discharge that comes directly from the mine portal to settling ponds before discharge. Neither ion exchange for uranium recovery nor barium chloride treatment for radium removal is used. Facilities 8 through 11 use ion exchange columns for uranium removal before discharge. Settling may or may not be used, depending on the suspended solids content of the particular discharge. Project number 10 removes radium prior to discharge. Radium concentrations in the combined effluent from two active mines in the Church-rock area (projects 8 and 4), both of which use settling ponds as the only treatment, have ranged from 1.9 to 8.9 pCi/l since 1975. In the first survey (EPA75), effluent from these same mines contained 30.8 and 7.9 pCi/l. The combined discharge from both mines was sampled by the U.S. Geological Survey in 1975, 1977, and 1978 (P. Frenzel, written communication) and by the EPA (EPA75) in 1975. Concentrations were 30, 14, 2.6, and 2.6 pCi/l, respectively.

It is apparent that there are marked temporal trends in mine water quality and quantity. Major factors responsible include changes in the dewatering rate accompanying shaft sinking versus actual ore production. Simultaneously, there are changes in the mineral quality and leaching rate of strata as the ore body is approached and then penetrated. Mining practices, oxidation of the ore body and possibly bacterial action may also assist in the solubilization of toxic stable and radioactive trace elements. Sample handling and analytical procedures can also markedly affect results. For example, if suspended solids are high and a sample is acidified prior to filtering, soluble radium, uranium, and other trace constituents typically

Table 3.39 Summary of average discharge and water quality data for underground uranium mines in the Colorado Plateau Region (Colorado, New Mexico, Utah) and a comparison with NPDES limits

Project	Discharge m ³ /min	Dissolved Radioactivity			Major and trace constituents, mg/ℓ							
		Total U, mg/ℓ	Ra-226, pCi/ℓ	Pb-210, pCi/ℓ	TSS	SO ₄	Zn	Ba	Cd	As	Mo	Se
<u>Utah</u> 1(a)	0.67	1.35	1.25		7.5							
<u>Colorado</u> 2	1.31	2.20	0.53		14.3	872	0.02	0.19	< 0.01	< 0.01	0.4	
3	0.06	0.25	10.00		144.9		0.065		0.003	0.055	0.054	
<u>New Mexico</u> 4	14.67	1.0	89 ^(b)	15	25.4	60.6		2.13		<0.005	< 0.01	0.03
5	3.79	0.67	23 ^(c)	33	2.6	213.7				0.011	0.24	0.008
6	1.89	0.02	14 ^(c)	15	51.5	744				0.005		0.004
7	0.95	0.18	0.1	0		1045				<0.005	0.05	0.002
8	0.18	4.2	1.9	9.7		67.2		0.88		<0.005	< 0.01	0.094
9(a)	0.82	1.9	4.7	16	1	675		0.17		0.011	0.45	0.407
10(a)	6.06	1.1	2.3	14	1.08	705				< 0.005	0.62	0.027
11(a)	0.216	2.6	4.3	14	2.2	837		0.66		0.012	0.79	0.036
Average	2.78	1.41	13.7	14.6	27.8	580		0.81		0.012	0.29	0.076
Standard Deviation	4.34	1.25	25.9	9.1	46.9	368		0.80		0.015	0.29	0.137

Table 3.39 (continued)

State	Summary of NPDES permit limits for daily average/daily maximum, mg/l except Ra-226, pCi/l								
	Dissolved Radium-226	Dissolved Uranium	Total Suspended Solids	Total Dissolved Solids	Zinc	Barium	Cadmium	Arsenic	Vanadium
<u>New Mexico</u>	3/10 10/30 Total Radium	2/4	50/150(day) ^(d) 20/30(month)		0.5/1.0				
<u>Utah</u>	3/10 and -/3	2/4 and -/2	20/30 ^(e)	NA/650 3500	0.5/1.0				
<u>Colorado</u>	3/10 and -/3	3/5 and 2/4	20/30	48990/ 122476 ^(f) kg/day	0.5/1.0	1/2 and -/1	0.05/0.1	1/2 and 0.5/1	5/10

(a) Average discharge rate per mine is shown. Two or more mines constitute the project.

(b) BaCl₂ treatment for radium removal faulty; repaired in late 1979.

(c) Values shown are for untreated water. BaCl₂ treatment now used.

(d) Applies to discharge associated with shaft construction.

(e) Maximum of 10 mg/l for 30-day period and 20 mg/l for 7-day period effective July 1, 1980.

(f) Receiving water standard.

Source: Chemical analyses from in-house studies (EPA75) and State of New Mexico (J. Dudley, Environmental Improvement Division, written communication). NPDES permit data from Regions VI, VIII (H. May, R. Walline, written communication). Other references include site-specific reports (EIS,ER) and company monitoring data.

will increase, as compared to samples that are filtered prior to acidification (Ka77). Therefore, development of "average" or "typical" trace element concentration data is questionable and may be erroneous without detailed knowledge of the many variables affecting the final results.

Despite the foregoing difficulties, available chemical data assembled in Table 3.39 provide much of the source term input data used in subsequent calculations. The reader should bear in mind that uranium concentrations are likely to be less than 3 mg/l simply because it is economically practical to use ion exchange recovery for concentrations greater than this level. Daily average radium-226 concentrations on the order of 3 pCi/l are specified in valid NPDES permits, and reliable data from USGS, EPA, and state sources reveal stream concentrations near the point of discharge to be on the order of 3 to 14 pCi/l in recent years. Therefore, the "average" radium-226 concentration of 13.7 pCi/l used in the subsequent modeling calculations is at least slightly conservative. Actual concentrations of stable elements (Zn, Ba, Cd, etc.) appear to be well below the NPDES limits, which were also developed from analysis of uranium mine effluent. Thus, it is presumed that the average values in Table 3.39 for these elements are reasonably correct. The variables of mine size, age, host rock, and water treatment (ion exchange, barium chloride, settling ponds) are reflected in the data. Water quality for mines examined in Utah and Colorado generally agrees with the New Mexico cases, with the exception of Project Number 3 mine, which is being dewatered and may, therefore, temporarily have excessive suspended solids. We recommend that the NPDES data for uranium mine discharges be evaluated and that additional compliance monitoring be conducted to confirm the quality of mine discharge. Such studies should focus on situations where mine water is being used for irrigation and stock watering.

Table 3.40 shows discharge and water quality characteristics for underground mines under construction and not yet producing ore. The first example involves water pumped from a deep mine shaft under construction. Considerable water is encountered above the ore body; water quality is good and representative of natural conditions; and suspended solids are high as a result of construction. The second case is similar except that flow is reduced, but radium and suspended solids concentrations are greatly elevated due to construction and possible ore body oxidation. The third case involves

Table 3.40 Water quality associated with underground mines in various stages of construction and operation

Project	Discharge m ³ /min	Total U mg/ℓ	Dissolved		Concentration, mg/ℓ				
			Ra-226 pCi/ℓ	Pb-210 pCi/ℓ	TSS	SO ₄	As	Mo	Se
<u>New Mexico</u>									
1. Underground mine shaft construction; dewatering	5.76	0.03	0.07	10	23.8	134	<0.005	0.01	0.003
2. Underground mine shaft construction; dewatering	1.73	<0.01	29	0	554	527	0.012	0.007	0.005
3. Underground mine; dewatering wells	1.43	0.08	0.2	0	1	144	<0.005	0.01	0.003
4. Underground mine recirculating leach solution from stopes (after ion exchange)	0	0.32	29	17	1.1	1060	<0.005	3.2	0.268

Source: J. Dudley, State of New Mexico, written communication, 1979.

dewatering wells used to dewater the ore body before mining. There is no oxidation and suspended solids are very low as is radium-226. Dissolved radium-226 in the ore body is on the order of 10 pCi/l or less in the natural state, but concentrations rise to 100 pCi/l or more after mining takes place, possibly due to oxidation and bacterial action in the workings (EPA75). Project Number 4, in the Ambrosia Lake district, is an inactive underground mine now used as a type of in situ leach facility. Mine water is recirculated through the workings. Leached uranium is selectively recovered using ion exchange. The process is a closed one, hence no effluent is involved. Water quality after uranium removal reflects the buildup in radium, lead-210, sulfate, molybdenum, and selenium.

3.4.3. Hydraulic and Water Quality Effects of Underground Mine Discharge

3.4.3.1 Runoff and Flooding in the Model Underground Mine Area

3.4.3.1.1 Study Approach

We chose to study an area of rather concentrated underground mining, similar to the Ambrosia Lake district of New Mexico. All of the mines in the district dewater to different degrees because the principal ore body is in the Westwater Canyon Member of the Morrison Formation, which is also a major aquifer. In the analysis, flows from some 14 active mines discharge to formerly dry washes and dissipate downstream by evaporation and, more importantly, infiltration. Suspended and dissolved constituents persist at the land surface and become available for resuspension and transport in surface floods with recurrence intervals of 2 to 25 years. Contaminated runoff from the sub-basin is then diluted in average annual flows of progressively larger streams and rivers of the region.

Similar to the analysis presented for surface mines in Wyoming, there is a three-basin hierarchy: sub-basin, basin, and regional basin (Fig. 3.16). These correspond to Arroyo del Puerto-San Mateo Creek, Rio San Jose and Rio Puerco, and the Rio Grande. Of these, the Rio Puerco is distinctly ephemeral. The Rio Puerco drains into the Rio Grande, which is perennial, due in large part to the heavily regulated flows and storage reservoirs. Because

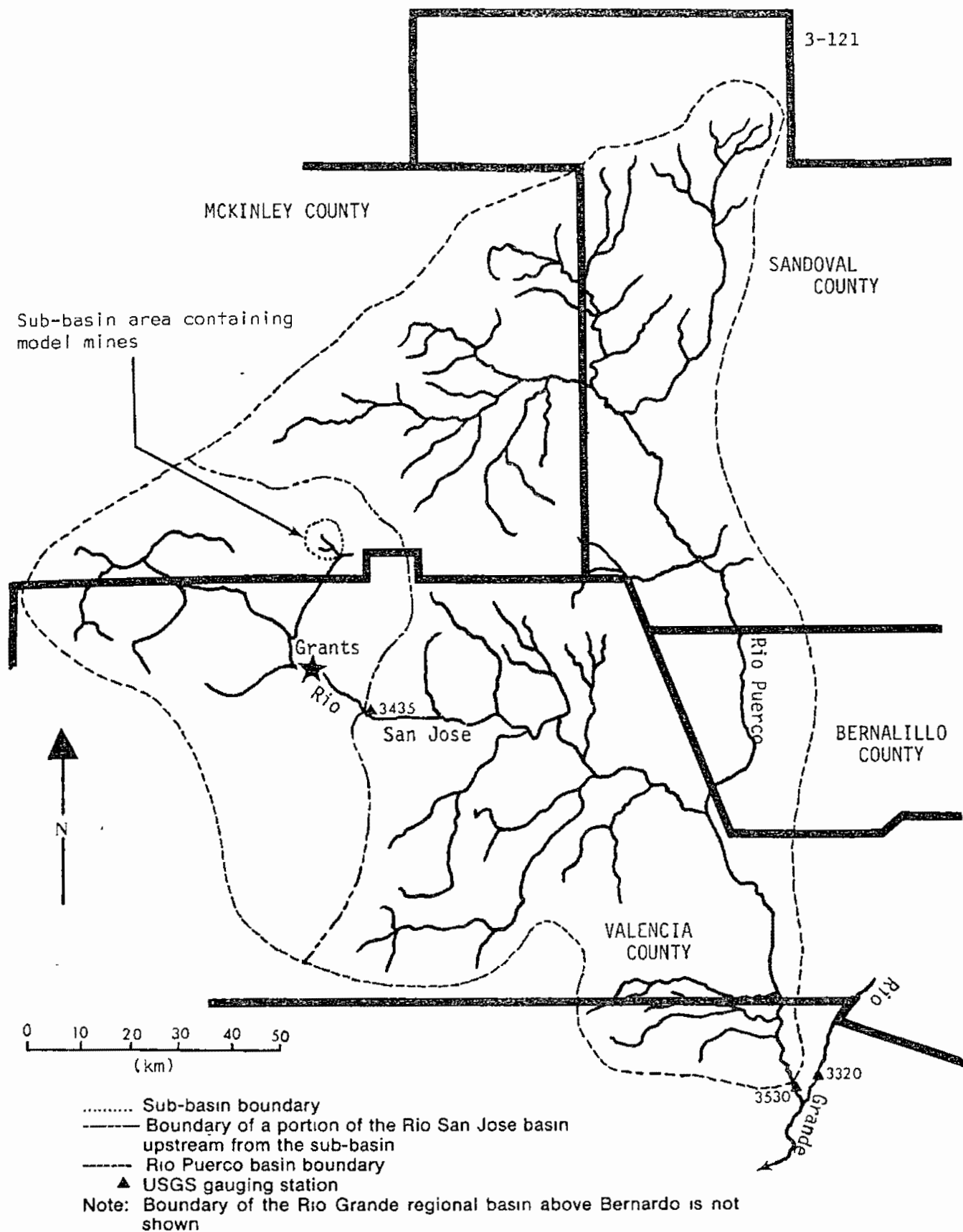


Figure 3 16 Sketch of sub-basin, basin, and regional basin showing orientation of principal drainage courses, areas of drainage, and location of mines in the New Mexico model area

the Rio Grande is the major regional river and the basis of extensive irrigation projects, it is included in the analysis. The mining area is well away from the Rio Grande Valley, and it is unlikely that noticeable changes in flow or water quality because of mining would occur.

Flow volumes for the sub-basin and open file USGS data (L. Beal, written communication, 1979) for flows in the basin and regional basin are used to transport and dilute contaminants originating in the mine effluent. It is initially assumed that all contaminants are available for transport by surface flow so as to deliberately create a worst-case situation. Section 3.4.3.2 reviews infiltration of water and solute for possible effects on groundwater.

We do not address the effects of seepage from settling ponds because such ponds are relatively small, tend to be self-sealing, and are well away from inhabited areas. Supposedly, settled solids from these ponds are removed and incorporated with uranium mill tailings. Limited field studies to determine whether such ponds cause groundwater contamination is warranted. In some instances, the ponds have synthetic liners, and leakage is expected to be minimal. The influence of mine dewatering (by wells, shafts, and pumping of mine workings) on groundwater quality or availability is not addressed primarily because of the lack of data. We strongly recommend further study of the hydraulic and groundwater quality effects of dewatering. This aspect of mining is coming under increased scrutiny by regulatory agencies at the State and Federal level because of the influence on water quality and availability.

In summary, our approach defines the quality and volume of mine water discharge; outlines hydrographic basins; and calculates flood flows for various return periods ranging from 2 to 25 years in the sub-basin. These flows are then diluted into the average annual flow in the basin and regional basin. The principal objective is to develop a rough estimate of contaminant loads resulting from mine discharge.

3.4.3.1.2 Description of Area

The Grants Mineral Belt of northwestern New Mexico is in the Navajo and Datil sections of the Colorado Plateau physiographic province (Fe31). Characteristic landforms in the study area include rugged mountains, broad, flat

valleys, mesas, cuestras, rock terraces, steep escarpments, canyons, lava flows, volcanic cones, buttes, and arroyos (Ki67; Co68). Elevations in the area range from 1,980 m at Grants to an average of 2,160 m near Ambrosia Lake. Just north of Grants is Mount Taylor, the highest point in the region. It rises from Mesa Chivato to an elevation of 3,471 m (Co68).

The study area has a mild, semiarid, continental climate. Precipitation averages 25.4 cm/year, and there is abundant sunshine, low relative humidity, and a comparatively large annual and diurnal temperature range. Average annual precipitation at Gallup, Bluewater, and Laguna is 27.12, 24.55, and 22.31 cm, respectively. In the higher elevations, the average is 51 cm or more because of thunderstorms in July, August, and September and snow accumulations in the winter months (Co68, Go61, Jo63). Only thunderstorms are significant in the lowlands. Heavy summer thunderstorms (40 to 70 in number) of high intensity and local extent can result in 5 cm of rain with local, damaging flash floods.

The watersheds of the Rio San Jose and Rio Puerco encompass 19,037 km². Most of the larger communities in the basin are located in the floodplain of the Rio Grande and principal tributaries. Extensive irrigation with surface water occurs in the watersheds of the Rio San Jose, Rio Puerco, and Rio Grande. In the sub-basin, there was no perennial flow before mining and, thus no irrigation, but increasing use is being made of the mine discharge, which is regarded as an asset in a water-short area. Subsequent sections summarize the surface water quantity at some of the principal gauging stations in the Middle Rio Grande Basin and the irrigated areas below these stations. Groundwater is used for essentially all public water supplies as the temperature, quality, and year-round availability are assured. Numerous wells scattered across the landscape, particularly in the stream valleys, are used for stock water and, to a lesser extent, for potable use on the scattered ranches and Indian settlements.

Under completely natural conditions, streams in the study area were distinctly ephemeral, and many of the smaller ones did not experience flow for periods of several years. The Rio Grande experiences peak flows in the April-June period when snowmelt and precipitation cause gradual rises to moderate discharge levels involving large volumes of flow and long durations. Peak discharge rates (volume per time) occur in the summer flash floods.

Construction of dams and conveyance channels to eliminate flooding problems has been extensive. In the tributaries such as the Rio San Jose and upper reaches of the Rio Puerco, there is considerable streamflow regulation to minimize flood damage and maximize use of available water for irrigation.

Conditions in the Ambrosia Lake district with respect to the type of mining operations and discharge of effluent to ephemeral streams are duplicated elsewhere in the Grants Mineral Belt. In the Churchrock district, two mines discharge to the Rio Puerco at rates of 4.7 to 15 m³/min. Most of the 4.7 m³/min discharge from one mine is now used in a nearby mill. At Mariano Lake, located between Ambrosia Lake and Churchrock, and at the Marquez and Rio Puerco mines east of Ambrosia Lake, mines are expected to discharge 0.8 to 4.5 m³/min to various ephemeral streams. Another large mine will soon discharge up to 5.3 m³/min northward into the San Juan River Basin. In the mid 1980's, construction is expected to begin on five large underground mining projects that will have a combined discharge on the order of 71 m³/min. Most discharge will be into the San Juan Basin, reflecting the trend of mines becoming deeper and requiring more dewatering as the mining center moves from the south flank of the San Juan Basin into more interior portions.

3.4.3.1.3 Estimate of Sub-basin Flood Flow

Since we use a dilution-model, emphasis is on flow volume rather than peak discharge rate in the sub-basin, basin, and regional basin hydrographic units. Gaging records from the U.S. Geological Survey WATSTORE system (L. Beal, written communication, 1979) provide average discharge rates for runoff events with various return periods and durations. The latter specify the time, in days, and the associated flow rate that will be equaled or exceeded. Flows for arbitrary periods of time ranging from 1 to 183 days are specified. Probability can be stated in terms of N-year recurrence interval. By combining discharge rate (volume per time) and time (partial duration), flow volume can be calculated.

In the ungaged sub-basin, runoff volumes associated with events having return periods of 2, 5, 10, and 25 years were calculated from regression equations developed by the USGS (Bo70). The equations were generated from multiple regression of discharge records from gaged basins against various basin characteristics. These are area (A), precipitation (P_a), longitude

at the center of the sub-basin (Lo), soils infiltration index (Si), and mean basin elevation (E_m). Through use of appropriate constants and coefficients (Bo70), flow volumes can be calculated for 1-day and 7-day events with return periods of 2, 5, 10, and 25 years. For the sub-basin, the basic equation has the following form:

$$FV = a A^{b_1} P_a^{b_2} Lo^{b_3} S_i^{b_4} E_m^{b_5} \quad (3.10)$$

where $A = 95 \text{ mi}^2$
 $P_a = 2.9 \text{ inches}$
 $Lo = 7.85 \text{ (longitude in decimal degrees minus 100)}$
 $S_i = 8.5$
 $E_m = 7.0 \text{ thousand feet}$

Table 3.41 contains the regression coefficients and total flow volume data. Short-term, 1-day and 7-day, events were of main interest because these would be expected to provide greater flushing of contaminants stored at or near the water-substrate interface in the streams receiving mine discharge.

The extent to which mine discharge transforms existing ephemeral streams into perennial ones is evaluated with a crude seepage and evaporation model (see Appendix H). The basic equations and approach are patterned after a similar analysis in the Generic Environmental Impact Statement on Uranium Milling (NRC79b).

Figure 3.16 shows the relationship of the sub-basin, basin, and regional basin boundaries and the principal drainage courses and gaging stations. The confluence of the Rio Puerco and Rio San Jose is shown approximately 55 km closer to the Rio Grande than is actually the case in order to simplify flow routing and to reduce the number of dilution calculations. Table 3.42 summarizes the key characteristics of these basins in terms of catchment area, discharge, and irrigated farmlands downstream from points where mine discharge might be tributary to the streams. Mine discharge occurs in the sub-basin which in turn discharges to the Rio San Jose and then to the Rio Puerco. No mine discharge and no significant runoff are associated with that portion of the basin tributary to San Mateo Creek between the Rio San Jose and the sub-basin. For modeling, flooding within and runoff from the sub-

Table 3.41 Total flow volume for sub-basin floods of 1- and 7-day durations and return periods of 2, 5, 10, and 25 years

Flood Volume	Regression Coefficients						Volume (m ³)
	a	b ₁	b ₉	b ₁₄	b ₁₅	b ₄	
FV 1,2 ^(a)	1.08×10^{-4}	0.931	1.83	-1.43	4.09	-	2.16×10^4
FV 1,5	1.27×10^{-3}	0.941	1.40	-1.89	4.07	-	6.23×10^4
FV 1,10	5.07×10^{-3}	0.953	1.17	-2.18	4.02	-	1.02×10^5
FV 1,25	2.39×10^{-2}	0.972	0.929	-2.51	3.95	-	1.76×10^5
FV 7,2	8.60×10^{-7}	0.965	2.36	-1.61	4.22	1.50	5.95×10^3
FV 7,5	2.99×10^{-4}	0.904	2.55	-2.09	3.53	-	8.79×10^3
FV 7,10	8.97×10^{-4}	0.910	2.37	-2.39	3.61	-	1.43×10^4
FV 7,25	3.06×10^{-3}	0.922	2.17	-2.76	3.68	-	2.26×10^4

(a) FV 1,2 indicates a flood of 1-day duration and a return period of 2 years.

Table 3.42 Summary of area, discharge, and irrigated acreage for the sub-basin, basin, and regional basin hydrographic units in New Mexico

	USGS Station Number	Area (km ²)	Period of Record Yrs.	Number of km ² Under Irrigation Below Station	Average m ³ /min Discharge (for Period of Record)	Average Annual Discharge (m ³) for Period of Record
<u>Sub-basin</u>	---	246	---	---	---	---
<u>Basins</u>						
Rio San Jose near Grants	3435	5957 (2927 non-contributing)	42	2.43+	11.09	5.83 x 10 ⁶
Rio Puerco near Bernardo	3530	19037 (2927+ non-contributing)	38	---	81.05	4.26 x 10 ⁷
<u>Regional Basin</u>						
Rio Grande at Bernardo	3320	49810 (7610 non-contributing)	41	N/C ^(a)	1649.35	86.69 x 10 ⁷

(a) N/C = Not Calculated.

basin is, in effect, routed without change in flow and quality and allowed to enter the Rio San Jose. Flow from the San Jose is further diluted in the Rio Puerco, then diluted again in the Rio Grande. In actuality, flow from the Ambrosia Lake district rarely, if ever, enters the Rio San Jose because flood volumes are small and infiltration losses are large. This departure from true conditions is justified within the context of the modeling approach used. Basically, the model draws from a specific area but does not attempt to closely duplicate its conditions. If a specific area were exactly represented, the model would still be incorrect to varying degrees for other areas, and the generic value of the assessment would depreciate.

Of special interest is the effect of contaminated flows on irrigation projects present on the Rio San Jose and Rio Grande. An extensive system of dams and conveyance channels regulates flow in the Rio Grande, and partial duration flow data are unavailable. Instead, the average annual flow volume is used to provide the final dilution estimate. For the sub-basin in which the mines are located, flood volumes are calculated using the USGS regression equations (Bo70). The maximum return period for which flows are calculated is 50 years. The remainder of this section first considers the flow or hydraulic aspects of the surface water pathway. Finally, several factors concerning the quality of runoff water are mentioned to balance conservatism and realism in the pathway analysis and, subsequently, in the health effects modeling to follow. The emphasis here is on surface water impacts, and we assume maximum transport for this pathway. The influence of infiltrating mine water is discussed in Section 3.4.3.2.

All of the streams, except the Rio Grande and certain reaches of the Rio San Jose are distinctly ephemeral under natural conditions. In the sub-basin, there is perennial flow because of mine discharge. In Fig. 3.17 are the average monthly and annual discharges for the Rio San Jose and the Rio Puerco in comparison to cumulative annual flow from 14 mines, each discharging at $2 \text{ m}^3/\text{min}$. The monthly data reveal pronounced seasonal variations approaching 1 to 2 orders of magnitude. The streams do not show the same seasonal variations, further attesting to varied patterns of runoff, irrigation diversion, and control features such as impoundments and conveyance/irrigation channels. Figure 3.18 shows the percentage of each month during

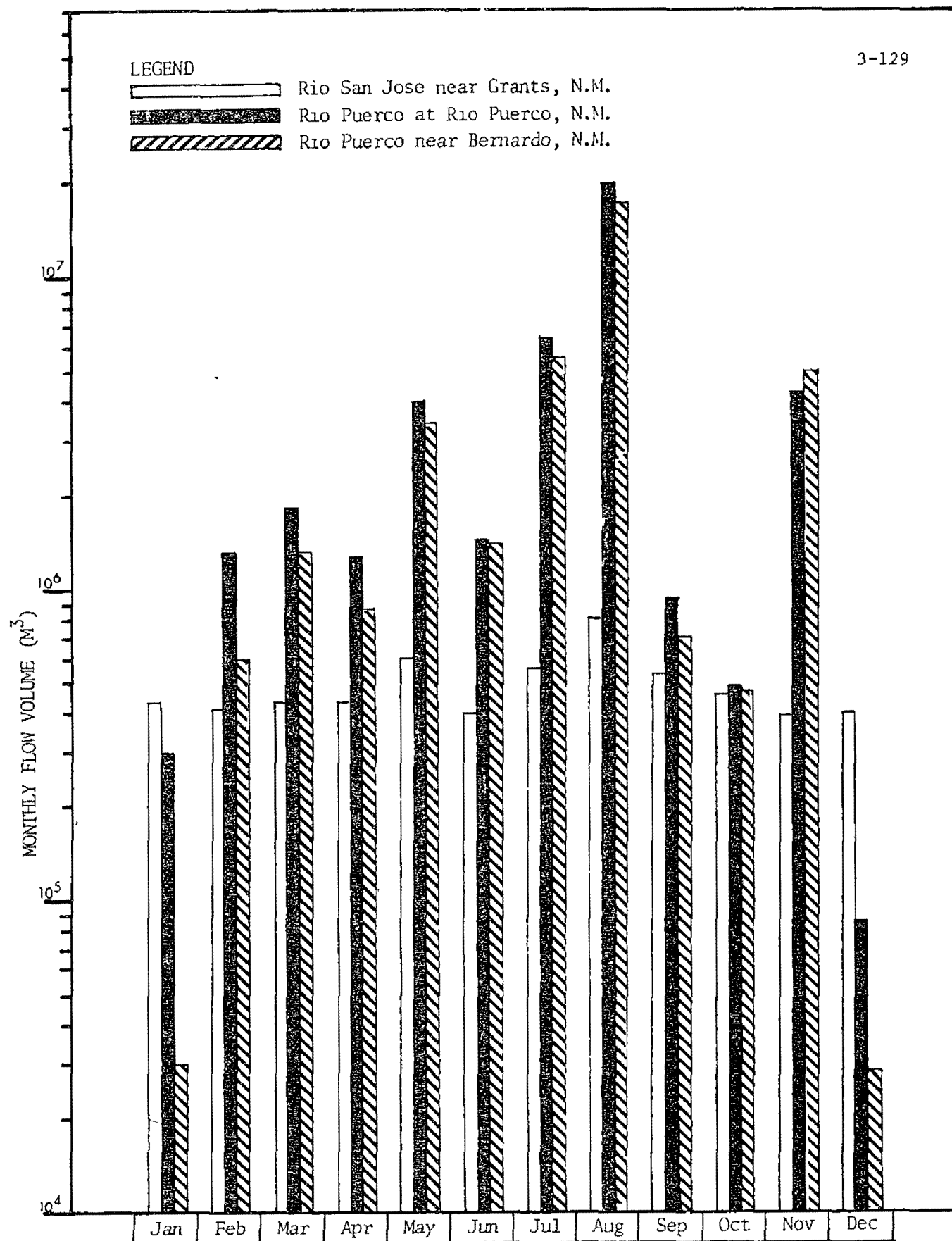


Figure 3 17 Average monthly flows for the period of record for the Rio San Jose and the Rio Puerco in New Mexico (Summarized from flow records provided by L. Beal, U S Geological Survey (Bureau))

which there is no flow in the Rio San Jose and Rio Puerco. The average period of annual or monthly no flow is as follows:

Rio San Jose near Grants:	0 Percent
Rio Puerco at Rio Puerco:	45 Percent
Rio Puerco at Bernardo :	71 Percent

It is also assumed that flow from the sub-basin reaches the first major stream, the Rio San Jose, with no change in flow or quality. Runoff is minimal in the lower reaches of San Mateo Creek because of internal drainage and considerable infiltration. Historical evidence indicates that only rarely, if ever, would flood runoff from Ambrosia Lake enter the Rio San Jose. In the interests of conservatism, total flow laden with contaminants is transported to the Rio San Jose. Dilution first occurs within the sub-basin and then, successively, in the Rio San Jose, Rio Puerco, and Rio Grande. The latter is the regional basin.

There is an infinite number of combinations of flood volumes and dilution volumes for the sub-basin, basin, and regional basin streams. Use of average annual discharge volumes in the receiving streams simplifies what would otherwise be a burdensome, confusing series of calculations. Flushing action from the sub-basin is handled on a probabilistic basis in terms of flow duration and return period. Concentration values are based on 14 mines, a loading period of two years, and flow and water quality data shown in Table 3.39. When, for example, 5-year or 10-year events are considered, it is conservatively assumed that events with shorter return periods do not occur. The accretion period remains constant (2 years), and only the return period and duration are varied, resulting in varying flow volumes. It is conceivable that contaminants could concentrate for 3, 4, or 5 years and then be flushed by a 2-year event, but this was not evaluated.

Minimum and maximum return periods for floods from the sub-basin were set at 2 and 25 years, respectively, for several reasons. The 2-year event, i.e., runoff volume over a duration of 1 day or 7 days and occurring on the average of every 2 years, is expected to occur rather frequently over the life of the mines (17 years). The intermediate-sized storms with return periods of 5 or 10 years would result in considerable contaminant transport,

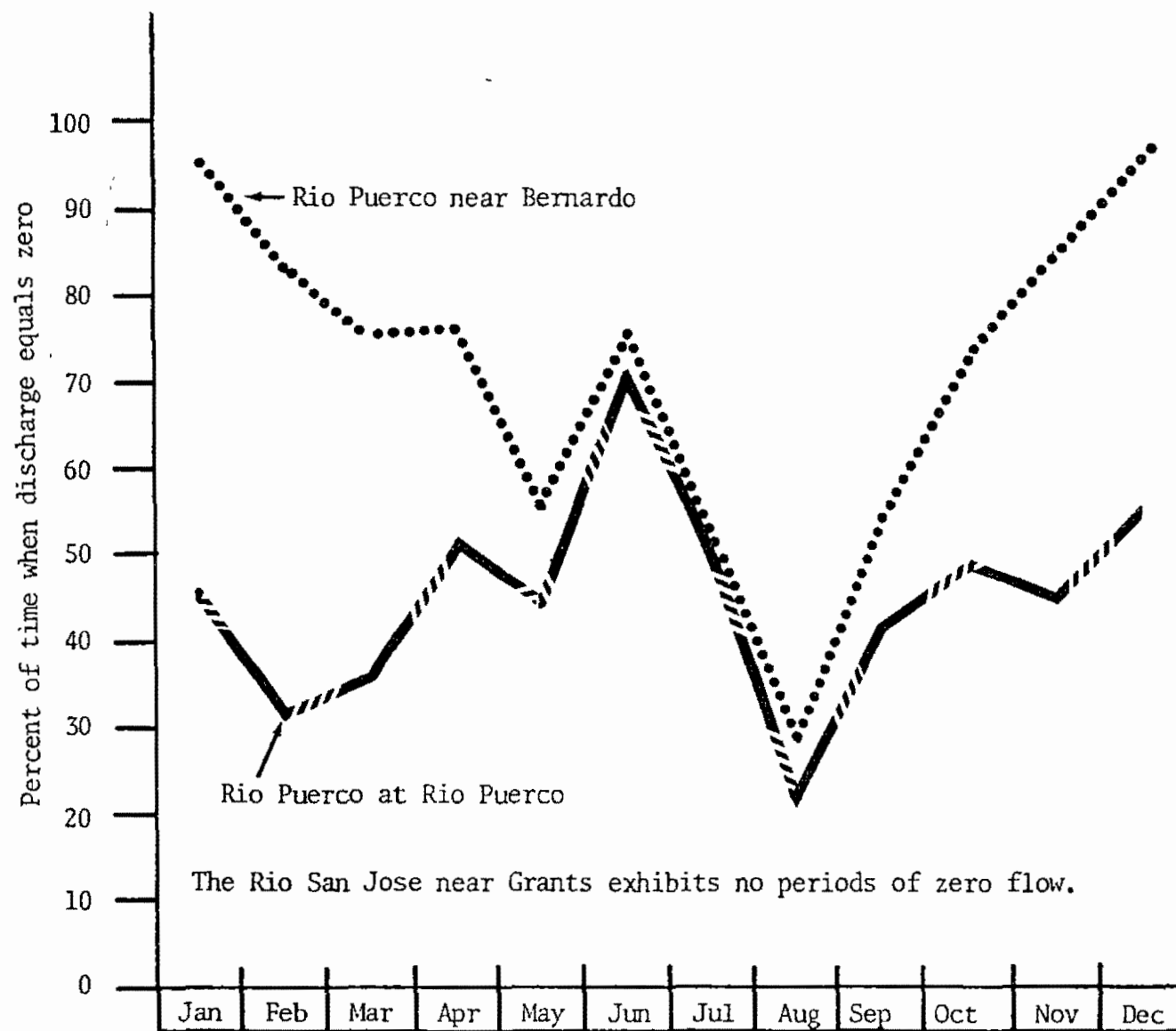


Figure 3.18 Periods of no flow in the Rio San Jose and Rio Puerco (Summarized from flow records provided by L. Beal, U.S. Geological Survey, Albuquerque)

but concentrations would be low owing to dilution and to annual or semiannual scouring provided by smaller floods. The 25-year event is a practical maximum expected to occur during the lifetime of the mining district. Still larger floods, with return periods of 50 or 100 years, can be calculated but are less important because of their infrequent occurrence. Figures 3.19 and 3.20 show calculated flow volumes from the sub-basin for 1-day and 7-day durations and return periods of 2 years to 50 years. The extreme range in flow volume is from $2.16 \times 10^4 \text{ m}^3$ to $2.55 \times 10^5 \text{ m}^3$.

Figures 3.19 and 3.20 show flow values in the Rio San Jose and Rio Puerco for 1-day and 7-day durations and return periods of 1 to 100 years. For the Rio San Jose, 1-day volumes range from $1.24 \times 10^4 \text{ m}^3$ to $1.68 \times 10^6 \text{ m}^3$. The mean annual discharge rate in the Rio San Jose is $11.09 \text{ m}^3/\text{min}$. Flow from the Rio San Jose enters the Rio Puerco where corresponding flows (1-day duration) range from 0.6×10^6 to $2.15 \times 10^7 \text{ m}^3$ at the point of inflow to the Rio Grande. Average daily discharge in the Rio Grande seasonally ranges from $8.87 \times 10^5 \text{ m}^3$ to $59.5 \times 10^5 \text{ m}^3$. Average annual flows rather than peak 1-day or 7-day flows were used in the subsequent calculations.

The maximum probability for peak runoff from the sub-basin and resulting contaminant transport is in the summer months, at which time the Rio Puerco has no flow about 22 to 75 percent of the time. Flow in the Rio San Jose and Rio Puerco from June through September ranges from 3.96×10^5 to $1.97 \times 10^7 \text{ m}^3$ per month for the period of record (Fig. 3.17).

3.4.3.1.4 Prediction of Sub-basin Water Quality

Table 3.43 outlines dilutions based on the foregoing discussion of flow patterns and discharges and considering only the 1-day sub-basin flood event with a 2-year recurrence interval. The dilution constant is the ratio of concentration in the receiving water to that in the contaminated (relatively) inflow. It is more commonly expressed as the dilution factor, which is the reciprocal. Thus, in the case of the sub-basin flood flow entering the mean annual flow of the Rio San Jose, there is a 271:1 dilution (Table 3.43).

With development of the foregoing (mine water) source term and surface water pathway, the remaining discussion emphasizes contaminant concentrations in surface water. This, in turn, serves as input data to health effects modeling for the water pathway. Chemical concentrations in the Rio

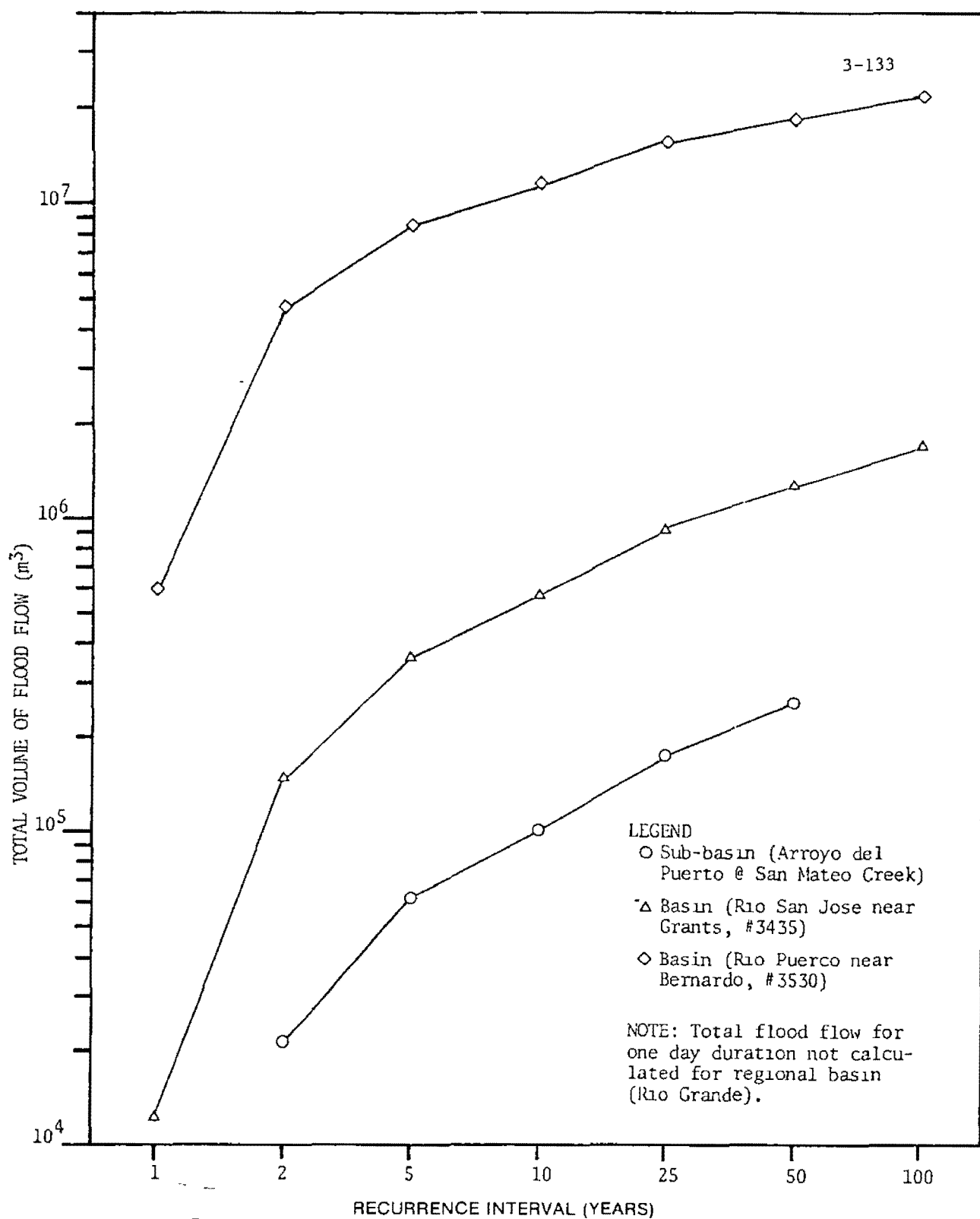


Figure 3 19 Total flow volumes in one-day periods for floods of various recurrence intervals in the sub-basin and basins in New Mexico (Summarized from flow records provided by L. Beal, U S Geological Survey, Albuquerque)

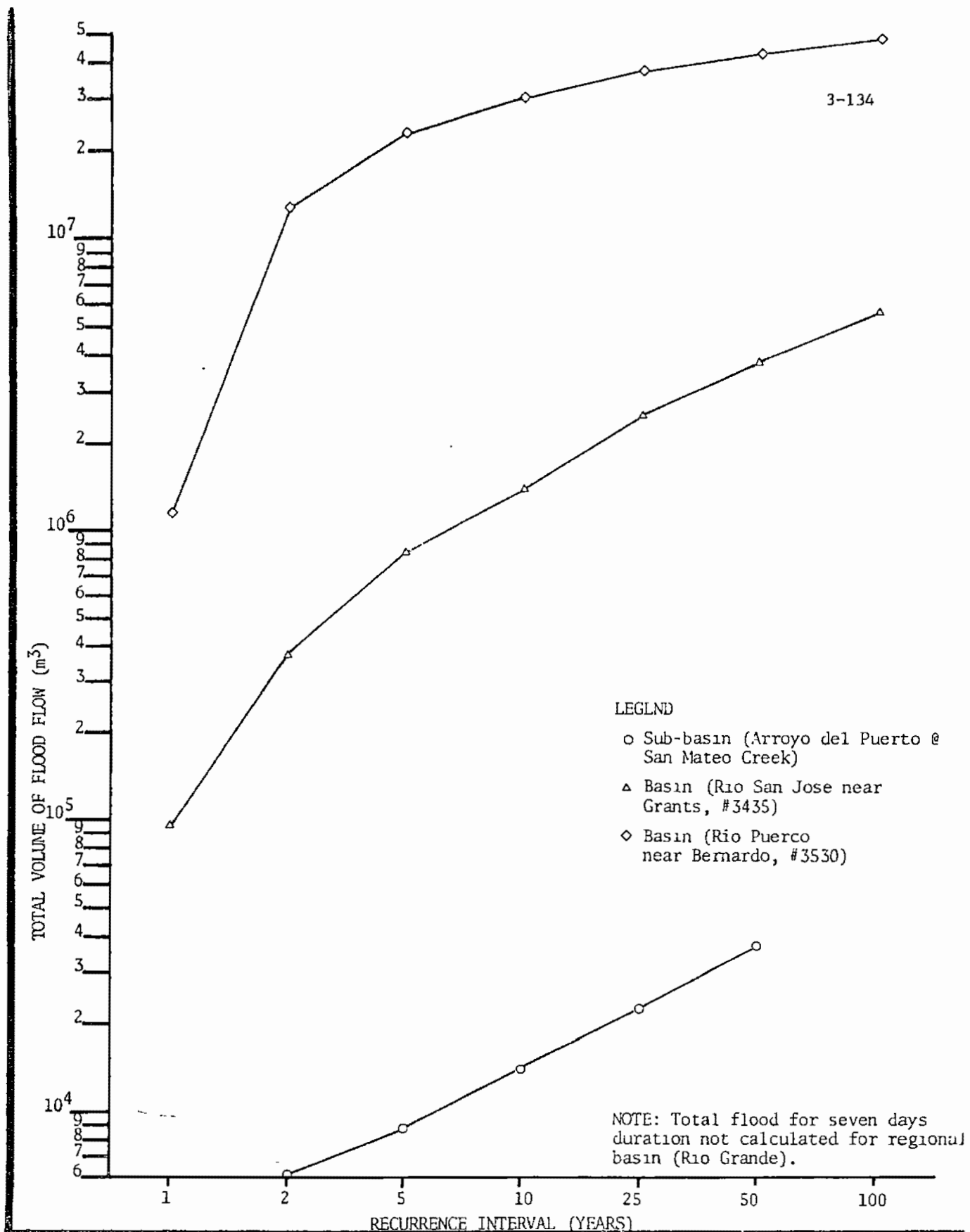


Figure 3-20 Total flow volumes in seven-day periods for floods of various recurrence intervals in the sub-basin and basins in New Mexico / Summarized

Table 3.43 Dilution factors for the Rio San Jose, Rio Puerco, and Rio Grande for 1-day flood flows with a 2-year recurrence interval

Hydrographic Basins	Flow Ratio (m ³ /m ³)	Dilution Constant	Dilution Factor
Rio San Jose near Grants ^(a)	$\frac{2.16 \times 10^4}{5.83 \times 10^6 + 2.16 \times 10^4}$	= 0.0037	271
Rio Puerco ^(b)	$\frac{2.16 \times 10^4}{4.26 \times 10^7 + 2.16 \times 10^4}$	= 0.00051	1973
Rio Grande near Bernardo ^(c)	$\frac{2.16 \times 10^4}{86.69 \times 10^7 + 2.16 \times 10^4}$	= 0.000025	40135

(a) Calculated using mean annual flow in the Rio San Jose (near Grants, NM) station:

$$\text{Dilution} = \frac{\text{Sub-basin flood flow}}{\text{Rio San Jose flow} + \text{Sub-basin flood flow}} .$$

(b) Assumes Rio San Jose enters the Rio Puerco at Bernardo:

$$\text{Dilution} = \frac{\text{Sub-basin flood flow}}{\text{Rio Puerco flow (includes Rio San Jose flow)} + \text{Sub-basin flood flow}} .$$

(c) Dilution =
$$\frac{\text{Sub-basin flood flow}}{\text{Rio Puerco flow} + \text{Rio Grande flow (at Bernardo)} + \text{Sub-basin flood flow}} .$$

San Jose, Rio Puerco (at Bernardo), and in the Rio Grande (near Bernardo) are shown in Table 3.44 along with 1-day and 7-day flood flow volumes from the sub-basin for return periods of 2, 5, 10, and 25 years. These flood volumes are diluted into the mean annual flow of the Rio San Jose (near Grants), Rio Puerco (at Bernardo), and Rio Grande (near Bernardo). The principal reason for using mean annual flow is that the radiation dose and health effects model (Section 6.0) stresses estimating average annual dose to the population over the duration of mining activity.

For example, the 1-day duration flood flow (with a 2-year return period) contains 1920 mg/ℓ uranium, which decreases to 7.09 mg/ℓ in the Rio San Jose and 0.973 mg/ℓ in the Rio Puerco. Because of the short duration of most floods in the sub-basin, there is little difference in flow volume and, thus, dilution between the 1-day and 7-day events. With progressive dilution downstream, the difference in size between sub-basin floods of varying durations and return periods becomes insignificant relative to the mean annual flow volumes of the basin and regional basin streams. As a result, concentrations tend to reach a minimum and remain unchanged at this degree of accuracy.

As in the case of the Wyoming surface-mine scenario, we assume that most contaminants in the mine water collect on or near the land surface and are available for transport. This assumption is open to question, but field data are scarce to support contentions as to the fraction of contaminant load that becomes unavailable. For example, extensive field studies along the Animas, San Miguel, and Dolores Rivers in Colorado concluded that "...once radium becomes a part of a stream's environment, it constitutes a relatively long-term and continuous source of water and aquatic biota contamination" (Si66). However, cessation of uranium mill discharges to the Colorado River tributaries effectively negated this source, which is now believed to be buried behind the Lake Powell and Lake Mead impoundments. Similarly, dissolved radium reverts to background levels of several picocuries per liter in natural streams receiving mine water in Colorado and New Mexico. Although it is likely that flood waters resuspend precipitates and sediments with sorbed radium, laboratory experiments (Sh64; Ha68) indicate that only minor re-solution takes place. This phenomenon is supported by recent surface water data collected in the Grants Mineral Belt of New Mexico (Ku79). Therefore, concentrations of dissolved radium in flood water are

Table 3.44 Annual contaminant loading from 14 uranium mines and resulting concentrations in sub-basin floods and in the average annual flow of the Rio San Jose, Rio Puerco, and Rio Grande

Contaminant concentration in mine effluent (mg/l except as noted)	Mass available for transport (kg/yr except as noted) (a)	1- and 7-day flood flow volumes (m ³) and contaminant concentrations associated with return periods of 2 to 25 years (b)							
		1-Day				7-Day			
		V ₂ =2.16x10 ⁴ C ₂	V ₅ =6.23x10 ⁴ C ₅	V ₁₀ =1.02x10 ⁵ C ₁₀	V ₂₅ =1.76x10 ⁵ C ₂₅	V ₂ =5947 C ₂	V ₅ =8794 C ₅	V ₁₀ =1.43x10 ⁴ C ₁₀	V ₂₅ =2.26x10 ⁴ C ₂₅
Total Uranium 1.41	1480	1920 7.09 0.973 0.0456	665 7.03 0.971 0.0456	406 6.98 0.970 0.0455	235 6.89 0.967 0.0455	6970 7.10 0.973 0.0456	4710 7.09 0.972 0.0455	2900 7.10 0.973 0.0456	1830 7.07 0.970 0.0456
Radium-226 13.7 pCi/l	0.00144 Ci/yr	1870 6.9 0.95 0.044	647 6.8 0.95 0.044	395 6.8 0.94 0.044	229 6.7 0.94 0.044	6780 6.9 0.95 0.044	4580 6.9 0.95 0.044	2820 6.9 0.95 0.044	1780 6.9 0.94 0.044
Lead-210 14.6 pCi/l	0.00153 Ci/yr	19800 73.1 10.0 0.470	6880 72.7 10.0 0.471	4200 72.2 10.0 0.471	2430 71.2 10.0 0.470	72000 73.4 10.0 0.471	48700 73.3 10.1 0.471	30000 73.4 10.1 0.472	19000 73.4 10.1 0.472
Cadmium 0.007	7	9 0.03 0.004 0.0002	3 0.03 0.004 0.0002	2 0.03 0.005 0.0002	1 0.03 0.004 0.0002	30 0.03 0.004 0.0002	20 0.03 0.004 0.0002	10 0.02 0.003 0.0002	9 0.03 0.005 0.0002
Arsenic 0.012	13	17 0.063 0.0086 0.00040	5.8 0.061 0.0085 0.00040	3.6 0.062 0.0086 0.00040	2.1 0.062 0.0086 0.00041	61 0.062 0.0085 0.00040	41 0.062 0.0085 0.00040	25 0.061 0.0084 0.00039	16 0.062 0.0085 0.00040
Selenium 0.076	80	100 0.38 0.053 0.0026	36 0.38 0.052 0.0026	22 0.38 0.052 0.0026	13 0.37 0.052 0.0026	376 0.38 0.053 0.0026	254 0.38 0.053 0.0026	156 0.38 0.053 0.0026	99 0.38 0.053 0.0026

Table 3.44 (continued)

Contaminant concentration in mine effluent (mg/l except as noted)	Mass available for transport (kg/yr except as noted) ^(a)	1- and 7-day flood flow volumes (m ³) and contaminant concentrations associated with return periods of 2 to 25 years ^(b)							
		1-Day				7-Day			
		$V_2=2.16 \times 10^4$ C_2	$V_5=6.23 \times 10^4$ C_5	$V_{10}=1.02 \times 10^5$ C_{10}	$V_{25}=1.76 \times 10^5$ C_{25}	$V_2=5947$ C_2	$V_5=8794$ C_5	$V_{10}=1.43 \times 10^4$ C_{10}	$V_{25}=2.26 \times 10^4$ C_{25}
Molybdenum 0.29	300	390 1.4 0.20 0.0093	130 1.4 0.19 0.0089	82 1.4 0.20 0.0092	48 1.4 0.20 0.0091	1400 1.4 0.20 0.0092	960 1.4 0.20 0.0093	590 1.4 0.20 0.0091	370 1.4 0.20 0.0092
Barium 0.81	850	1100 4.1 0.56 0.026	380 4.0 0.55 0.026	230 4.0 0.55 0.026	140 4.1 0.58 0.027	4000 4.1 0.56 0.026	2700 4.1 0.56 0.026	1700 4.2 0.57 0.027	1100 4.2 0.58 0.027
Zinc 0.043	45	58 0.21 0.029 0.0014	20 0.21 0.029 0.0014	12 0.21 0.029 0.0014	7.2 0.21 0.030 0.0014	210 0.21 0.029 0.0014	140 0.21 0.029 0.0014	88 0.22 0.030 0.0014	56 0.22 0.030 0.0014
Sulfate 580	1.22×10^5	1.58×10^5 584 80 3.8	5.48×10^4 580 80 3.8	3.35×10^4 574 80 3.7	1.94×10^4 568 80 3.8	5.74×10^5 586 80 3.8	3.88×10^5 584 80 3.8	2.38×10^5 582 80 3.7	1.51×10^5 584 80 3.8
Total Suspended Solids 27.8	29000	38000 140 19 0.90	13000 140 19 0.89	8000 140 19 0.90	4600 130 19 0.89	140000 140 20 0.92	92000 140 19 0.89	57000 140 19 0.90	36000 140 19 0.89

(a) Mass values shown are on an annual, per-mine basis.

(b) V_r and C_r refer respectively to flood volume, in cubic meters, and concentration in runoff for an r-year flood. Concentrations are in mg/l, except for radium-226 and lead-210, which are in pCi/l. Concentrations shown are from accretion or loading in the sub-basin for 2, 5, 10, 25 years, yielding the first value shown in each set. The next three values below this initial value represent, in downward order, concentrations in the flood flow as diluted by the mean annual flow in 1) the Rio San Jose near Grants (5.83×10^6 m³), 2) the Rio Puerco at Bernardo (4.26×10^7 m³), and 3) the Rio Grande near Bernardo (86.69×10^7 m³).

Note.--Assumptions: Mines discharge continuously at a rate of 2.0 m³/min. Concentrations are the average of those shown in Table 3.39. Except for radium and sulfate, all suspended and dissolved contaminants remain in or on the stream sediments and are mobilized by flood flow. Twenty percent of the sulfate and 10 percent of the radium are available for resolution.

arbitrarily set at 0.00144 Ci/yr or 10 percent of the annual loading from the model mine.

Sulfate is also considered an important exception in the total "transport" concept. Because sulfate can be a highly mobile anion, it is assumed that 80 percent of the load enters the shallow groundwater reservoirs and 20 percent is available for solubilization and chemical transport in surface flows. No distinct pattern of groundwater contamination from mine water, per se, was documented in an earlier Grants Mineral Belt survey (EPA75), but recent data from the State indicate groundwater deterioration as a result of mine drainage (J. Dudley, New Mexico Environmental Improvement Division, oral communication, 1979). It is likely that considerable fractionation of other stable and radioactive trace elements occurs, but field data specific to the uranium mining regions are quite scarce, with the exception of Texas (He79), where only stable elements were studied. Because of our imperfect, non-predictive understanding of trace element transport in aqueous systems, our analysis assumes total transport for most constituents in lieu of numerous, equally unfounded assumptions for resuspension factors, fractionation, etc. Floods of 1-day and 7-day duration and return periods of 2, 5, 10, and 25 years are arbitrarily selected as providing the necessary flushing action associated with intense, short-term runoff events. It is likely that storms of shorter (less than 1-day) duration and possibly greater discharge rate also transport contaminants. The flow volume and thus the dilution cannot be estimated for these events.

Calculated water quality in basin and regional basin streams is shown in Table 3.45 along with established and suggested standards for selected contaminants. For uranium, concentrations in the basin exceed the suggested limits based on chemical toxicity and radiotoxicity. Radium-226/228 exceeds the standard in the basin but is well below the standard for the regional basin. The same is true for sulfate, cadmium, arsenic, barium, and selenium. Zinc is the only contaminant consistently below the potable and irrigation water standards. As in the case of the surface mine scenario for Wyoming, uranium is apparently well above suggested limits and warrants further study, as do the stable toxic elements in the basin area(s) closest to the mining centers.

With the exception of radium-226 and sulfate, the concentrations of radionuclides and other parameters shown in Tables 3.44 and 3.45 reflect no

Table 3.45 Comparison of potable and irrigation water standards and surface water quality affected by underground mine drainage

Parameter	Range of contaminant concentrations in flood flow affected by mine discharge (a)				Potable water standards (mg/l) (b)			Irrigation (c) Recommendations for maximum concentration for continuous use on all soils (mg/l)
	Basin Min.	Max.	Regional Min.	Basin Max.	Maximum Permissible Concentration	Recommended Concentration	Limiting Concentration	
Total U	6.9	7.1	0.045	0.046	0.015/3.5/0.2 ^(d)			---
Ra-226 + 228	6.7	6.9	0.044	0.044	----	5 pCi/l	---	5 pCi/l
TSS	130	140	0.89	0.92	---	---	---	---
Sulfate	574	584	3.7	3.8	---	250	---	200
Zn	0.21	0.22	0.0014	0.0014	---	5.0	---	2.0
Cd	0.03	0.03	0.0002	0.0002	0.01	---	---	0.010
As	0.061	0.063	0.00039	0.00041	0.05	0.01	---	0.10
Ba	4.0	4.2	0.026	0.027	1	---	---	---
Se	0.37	0.38	0.0026	0.0026	0.01	---	---	0.02

(a) Concentrations in milligrams per liter, except Ra-226 -228 which are in picocuries per liter. Data shown apply to the Basin (Rio San Jose near Grants) and Regional Basin (Rio Grande near Bernardo) streams (Table 3.44).

(b) Sources: U.S. Environmental Protection Agency (EPA76) and, in the case of uranium, suggested guidance from the National Academy of Sciences (NAS79) to the USEPA and from USEPA, (Office of Drinking Water) to the State of Colorado (La79).

(c) Source: (NAS72).

(d) 0.015 mg/l: Suggested maximum daily limit based on radiotoxicity for potable water consumed at a rate of 2 liters per day on a continuous basis
 3.5 mg/l: Suggested maximum daily limit based on chemical toxicity and intake of 2 liters in any one day
 0.21 mg/l: Suggested maximum daily limit based on chemical toxicity and intake of 2 liters per day for 7 days

reductions for ion exchange, precipitation, or sorption. Rather, a simple dilution model is used in which the mass loading from mine discharge is calculated as the product of concentration and discharge (volume). There are problems with this approach. In some cases, the calculated concentrations in flood waters probably exceed the solubility limits, as in the case of sulfate in the presence of barium. In other instances, precipitation of barium sulfate or iron and manganese hydroxides might greatly reduce the concentration of radium and uranium, both of which would coprecipitate. Thus the stream concentrations shown in Table 3.44 are probably high (conservative). To improve the analysis, additional comparisons or parallels were drawn using mill tailings solutions and stream water quality as affected by mine drainage and a mill tailings spill.

Contaminant concentrations in uranium mill tailings liquids provide an upper limit estimate of runoff concentrations insofar as the solvent action of tailings solutions maximize dissolution of minerals present in the ore (J. Kunkler, USGS, written communication, 1979). Table 3.46 is a compilation of mill tailings water quality data from numerous previous reports and summarized by EPA and USGS staff (Ka79; Ku79). It is apparent that there are wide variations as a function of mining region and whether an acid or alkaline leach mill circuit is used. The Nuclear Regulatory Commission (NRC79b) assumption for the composition of a "typical" acid leach mill is shown along with other average or representative analyses. A conservative (worst quality) analysis for uranium mill pond water quality is estimated as follows (Table 3.47) and compared to the average concentrations calculated from the mixing of mine effluent and flood volumes (Table 3.44).

The data in Table 3.47 suggest that calculated concentrations in the sub-basin almost without exception exceed those in uranium mill tailings solutions. Thus, the calculated values are probably erroneously high. Calculated concentrations in flood waters of the basin and regional basin streams are considerably less and are in rough agreement with field data, at least for the stable constituents. Radium-226 and lead-210, however, still seem excessively high considering the various natural processes of sorption, precipitation, and so on. To understand the degree to which natural streams transport contaminants, we reviewed water quality data from selected New Mexico streams receiving mine drainage.

Table 3 46 Radiochemical & stable element/compound water quality for selected acid & alkaline leach uranium mill tailings ponds in the United States

Tailings Pile Location	U (mg/l)	Th-230	Ra-226 (pCi/l)	Pb-210	Po-210	As	Mn	Cu	Se	Mo	V (mg/l)	SO	Na	Fe	TDS	NH	Ca	NO	Cl
1. Split Rock, WY (acid)	10.5	41600	4800	---	940	1.1	15.5		0.2	1	0.05			280	11810	374	560	43.5	65
2. Canon City, CO (acid)	---	---	---	---	---	10.1	25.0	18	0.6	190	7.1	34000	19000	280	77400	---	380	140	6500
3. Moab, UT	2.0	50	100	---	---	7.0	---	--	---	---	---	100	---	---	150000	---	---	---	300
4. United Nuclear, 14 NM (acid)(a)		---	38	---	---	---	50	3	0.005	3	30	---	300	1000	---	---	700	---	---
5. Anaconda Inj. Well Feed, NM	130	---	53	---	---	---	340	--	0.03	---	6.3	4900	1200	---	---	69	---	7.4	---
6. Kerr-McGee, NM (acid)(a)	32	---	58	---	---	---	30	5	0.18	7	10	---	500	1000	---	---	300	---	---
7. UN-HP, Grants, NM (alkaline)	150	---	52	---	---	---	---	--	0.92	70	6.8	4300	4300	---	---	4.4	---	4.4	2
8. Humecca, WY (acid)	68.4	110	240	---	---	---	0.4	0.1	---	---	---	6500	11700	0 5	---	460	---	16	16000
9. USNRC-Uranium Milling EIS(acid)	8 0	150000	400	400	400	0.2	500	50	20	100	0.1	30000	500	1000	35000	500	500	---	300
10. Representative acid millpond in New Mexico (a)	---	---	---	---	---	---	---	--	---	---	---	44000	---	170	---	300 ^(b)	---	---	1800
"Average" (Exclusive of 9 and 10)	58	13920	760	---	---	6	160	6.5	0.32	54	10	10000	6200	510	80000	227	485	40	1700
Maximum value:																			
"Average" versus NRC GEIS	58	150000	760	400	400	6	500	50	20	100	10	30000	6200	1000	80000	500	500	40	1700

(a) Source: Ku79.
(b) Ammonium ion.

Table 3.47 Summary of flood runoff water quality and
uranium millpond quality

Parameters	Concentration in uranium mill tailings solution	Concentration in flood waters of the sub-basin ^(a)	Concentration in flood waters of the Rio San Jose ^(a)
Uranium (mg/l)	58	235 - 6970	7
Radium-226 (pCi/l)	760	229 - 6780	6.9
Lead-210 (pCi/l)	400	2430 - 72000	73
Polonium-210 (pCi/l)	400	NC ^(b)	NC
Arsenic (mg/l)	6	2.1 - 61	0.062
Manganese (mg/l)	500	NC	NC
Copper (mg/l)	50	NC	NC
Selenium (mg/l)	20	11 - 220	0.34
Molybdenum (mg/l)	100	48 - 2400	1.4
Vanadium (mg/l)	10	NC	NC
Sulfate (mg/l)	30,000	9.7×10^4 - 2.87×10^6	2901

(a) Refer to Table 3.44.

(b) Not calculated.

The USGS, by water sampling in the Churchrock area of New Mexico (J.L. Kunkler, USGS, written communication, 1979), determined water quality in an ephemeral stream receiving rather large and continuous mine discharges. Data are also available from the Schwarzwald Mine near Golden, Colorado (EPA72). Until 1972, this mine discharged effluent high in uranium, radium, and trace elements to Ralston Creek and subsequently to two lakes/reservoirs used for irrigation and potable supply (Section 3.2.3.2.1).

The way in which surface runoff water quality is created or affected by mine discharge is complex. In the Churchrock area, numerous water quality changes occur as the mine discharges flow toward Gallup (Fig. 3.21 and Table 3.48). As in other uranium mining areas in New Mexico, stream volume constantly decreases with flow distance, but water quality changes are erratic. Infiltration, discussed in more detail in the following section of the report and in Appendix H, amounts to about 90 percent or more of the water loss. The balance is by evaporation. On a percentage basis, similar losses occur in the principal drainage courses in Ambrosia Lake. Dissolved Ra-226 decreases from 30 to 0.88 pCi/l in a reach of 9.2 km and, on a later date, from 14 to 0.95 pCi/l in a distance of 26.7 km. Based on the limited flow and water quality data, it appears that radium is strongly sorbed onto the stream sediments. In October 1975, soluble uranium decreased from 1150 to 740 $\mu\text{g/l}$ in the reach immediately below the mine discharges, yet in July 1977 and May 1978 uranium increased in the downstream direction from 580 to 860 $\mu\text{g/l}$ and from 970 to 2800 $\mu\text{g/l}$. These changes bear no consistent relation to fluctuations in dissolved or suspended solids along the flow path. Both of the latter parameters appear to increase in the direction of flow and may be a result of flash floods in lower reaches of the basin. Uranium appears to undergo little change and may actually increase in the downstream direction. Of the stable trace elements, vanadium, selenium, iron, molybdenum, and zinc show no consistent change with distance.

A third approach used to assess surface runoff quality involved a brief review of some of the data collected to monitor a July 1979 tailings accident in New Mexico. The mill tailings dam at the Churchrock mill breached and dumped 223,000 m^3 of liquid and 1,000 metric tons of solids into the Rio Puerco drainage system. The catastrophe immediately spurred numerous water quality studies by State and Federal agencies. Numerous inter-

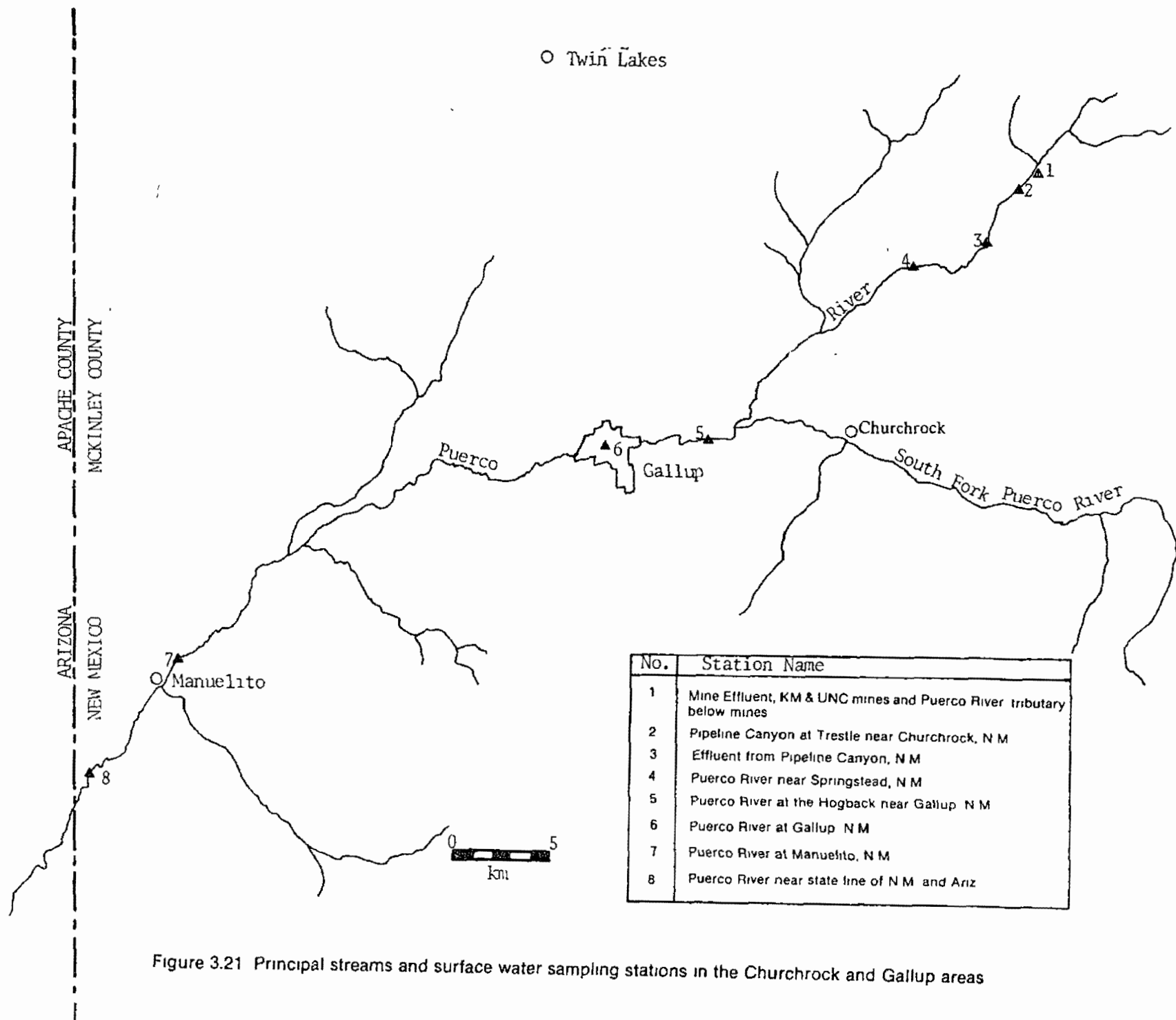


Figure 3.21 Principal streams and surface water sampling stations in the Churchrock and Gallup areas

Table 3.48 Flow and water quality in the Puerco River near Churchrock and Gallup, New Mexico

Location and (Station Number)	m ³ /min	U nat. μg/l as U ₃ O ₈	Ra-226 pCi/l	Total Solids, mg/l		Suspended solids, metric tons per day	Concentrations μg/l									
				Dissolved	Suspended		Ba	Cd	Cr	Pb	Mo	V	Zn	Se	As	Fe
<u>Oct. 16, 1975</u>																
Puerco River tributary below mines - (1)	14.5	1150	30	430	410	9.5	-	-	-	-	-	21	-	27	-	20
Puerco River near Springstead, NM -(4)	12.4	740	0.88	480	1600	8.67	-	-	-	-	-	13	-	25	-	30
Puerco River at Gallup, NM - (6)	5.11	---	0.52	640	2300	5.14	-	-	-	-	-	5.7	-	26	-	40
Puerco River at Manuelito - (7)	6.8(est)	540	0.25	800	2800	---	-	-	-	-	-	-	-	-	-	-
<u>July 6, 1977</u>																
Puerco River tributary below mines - (1)	11.55	580	14	410	260	---	800	1	0	6	-	-	0	25	1(3)	10
Puerco River at the Hogback, near Gallup, NM - (5)	6.47	860	0.95	520	15000	---	100	1	0	11	-	-	50	20	1(19)	80
Puerco River near State line (NM/AZ) - (8)	15.5	83	0.27	600	44000	---	1700	4	0	2	-	-	30	5	6(7)	90

Table 3.48 (Continued)

Location and (Station Number)	m ³ /min	U nat.	Ra-226 pCi/l	Total Solids, mg/l		Suspended solids, metric tons per day	Concentrations $\mu\text{g/l}$									
		$\mu\text{g/l}$ as U ₃ O ₈		Dissolved	Suspended		Ba	Cd	Cr	Pb	Mo	V	Zn	Se	As	Fe
<u>May 25, 1978</u>																
Effluent from Kerr McGee and United Nuclear Mines, Churchrock, NM- (1)	10.9	807	2.6	---	---	---	-	-	-	-	12	19	-	4	-	-
Effluent from Pipeline Canyon, NM - (3)	9	2800	1.5	---	---	---	-	-	-	-	820	28	-	110	-	-
Puerco River near Springstead, NM - (4) (sampled 5/18/78)	10.88	1100	0.8	---	---	---	-	-	-	-	12	16	-	0	-	15
<u>July 11-12, 1978</u>																
Pipeline Canyon at trestle near Churchrock, NM - (2)	14.45	940	8.6	---	---	---	-	-	-	-	230	11	-	-	-	540
Effluent from Pipeline Canyon, NM - (3)	14.3	1120	1.3	---	---	---	-	-	-	-	260	6	-	11	-	70
Puerco River near Springstead, NM - (4)	---	1130	2.2	---	---	---	-	-	-	-	240	9	-	13	-	40

Source: New Mexico District office of the U.S. Geological Survey (Peter Frenzel, written communication, 1979 and Kunkler, 1979).

pretations of the data have led to some confusion, compounded in some instances by inconsistent sample collection and preservation. However, several general findings seem true. Dissolution of stable and radioactive trace contaminants in flood waters does not seem significant providing that pH of the flood is in the range of 4 to 7. After several days, the mill tailings liquid was diluted and neutralized and contaminant concentrations decreased -- sometimes to levels lower than before the accident (J. Kunkler, USGS, written communication, 1979). At a downstream sampling station near Gallup, some 30 kilometers from the spill, dissolved uranium and radium-226 about 36 hours after the spill were 3.1 mg/l and 0.95 pCi/l, respectively. Suspended sediments contained 19 ppm uranium and 0.72 pCi/g radium-226. For the latter, this is less than background.

The surface water quality data pertaining to discharge of mine effluents and to the July 1979 spill seem to indicate rapid and thorough removal of radium-226 as a result of sorption, precipitation, pH adjustment, etc. However, stream sediment analyses in the Grants Mineral Belt are scarce, and there are no analyses of suspended solids in flood waters. Stream-bed sediment analyses by the USGS indicate less sorbed radium-226 and uranium than expected (Ku79). During this spill incident, uranium and selenium were relatively mobile in surface streams.

From the foregoing review of the literature and field data and preliminary calculations of runoff quality (Table 3.44), the following general conclusions are offered:

1. Radium-226 is removed from surface water in the New Mexico study area at rates of 0.5 to 3 pCi/l per kilometer of stream. Final concentrations are on the order of 0.25 pCi/l. Resolution in successive surface flows occurs, but it is not significant.
2. Uranium and certain stable trace elements, such as selenium, vanadium, molybdenum, and iron, show no consistent reduction with flow distance and may show an increase, at times.
3. Considerable more data collection is needed to understand the fate of dissolved and suspended contaminants from mine drainage. The present data base is rather limited in terms of sampling frequency, variety of contaminants measured, and types of measurements, for example, suspended solids analyses for flood waters.

4. With the exception of radium-226, the preliminary calculations of runoff quality in Table 3.44 are believed to be a first approximation of field conditions. Additional studies specific to the principal mining districts are needed.

5. Dissolved radium-226 concentrations in runoff are believed to be several picocuries per liter or less under natural conditions.

6. Uranium is fairly mobile and probably the most significant radionuclide in uranium mine effluent.

3.4.3.2 Impacts of Seepage on Groundwater

The principal use of groundwater in the immediate area of the mines is for stock water. Wells in the highland areas are typically one to two hundred meters deep and completed in underlying bedrock strata (Co68; Ka75). Contamination of such wells by mine discharge is considered extremely unlikely. Shallow wells are few in number and located along major drainages that are typically ephemeral. Such shallow wells are susceptible to contamination if located downgrade from mine discharges. Municipal water supplies are usually developed from wells because groundwater is consistently available and has acceptable suspended and dissolved mineral contents. The aquifers tapped by municipal wells are mostly either quaternary lava flows or deeper mesozoic sandstone and carbonate sequences. Considering the distance from the mining centers to the communities and the hydrogeologic conditions, it is unlikely that mining will cause measurable deterioration of municipal water quality. The greatest likelihood for contaminated groundwater is in the shallow, alluvial aquifer beneath streams receiving mine drainage. It is extremely unlikely that water quality in deeper, artesian aquifers will be adversely affected by mine discharge or overland flow affected by solid wastes. Shallow wells in these locations have been constructed in the past, but there are only a few and they are used for stock watering. It is possible that recharge of substantial quantities of mine water to the shallow aquifer will encourage additional use of it, in which case water quality will be of concern.

Table 3.49 shows average and extreme concentrations of various common and trace constituents in groundwater and other measures of water quality. The data are composited from a previous study (EPA75) and from unpublished

analyses by the New Mexico Environmental Improvement Division (J. Lazarus, NMEID, oral communication, 1979). We have categorized the data according to principal aquifers, which are in areas where the groundwater is not believed to be contaminated by mining. Because it is common for a well to tap more than one aquifer, the differences in water quality in Table 3.49 are approximate at best. The data reveal no sharp differences in water quality amongst the three major aquifers. The San Andres Limestone, a major aquifer for municipal and industrial uses in the Grants and Milan areas, has equal or greater concentrations of most constituents as compared to the Westwater Canyon Member and Gallup Sandstone units, which are closely associated with uranium mineralization.

Theoretical analysis of radionuclide transport in groundwater beneath and adjacent to a uranium mill tailings pond reveals very limited migration of radionuclides in groundwater (Se75). Using a seepage rate of 4×10^{-7} cm/sec and a 10 percent loss of soluble radionuclides, numerical solutions for steady state flow and transport into unconsolidated sand for periods of 5 years and 20 years reveal up to several meters movement of radium-226, thorium-230/234, uranium, and lead-210 after 20 years of leaching. For example, radium in groundwater to a depth of 3 meters is 10 percent of that in the tailings pond. Because the other isotopes tend to have even greater sorption, migration distances are further reduced. Although field studies at three uranium mill tailings piles in the Grants Mineral Belt substantiate only local migration of radionuclides (EPA75), extensive lateral migration of stable chemical species has been observed at uranium mills in Colorado, Wyoming, and Washington (Ka79, Ka78a, He79). For example, with respect to the old Cotter uranium mine at Canon City, Colorado, the Colorado Department of Health has stated in its Final Executive Licensing Summary, August 17, 1979, that "contamination attributed to tailings liquid was observed in an off-site water well ten years after the mill began depositing tailings, a [migration] rate of over five hundred feet per year." With respect to the same site, one researcher has stated that "the soluble uranium content of Lincoln Park ground waters is highly elevated with respect to Arkansas River water and exceeds suggested thresholds below which ecological and health effects are not expected. Molybdenum concentrations in these ground waters greatly exceed irrigation standards as well as the ALG based on health and ecological effects...." (Dr79). Near neutral pH and relatively low concen-

Table 3.49 Groundwater quality in principal aquifers in the Grants Mineral Belt, New Mexico

Parameter	Aquifer		
	San Andres Limestone	Westwater Canyon Member, Morrison Fm. and Gallup Sandstone	Quaternary Alluvium, Tertiary Volcanics, and Chinle Formation
pH	7.2 ^(a) (6.9 - 7.5)	7.9 (6.7 - 9.15)	7.6 (6.25 - 8.8)
Spec. cond. μmhos/cm	1900 (720 - 3500)	1800 (550 - 4250)	1715 (700 - 4000)
TDS	1680 (490 - 4500)	1160 (340 - 2300)	1240 (490 - 3800)
Cl mg/l	98 (< 0.2 - 270)	15 (0 - 98)	57 (6.2 - 260)
Se mg/l	0.31 (0.01 - 1.52)	0.02 (0.01 - 0.13)	0.59 (0.02 - 1.06)
V mg/l	0.88 (0.4 - 1.3)	0.3 (0.3 - 0.3)	0.55 (0.3 - 1.3)
Radium-226 pCi/l	0.47 (0.11 - 1.92)	0.71 (0.07 - 3.7)	0.22 (0.05 - 0.72)
Uranium, mg/l	1.31 (0.04 - 2.6)	0.35 (0.02 - 1.0)	4.72 (0.07 - 14)
Th-230, pCi/l	0.12 (0.017 - 0.52)	0.030 (0.015 - 0.053)	0.212 (0.018 - 0.65)
Th-232, pCi/l	0.11 (0.0053-0.54)	0.015 (< 0.01-0.036)	0.123 (0.0094-0.99)
Po-210, pCi/l	0.75 (0.070 - 2.3)	0.42 (0.19 - 0.79)	0.193 (0.010 - 0.55)

(a) Mean and range of values shown.

Note.--Selenium, vanadium, and uranium values for the limestone and alluvium/chinle aquifers are based on 4 to 5 analyses and must be regarded as tentative.

trations in mine effluents, together with low hydraulic heads, indicate short migration distances in groundwater for radionuclides and most stable trace elements in mine effluents.

Discharge of water pumped from mines to arroyos has both hydraulic and water quality impacts on shallow groundwater in the alluvial aquifer. The seepage model (Appendix H) and scattered field measurements in the Grants Mineral Belt substantiate that significant groundwater recharge is associated with mine discharge. Water quality effects on groundwater are poorly documented, however. We do not address the influence of impoundments used to remove suspended solids from mine effluents before discharge. Seepage water losses from such impoundments are believed to be small, especially when compared to infiltration losses in the arroyos and open fields receiving most of the wastes not piped to mills for process water. The impoundments are rather small and tend to become self-sealing due to settlement of fines. In at least one instance in Ambrosia Lake, the mine pond is lined to prevent seepage.

Unpublished flow and water quality data from the U.S. Geological Survey (P. Frenzel, written communication, 1979) document conditions in the Rio Puerco drainage near Churchrock and Gallup, New Mexico. Figure 3.21 shows the sampling station locations, and the chemical data are in Table 3.47. From October 1975 gaging data, seepage and evaporation reduce flow $9.39 \text{ m}^3/\text{min}$ in a reach of 30.2 km, a loss of $0.31 \text{ m}^3/\text{min}/\text{km}$. Conservatively assuming 20 percent of this is by evaporation, seepage is $7.5 \text{ m}^3/\text{min}$ or $3.94 \times 10^6 \text{ m}^3/\text{yr}$. Gaging data for July 1977 and May 1978 similarly indicate average bed losses of $0.24 \text{ m}^3/\text{min}/\text{km}$. In the Ambrosia Lake district (data not shown), discharges (to San Mateo Creek and Arroyo del Puerto) from about a dozen mines total about $10.8 \times 10^6 \text{ m}^3/\text{year}$, and the total length of perennial stream is about 15 kilometers. Assuming an average stream width of one meter and the above evaporation rate, evaporation and infiltration are $0.06 \text{ m}^3/\text{min}$ and $7.54 \text{ m}^3/\text{min}$, respectively. In this case, infiltration amounts to 99 percent of total loss. Dissolved solids range from 520 to 1231 mg/l (mean 743 mg/l), and Ra-226 ranges from 0.2 to 23 pCi/l (mean 6.6 pCi/l). Selenium and molybdenum average 0.010 and 0.22 mg/l, respectively.

Considering these two areas, evaporation averages about 4 percent of mine discharge versus the value of one percent calculated in Appendix H. Obviously, increased evaporation is accompanied by decreased infiltration.

Infiltration ranges from at least 90 percent to perhaps 99 percent of mine discharge, or from 1.8 to 1.98 m³/min per mine. The foregoing field data and the more theoretical approach used in Appendix H show reasonable agreement on the relative amounts of infiltration and evaporation. We conclude then that most of the mine effluent infiltrates within relatively short distances of the mine(s) and recharges the shallow water table. The dissolved, generally nonreactive contaminants such as chloride and sulfate are expected to reach the water table, but reactive contaminants such as radium-226 and most trace metals would sorb or precipitate in the soil (substrate) in the course of infiltration.

The influence of mine discharge on groundwater quality beneath formerly ephemeral streams now receiving the discharge is currently under investigation by the New Mexico Environmental Improvement Division. Monitoring wells have been installed at several locations along the Rio Puerco (west) in the Churchrock area and San Mateo Creek in the Ambrosia Lake district. Table 3.50 summarizes partial results of samples taken in the last 12 to 18 months. In the Ambrosia Lake district, marked deterioration in water quality between the Lee Ranch and Sandoval Ranch stations on San Mateo Creek is a result of either natural causes and (or) mine drainage from a nearby deep underground uranium mine. Between Sandoval Ranch and Otero Ranch even more pronounced changes occur. In this short reach of 2.5 km, contaminated flows from uranium mines, ion-exchange plants, and seepage from an acid leach uranium mill enter Arroyo del Puerto, a tributary of San Mateo Creek. Additional study of surface water quality in the Arroyo del Puerto is recommended to further characterize the obviously interconnected surface water and groundwater systems.

In the Churchrock area, drained by the Rio Puerco, groundwater quality changes in the downstream direction are not readily apparent (Table 3.50). Although there is an acid leach mill also adjacent to the Rio Puerco tributary receiving the mine discharges, the mill is relatively new (1978 start-up) and may not yet influence stream quality. Most of the discharge from one of the two mines is used as mill feed water, thereby causing decreased discharge from the mines to the stream. Nevertheless, the reach of the perennial stream is increasing, indicating infiltration of remaining mine effluent and addition of water to storage in the shallow aquifer. Storage changes have been confirmed by static groundwater level measurements in the area east of

Table 3.50 Groundwater quality associated with the San Mateo Creek and Rio Puerco (west) drainages in the Grants Mineral Belt, New Mexico

Station	Sulfate (mg/ℓ)	Molybdenum	Selenium (μg/ℓ)	Uranium
<u>San Mateo Creek</u>				
Lee Ranch	125.7	<10	< 5	<10
Sandoval Ranch	225-274	103-235	4-14.7	293-400
Otero Ranch	463-989	350-516	33-59	680-860
<u>Rio Puerco (west)</u>				
Hwy. 566 Bridge on N. Fork Rio Puerco	101-223	<10-284	20-22	530-760
Rio Puerco at Fourth St. Bridge, Gallup	163-244	<10-215	9-26	550-625

Source: Based on unpublished 1978 data developed by the New Mexico Environmental Improvement Division (J. Lazarus, oral communication, 1979).

Gallup. A massive spill of mill tailings into the Rio Puerco occurred in July 1979 and will complicate water quality investigation, insofar as the mine and mill influences are now superimposed in terms of both solid and liquid waste loadings in the watershed. The tailings "flood," estimated to contain about 360,000 m³ of fluid and 1000 MT of solids, was traced into Arizona.

In summary and considering the high volume of dilute mine discharges,

which are enriched in certain stable and radioactive toxic trace elements (EPA75; Hi77), we recommend that water quality effects of mine discharge be very carefully evaluated in at least a few selected areas. Available stream-flow data indicate that infiltration is the principal means of disposal, yet the water quality data base, in particular, is rather weak to assess whether adverse impacts are likely. It is expected that future discharges in the Churchrock area alone will amount to about $40 \text{ m}^3/\text{min}$ and will contain less than 400 mg/l dissolved solids, most of which is sodium and bicarbonate. Dissolved concentrations of uranium, radium, iron, selenium, and vanadium are elevated relative to drinking water limits and infiltration of uranium, selenium, and possibly other stable elements warrants study. Use of settling ponds and barium chloride treatment greatly reduces the suspended solids, uranium, and radium concentrations. The final composition and ultimate disposal of pond sediments and added chemicals is essentially undocumented and bears additional investigation. Lastly, mine dewatering creates marked regional cones of depression and reduces the flow of water to existing supply wells and the baseflow component in major drainage systems such as the San Juan River (Ly79).

3.4.4 Gases and Dusts from Mining Activities

3.4.4.1 Radon-222 in Mine Exhaust Air

Unlike surface mines, large capacity ventilating systems are required in underground uranium mines, primarily to dilute and remove Rn-222 that emanates from the ore (Section 1.3.3). Ventilation rates vary from a few hundred to a few hundred thousand cubic meters of air per minute, and measured Rn-222 concentrations in mine vent air range from 7 pCi/l to $22,000 \text{ pCi/l}$ (Ja79b). The concentration of Rn-222 in mine exhaust air varies depending upon ventilation rate, mine size (volume) and age, grade of exposed ore, size of active working areas, rock characteristics (moisture content and porosity), effectiveness of bulkhead partitions, barometric pressure, ore production rates, and mining practices. The emanation of Rn-222 dissolved in water that seeps into most mines may also contribute to Rn-222 in the exhaust air.

Because of the numerous variables that affect Rn-222 concentrations in mine air, it is difficult to confidently model radon releases from underground mines. A useful model would be one that would relate radon emissions to the production of U_3O_8 . Measurements relating radon emissions to ore production have been made at seven underground uranium mines in New Mexico (Ja79b). The results of these measurements varied at the different mines from 1,380 to 23,500 Ci Rn-222 per AFR*, with an average rate of 4,300 Ci Rn-222 per AFR. The higher emission rates were noted to occur at the older mines. This was believed due to larger surface areas of exposed ore and sub-ore in the older mines. That is, inactive mined-out areas increase with mine age, and the ceiling, floors, and walls of these areas still contain certain amounts of ore and sub-ore. Radon emanating from these surface areas tend to increase the Rn-222 content of exhausted mine air unless these inactive areas of the mine (rooms, stopes, drifts, etc.) are effectively sealed. Because the radon emission factor is so variable in terms of Ci per AFR, an emission rate based on cumulative U_3O_8 mined has been proposed for modeling purposes (Ja79b). It is believed that this relationship would reduce the apparent dependence of the emission rate on the mine age. However, data are not presently available to make this latter correlation.

Although the average measured Rn-222 exhaust factor of 4,300 Ci/AFR is tentative and may be improved by studies in progress (Ja79b), it is the only value currently available for modeling purposes and will, therefore, be used in the present assessment. Assuming that 1 AFR is equivalent to 245 MT** of U_3O_8 (Ja79b), 0.017 Ci of Rn-222 will be released from the mine vents per metric ton of 0.1 percent grade ore mined. This emission rate will include all underground sources, i.e., emanation from exposed ore and blasting, slushing, loading, and transporting ore bearing rock. Radon-222 emissions were estimated for the two model underground mines by multiplying their

*AFR = Annual Fuel Requirement for a 1000 MWe LWR.

**The AFR value on which the exhaust factor was based.

respective annual ore capacities by the above emission rate. Table 3.51 lists the results.

The estimated annual radon release computed for the average underground mine is compared below with releases reported elsewhere. Agreement is reasonably good.

<u>Source</u>	<u>Annual Release of Rn-222, Ci</u>
This Study	306
Tr79	289 - 467 ^(a)
TVA78a	1577
TVA78b	180
TVA79	215
Th79	87

(a) Adjusted for 0.1 percent ore grade.

By properly capping the exhaust vents and sealing the shaft and mine entrance, radon emission rates from inactive mines will be a negligible fraction of the radon release rate that occurs during active mining.

3.4.4.2 Aboveground Radon-222 Sources

Radon-222 will be released from the following aboveground sources.

1. Dumping ore, sub-ore, and waste rock from the ore skip into haul trucks and unloading them on their respective piles.
2. Reloading ore from the stockpile after a 41-day residence time.
3. Emanation from waste rock, sub-ore, and ore storage pile surfaces.

The annual quantities of Rn-222 released by sources 1 and 2 were estimated using the following factors and assumptions.

- Radon-222 is in secular equilibrium with U-238.
- The density of ore, sub-ore, and waste rock is 2.0 MT/m^3 .
- Annual production rates of ore and sub-ore are equal and assumed to be $1.8 \times 10^4 \text{ MT}$ at the average mine and $2 \times 10^5 \text{ MT}$ at the average large mine (Sections 3.4.1.1 to 3.4.1.3).
- The production rate ratio of ore to waste rock is 9.1:1 (Section 3.4.1.1).
- All Rn-222 present is available for release, 0.00565 Ci/m^3 per

Table 3.51 Estimated annual radon-222 emissions from underground uranium mining sources

Source	Average Mine ^(a) , Ci/yr	Average Large Mine ^(b) , Ci/yr
<u>Underground</u>		
Mine vent air	306	3,400
<u>Aboveground</u>		
Ore loading and dumping	1.4	15.3
Sub-ore loading and dumping	0.5	5.3
Waste rock loading and dumping	0.003	0.03
Reloading ore from stockpile	1.4	15.3
Ore stockpile exhalation	6.3	53
Sub-ore pile exhalation	61	338
Waste rock pile exhalation	<u>0.5</u>	<u>2.6</u>
Total	377	3830

(a) Annual production of ore and sub-ore = 1.8×10^4 MT, waste rock = 2.0×10^3 MT.

(b) Annual production of ore and sub-ore = 2×10^5 MT, waste rock = 2.2×10^4 MT.

percent U_3O_8 (Ni79), with an emanation coefficient of 0.27 (Au78, Tanner, A.B., Department of Interior, Geological Survey, Reston, VA., 11/79, personal communication).

The quantities of U_3O_8 present in ore, sub-ore, and waste rock are 0.10 percent, 0.035 percent and 0.0020 percent, respectively (Sections 3.4.1.1 to 3.4.1.3).

Substituting the above values into the following equation yields the Rn-222 releases given in Table 3.51 for the average mine and the average large mine.

$$\text{Rn-222 (Ci/yr)} = (\text{percent } U_3O_8) \left(\frac{0.00565 \text{ Ci}}{\text{m}^3 \cdot \text{percent}} \right) (0.27) \left(\frac{\text{m}^3}{2.0 \text{ MT}} \right) \times (\text{Production Rate, } \frac{\text{MT}}{\text{yr}}) \quad (3.11)$$

These releases are maximum values since very little time will have elapsed between the underground (blasting, slushing, loading, etc.) and surface operations. A significant amount of the radon that is available for release will emanate during the underground operations and invalidate the first assumption above concerning radioactive equilibrium. Nevertheless, these estimated maximum releases are very small in comparison to the radon released from the mine exhaust vents.

The emanation of Rn-222 from waste rock, sub-ore, and ore piles is based on an exhalation rate of $0.092 \text{ Ci/m}^2 \cdot \text{yr} \cdot \text{percent } U_3O_8$ (Ni79) and ore grades of 0.002 percent, 0.035 percent, and 0.10 percent, respectively. Surface areas of the ore piles (Table 3.37), sub-ore piles (Table 3.38), and waste rock piles (Table 3.36) were used in these calculations. Applying these parameters, the annual Rn-222 emissions from the waste rock, sub-ore, and ore piles at the average mine and average large mine were computed. Table 3.51 gives the results. Total annual Rn-222 emissions during underground mining operations is the sum of the releases from all sources considered: 377 Ci from the average mine and 3830 Ci from the average large mine. More than 80% of the Rn-222 emissions results from the mine vent air.

3.4.4.3 Dusts and Fumes

Vehicular emissions resulting from the combustion of hydrocarbon fuels in gasoline and diesel-powered equipment are considerably less at underground mines than at surface mines (Section 3.3.4.1). The principal emissions are

particulates, sulfur oxides, carbon monoxide, nitrogen oxides, and hydrocarbons. The quantity of these combustion products released to the atmosphere depends on the number, size, and types of equipment used, all of which are directly related to ore production.

EPA has estimated the following emissions from mining 1350 MT of ore per day from an underground mine (Re76).

<u>Pollutant</u>	<u>Emissions per Operating Day, Kg/d</u>
Particulates	2.4
Sulfur Oxides	5.0
Carbon Monoxide	41.9
Nitrogen Oxides	68.1
Hydrocarbons	6.9

Assuming a 330 operating-day year (Ni79), these emissions were adjusted according to the annual ore production of the average mine (1.8×10^4 MT) and the average large mine (2×10^5 MT). Table 3.52 lists the total airborne combustion product emissions. These emissions are small compared to those at surface mines (Table 3.30). For example, these estimates indicate that the emissions of combustion products at the average surface mine are more than 100 times greater than those at the average underground mine.

At underground mines, dust is produced by both underground and surface operations. No measurements have been made of dust concentrations in mine exhaust air. Because underground mines are wet, which greatly reduces dust production, and since a large portion of the dust produced would probably deposit underground, dust emissions from underground operations are probably relatively small. Hence, dust emissions from underground operations will not be assessed.

Aboveground sources of dust include dumping ore, sub-ore, and waste rock from the skip into haul trucks; dumping these materials onto their respective piles; reloading ore from the stockpile; using dirt haul roads by vehicular traffic; and dust suspended by the wind from the waste rock, sub-ore, and ore piles. These sources will be assessed as was done previously for surface mines (Section 3.3.4.1).

Table 3.52 Estimated air pollutant emissions from heavy-duty equipment at underground uranium mines

Pollutant	Emissions, Kg/yr ^(a)	
	Average Mine ^(b)	Average Large Mine ^(c)
Particulates	32	350
Sulfur oxides	67	740
Carbon monoxide	560	6,210
Nitrogen oxides	910	10,100
Hydrocarbons	92	1,020

(a) Based on Re76 and 330 operating days per year.

(b) Annual ore production = 1.8×10^4 MT.

(c) Annual ore production = 2×10^5 MT.

Dust emissions will vary over a wide range depending upon moisture content, amount of fines, number and types of equipment operating, and climatic conditions. Because ore is generally wet, the relative amounts of dust produced from its mining and handling are usually small. The following emission factors were selected from those suggested by the EPA for loading and dumping operations (Hu76, Ra78, Da79):

truck loading = 2.5×10^{-2} kg/MT; and

truck dumping = 2.0×10^{-2} kg/MT.

Average annual dust emissions were estimated for the aboveground mining activities by applying these emission factors to the ore, sub-ore, and waste rock production rates of the average mine and average large mine. Table 3.53 lists the results. One-half the emission factor values were applied to ore and sub-ore because they are generally wet, except when reloading ore from the stockpile. In that case, it is assumed to have dried during the 41-day residence period (Section 3.4.1.2). Also, the emission factor for truck loading was assumed valid for loading the haul trucks from the mine skip. The dust emission for truck dumping may be high since it was based on dumping of aggregate, which would have a smaller particle size distribution than the ore, sub-ore, or waste rock (Hu76).

The movement of heavy-duty trucks is a large source of dust at most uranium mines. The magnitude of this source depends upon a number of factors, including the particle size distribution and moisture content of the road bed material, vehicular speed and distance traveled, and meteorological conditions. Emission factors for heavy-duty haul trucks (1.15 kg/VKmt) and light duty vehicles (1.03 kg/VKmt) are the same as those computed for these vehicles at surface mines (Section 3.3.4.1). Dust emissions for the movement of heavy-duty haul trucks were estimated using the appropriate emission factor and assuming --

- . 31.8 MT truck capacities;
- . round-trip haul distances of 1.61 km to the ore and sub-ore piles and 3.22 km to the waste rock pile; and
- . the annual production rates given in Sections 3.4.1.1, 3.4.1.2 and 3.4.1.3.

Table 3.53 lists the results.

Additional dust emissions will occur from light-duty vehicular traffic along access roads. Using the emission factor derived in Section 3.3.4.1 (1.03 kg/VKmt) and assuming that there are 16 km of access roads traveled 4 times a day during the 330 operating days per year, about 22 MT of dust will be produced from this source annually. Emissions that occur during haulage road maintenance is relatively small and will not be considered.

Heavy-duty, haul truck traffic at underground uranium mines produces considerably less dust than at surface mines. This is to be expected because of the vast quantities of overburden that must be transported as well as larger ore and sub-ore capacities at surface-type mines.

The dust emissions computed above for transportation assume no effective dust control program. But, haul roads are normally sprinkled routinely during dry periods, and stabilizing chemicals are applied to roadways, usually to the ore haul roads. Dust emissions along haul roads can be reduced by 50 percent from sprinkling and up to 85 percent by the application of stabilizing chemicals (EPA77b, Da79).

Table 3.53 also lists average annual dust emissions caused by wind erosion of waste rock, sub-ore, and ore piles at the model underground mines. Emission factors, computed in Appendix I, are 2.12 MT/hectare-yr for waste rock and sub-ore piles and 0.040 kg/MT for the ore stockpiles. The first emission factor was multiplied by the waste rock and sub-ore pile surface

Table 3.53 Estimated average annual dust emissions from underground mining activities

Source ^(d)	Dust Emissions, MT/yr					
	Average Mine ^(a)			Average Large Mine ^(b)		
	Ore ^(c)	Sub-ore ^(c)	Waste Rock	Ore ^(c)	Sub-ore ^(c)	Waste Rock
Loading truck from skip at mine shaft	0.23	0.23	0.05	2.5	2.5	0.6
Truck dumping at piles	0.18	0.18	0.04	2.0	2.0	0.4
Reloading stock-piled ore ^(e)	0.45	NA ^(f)	NA	5.0	NA	NA
Wind suspended dust from piles	0.72	4.0	0.57	8.0	22	3.0
Transportation ^(g)	1.0	1.0	0.23	11.6	11.6	2.6

(a) Based on annual production rates of 1.8×10^4 MT of ore and sub-ore, and 2.0×10^3 MT of waste rock.

(b) Based on annual production rates of 2×10^5 MT of ore and sub-ore, and 2.2×10^4 MT of waste rock.

(c) Assumed wet.

(d) Aboveground activities.

(e) Assumed dry.

(f) NA - Not applicable.

(g) Dust emissions from heavy-duty, vehicular traffic along ore, sub-ore, and waste rock haul roads.

areas given in Tables 3.36 and 3.38, respectively, while the second factor was multiplied by the annual ore production.

Table 3.54 shows annual contaminant emissions caused by mining activities (loading and dumping) according to source location, at the mine shaft and at the piles. Contaminant emissions were computed by multiplying the total annual dust emissions at each pile (Table 3.53) by the respective contaminant concentrations in each source--waste rock (Section 3.4.1.1; Table 3.16), sub-ore (Section 3.4.1.3; Table 3.19), and ore (Section 3.4.1.2; Table 3.19). Contaminant emissions at the site of the mine shaft were computed by multiplying the annual dust emissions of ore, sub-ore, and overburden (loading truck from skip - Table 3.53) by their respective contaminant concentrations. The three products of the multiplication were then summed to give the values listed in the 4th and 8th data columns of Table 3.54. The health impact of the sources at each location will be assessed separately in Section 6.1.

Annual contaminant emissions due to wind suspension and transport of dust are listed in Table 3.55. These values were computed by multiplying the annual mass emissions (Table 3.53) by the contaminant concentrations in waste rock, sub-ore, and ore listed in Sections 3.4.1.1, 3.4.1.3, and 3.4.1.2, respectively. The uranium and uranium daughter concentrations in dusts from all sources were also multiplied by an activity ratio (dust/source) of 2.5 (Section 3.3.1.2). Although some metals may also be present as secondary deposits, it was believed that there were insufficient data to justify multiplying their concentrations by the 2.5 ratio.

The dust emissions from vehicular traffic listed in Table 3.53 (transportation) were summed with that produced by light vehicular traffic (22 MT/yr) and considered one source of emissions. Concentrations of contaminants in haul road dust have not been measured and are not known. Some spillage of ore and sub-ore along haul roads will undoubtedly raise uranium levels in roadbed dust. As an estimate, uranium and daughter concentrations in the dust were considered to be twice background, 8ppm (2.7 pCi/g), while concentrations of all other contaminants were considered to be similar to those in the waste rock (Section 3.4.1.1). Table 3.56 shows the annual emissions computed with these assumptions.

Table 3.54 Average annual emissions of radionuclides (μCi) and stable elements (kg) from mining activities at the model underground mines

Contaminant	Average Underground Mine ^(a)				Average Large Underground Mine ^(a)			
	Waste Rock	Sub-ore	Ore	Mine	Waste Rock	Sub-ore	Ore	Mine
	Pile Site	Pile Site	Pile Site	Site	Pile Site	Pile Site	Pile Site	Site
Arsenic	0.0004	0.02	0.05	0.04	0.004	0.17	0.60	0.44
Barium	0.012	0.17	0.58	0.44	0.12	1.8	6.4	4.8
Cobalt	NR ^(b)	0.003	0.01	0.007	NR	0.03	0.11	0.08
Copper	0.0007	0.01	0.04	0.03	0.007	0.12	0.43	0.32
Chromium	< 0.002	0.004	0.01	< 0.01	< 0.02	0.04	0.14	< 0.13
Iron	0.24	2.8	9.9	7.5	2.4	3.1	110	82
Mercury	< 0.0003	ND ^(c)	ND	< 0.001	< 0.003	ND	ND	0.005
Potassium	0.28	4.5	16	12	2.8	50	175	129
Magnesium	NR	0.63	2.2	1.6	NR	7.0	25	18
Manganese	0.02	0.17	0.60	0.47	0.19	1.9	6.7	5.1
Molybdenum	0.0001	0.02	0.07	0.05	0.001	0.23	0.81	0.58
Nickel	NR	0.004	0.01	0.009	NR	0.04	0.14	0.10
Lead	0.0009	0.01	0.05	0.04	0.009	0.16	0.55	0.40
Selenium	0.0001	0.02	0.07	0.05	0.001	0.22	0.77	0.55
Strontium	0.006	0.02	0.08	0.07	0.06	0.26	0.91	0.74
Vanadium	0.004	0.25	0.89	0.65	0.04	2.8	9.9	7.1
Zinc	0.0008	0.005	0.02	0.01	0.008	0.06	0.20	0.16
Uranium-238 and each daughter	0.6	45	450	222	6	495	4,990	2,410
Thorium-232 and each daughter	0.04	0.4	6.3	2.8	0.4	4	70	31

(a) Mass emissions from Table 3.53.

(b) NR - Not reported.

(c) ND - Not detected.

Table 3.55 Average annual emissions of radionuclides (μCi) and stable elements (kg) in wind suspended dust at the model underground mines

Contaminant	Average Large Underground Mine			Average Underground Mine		
	Waste Rock Pile	Sub-Ore Pile	Ore Stockpile	Waste Rock Pile	Sub-Ore Pile	Ore Stockpile
Arsenic	0.03	1.9	0.69	0.005	0.34	0.06
Barium	0.87	20	7.4	0.17	3.7	0.66
Cobalt	NR ^(a)	0.35	0.13	NR	0.06	0.01
Copper	0.05	1.3	0.49	0.01	0.24	0.04
Chromium	< 0.15	0.44	0.16	< 0.03	0.08	0.01
Iron	18	345	126	3.4	63	11
Mercury	< 0.02	ND ^(b)	ND	< 0.005	ND	ND
Potassium	21	550	200	4.0	100	18
Magnesium	NR	77	28	NR	14	2.5
Manganese	1.5	21	7.7	0.28	3.8	0.69
Molybdenum	0.008	2.5	0.92	0.001	0.46	0.08
Nickel	NR	0.44	0.16	NR	0.08	0.01
Lead	0.07	1.7	0.62	0.01	0.31	0.06
Selenium	0.006	2.4	0.88	0.001	0.44	0.08
Strontium	0.45	2.9	1.0	0.09	0.52	0.09
Vanadium	0.30	31	11	0.06	5.6	1.0
Zinc	0.06	0.64	0.23	0.01	0.12	0.02
Uranium-238 and each daughter	45	5,450	5,700	9	990	513
Thorium-232 and each daughter	3	44	80	0.6	8	7.2

(a) NR - Not reported.

(b) ND - Not detected.

Table 3.56 Average annual emissions of radionuclides (μCi) and stable elements (kg) from vehicular dust at the model underground mines

Contaminant	Average Large Underground Mine ^(a)	Average Underground Mine ^(b)
Arsenic	0.43	0.22
Barium	14	7.0
Copper	0.86	0.44
Chromium	<2.4	<1.2
Iron	287	145
Mercury	<0.38	<0.19
Potassium	335	170
Manganese	23	12
Molybdenum	0.12	0.06
Lead	1.1	0.53
Selenium	0.10	0.05
Strontium	7.2	3.6
Vanadium	4.8	2.4
Zinc	0.96	0.48
Uranium-238 and each daughter	129	65
Thorium-232 and each daughter	48	24

(a) Mass emissions = 47.8 MT/yr.

(b) Mass emissions = 24.2 MT/yr.

3.5 In Situ Leach Mining

Because in situ leaching of uranium (see general description in Section 1.3.4) is in its infancy, a data base for performing a detailed generic environmental assessment does not presently exist. The fact that the parameters for assessing this process are so site specific and depend upon operational procedures further impedes a generic assessment. Current research projects may help to resolve many of the present uncertainties and provide the data needed to better quantify the potential source terms (La78).

In view of the expected future expansion of this uranium mining method (Section 1.3.4), a qualitative assessment that can be modified later when additional data become available was deemed necessary. This assessment was possible because of recent laboratory experiments and field measurements at pilot-scale plants (Wy77, Ka78b, NRC78b, Tw79).

Similar to other uranium mining methods, in situ leaching also produces liquid, solid, and airborne wastes. However, the quantities of these wastes and their characteristics differ considerably from those produced at surface or underground mines. Also, because the recovery, drying, and packaging of the U_3O_8 produced is often performed at the mine site, wastes from these processes should probably be included in the mine assessment.

This assessment uses the parameters of a hypothetical "typical" commercial-sized in situ solution mine. Unlike surface or underground mines, relatively few in situ facilities exist, and they are all somewhat different because of site specificity and the rapid development of new or modified techniques. The following parameters for the hypothetical mine were based upon those of the Highland, Crownpoint, and Irigaray uranium projects and those reported by Kasper et al. (1978) (Wy77, NRC78b, TVA78b).

The Hypothetical In Situ Solution Mine

- (1) Size of deposit = 52.6 hectares
- (2) Average thickness of ore body = 8 m (Ka78b, NRC78b)
- (3) Average ore grade = 0.06 percent U_3O_8 (Ka78b, Tw79)
- (4) Mineralogy = Sandstone

- (5) Ore density = 2 MT/m³
- (6) Ore body depth = 153 m
- (7) Mine life = 10 years (2-yr leach period in each of 5 sectors)
- (8) Well pattern = 5 spot (NRC78b, TVA78b, Ka78b)
 - Injection wells = 260
 - Production wells = 200
 - Monitoring wells = 80
- (9) Annual U₃O₈ production = 227 MT (Wy77, NRC78b, Ka78b)
- (10) Uranium leaching efficiency = 80 percent (Ka78b)
- (11) Lixiviant = Alkaline
- (12) Lixiviant flow capacity = 2,000ℓ/min (Ka78b, Wy77, NRC78b)
- (13) Lixiviant bleed = 50ℓ/min (2.5 percent) (Wy77, NRC78b, TVA78b)
- (14) Uranium in Lixiviant = 183 mg/ℓ (TVA78b, Ka78b, NRC78b)
- (15) Calcite (CaCO₃) removal required = 2 kg calcite per kg U₃O₈ (Wy77)

The solid, liquid, and airborne wastes generated by this facility are described below. Wastes and quantities generated, as well as operations and procedures selected, will naturally differ to varying degrees from those at some operating sites.

3.5.1 Solid Wastes

The quantity of solid wastes generated depends upon the leachate, the ore body, and operational procedures that effect the mobilization of ore constituents. Little information is available on the quantities of solids generated because of this site dependence, the newness of the process, and the apparent relatively small quantities that are produced. Examples of solid wastes that might be expected to be generated by the alkaline leach process are listed below:

- (1) Materials filtered from the lixiviant line
- (2) Sediments from the surge tanks
- (3) Calcium carbonate from the calcium control unit

- (4) Barium sulfate from the contaminant control in the elution/precipitation circuit of the recovery process
- (5) Materials deposited in the evaporation ponds
- (6) Drill hole residues
- (7) Solids from aquifer restoration

Sources 1 and 2

No information concerning quantities of solids from these two sources could be found in the literature, but they are described as being relatively small compared to other sources (NRC78b). These wastes are transferred to evaporation ponds and retained beneath a liquid seal.

Source 3

One of the larger sources of solids is the calcium control unit (Wy77). Calcite, CaCO_3 , which is removed prior to injection of the refortified lixiviant, coprecipitates radium and any residual uranium. It has been reported that the amount of calcite produced is less than 2.8 kg per 1 kg of U_3O_8 recovered (Wy77). Assuming this ratio to be 2.0, and if Ra-226 is in secular equilibrium with U-238 in the ore, and 2.5 percent is solubilized by the lixiviant (Wy77, NRC78b), 454 MT of calcite will be produced annually and contain a total of 1.6 Ci of Ra-226. Also, calcite has been observed to contain between 1 to 2 percent U_3O_8 by weight (Wy77). Assuming an average of 1.5 percent U_3O_8 , about 1.9 Ci (6.8 MT U_3O_8) of U-238 may also be present in the calcite waste.

Radium-226 and its daughter, Rn-222, are probably the most radiologically significant radionuclides associated with uranium mine wastes, and the small amount of Ra-226 retrieved by in situ leaching is a distinct advantage. Conventionally mining the quantity of ore assumed for the hypothetical in situ mine would contribute 64 Ci of Ra-226 per year to the surface. Because of the insolubility of RaSO_4 , acid lixiviants containing H_2SO_4 mobilize even less radium than alkaline lixiviants. It is reported that the latter mobilizes up to 4.5 times the radium as acid leach solutions (Wy77).

If practical, the calcite waste is transferred to the mill to recover the coprecipitated uranium. Otherwise, the waste is transferred to an evaporation pond and retained beneath a liquid seal to minimize atmospheric dispersion and radon emanation.

Source 4

If necessary, the sulfate concentration in the eluant circuit of the uranium recovery unit may be controlled by the precipitation of BaSO_4 . There are no data on the contaminant levels expected in the BaSO_4 waste, although less than 730 MT per year are anticipated (Wy77). These wastes are impounded beneath a liquid seal of an evaporation pond.

Source 5

An assortment of precipitation compounds will be produced by evaporative concentration of impounded waste solutions. The principal products expected are alkali chlorides, carbonates, and sulfates. The quantity of solids produced by this mechanism and their rate of accumulation on the pond bottom has not been reported.

Source 6

Residues produced from drilling the numerous wells required for in situ leaching constitute another solid waste. The hypothetical in situ leaching facility defined above requires a total of 540 wells drilled to a depth of 153 m: 200 production, 260 injection, and 80 monitoring wells. A diameter of 10.2 cm will be assumed for all wells, although 5.1 cm, 12.7 cm, and 15.2 cm diameter wells have been used (Wy77). To accommodate a concrete and steel casing, a drill hole of approximately 20 cm will be required. The residue from drilling the monitoring wells will consist mostly of barren rock; however, an equivalent of an 8-m section of each injection and recovery well will contain 0.06 percent grade ore. Hence, drill hole residues will consist of 4,960 MT of barren waste rock and 230 MT of ore containing 138 kg of U_3O_8 . These wastes are in relatively small quantities and should be manageable. The waste rock and ore, if mixed and stored in a 2-m-high rectangular pile, would only cover an area of about 0.15 hectares and average 0.0027 percent U_3O_8 .

Source 7

During the active mining period, all solid wastes are generally retained beneath a liquid seal in lined evaporation ponds to minimize atmospheric dispersion and radon emanation. A plan for the final disposal of solid wastes has not been determined. Suggested procedures are to transport the wastes to a conventional uranium mill for further treatment to recover any U_3O_8 present, treat the effluent as mill wastes, construct long-term tailings ponds on the site, or ship the wastes to a licensed off-site burial ground. Solid wastes probably comprise the least significant type waste relative to health and the environment. Solid wastes generated from reclamation procedures will be discussed in Section 3.5.5.

3.5.2 Associated Wastewater

Water flushed through the leached area when restoring the well field is the largest source of wastewater (see Section 3.5.5). The principal sources of wastewater generated by the hypothetical facility during the leaching and recovery operations are as follows:

- (1) Lixiviant bleed -- barren lixiviant removed from the leach circuit to produce a net inflow into the well-field area and to control contaminant concentrations
- (2) Resin wash -- water to wash resin of excess NH_4Cl used to regenerate the resin. Lixiviant bleed is sometimes used for this operation, and it reduces the total quantity of wastewater produced (Ka78b)
- (3) Eluant bleed -- barren eluant removed to control salt accumulation, principally NaCl and Na_2CO_3 , and maintain proper volume
- (4) Well cleaning -- water used to flush injection wells to prevent clogging

The sources of wastewater and the quantities produced vary at different sites, depending upon the lixiviant and recovery circuit chemistry as well as the production rates. However, estimates were made of the quantities of wastewater generated by the four principal sources for the hypothetical in situ facility, and they are listed in Table 3.57. It is assumed that waste from backwashing the sand filters is lixiviant bleed waste water and does not contribute to the total wastewater generated. The total volume of wastewater estimated to be generated is $8.43 \times 10^4 \text{ m}^3/\text{yr}$. Assuming the evaporation ponds are 3.05 m deep with a 0.604 m freeboard (Wy77) and a natural evaporation rate of 142 cm/yr (TVA78b), a pond capacity of $34,770 \text{ m}^3/\text{yr}$ which would encompass a surface area of about 1.4 hectares/yr would be required. Using evaporation data assumed for the Irigaray Uranium Project, about 75 percent of the annual wastewater inventory would evaporate, which would leave $2.11 \times 10^4 \text{ m}^3/\text{yr}$ and require a surface area of 0.85 hectares/yr. If necessary, the pond size can be reduced by using mechanical evaporators.

Table 3.57 Estimated quantities of wastewater produced by an
in situ leaching operation

Source	Flow Rate, (ℓ/min)	Annual Accumulation, (m^3/yr)
Lixiviant bleed (2.5%)	50	2.63×10^4
Resin wash ^(a)	26	1.37×10^4
Eluant bleed	17	8.9×10^3
Well cleaning ^(b)	--	3.54×10^4
Total		8.43×10^4

(a) This may be included in the lixiviant bleed.

(b) Assumes 260 injection wells flushed twice each month with 5680 liters of water.

Source: Data from Wy77 and Ka78b proportioned to an annual U_3O_8 production of 227 MT and a lixiviant flow of 2000 ℓ/min ; aquifer restoration is excluded (Section 3.5.5).

The liquid wastes are generally brines. They contain large amounts of sodium chloride consisting of 1,500 to 5,000 mg/l total dissolved solids (TDS), trace metals ranging from 0 to 10 mg/l, and small quantities of radioactivity. The quantities of contaminants generated each year were estimated for the hypothetical solution mine by using the annual mass emissions estimated for the Highland Uranium Project and adjusting the flow rates to predict the concentrations (NRC78b). Table 3.58 lists these estimated concentrations and annual emissions. Because the contaminants from the lixiviant bleed were not included in the source document, the trace metals that are mobilized by the leachate do not appear in the tabulation, and Ra-226 presence is grossly underestimated (Table 1.7, Section 1.3.4). Considering possible trace metal concentrations and their toxicities, their presence in the lixiviant bleed wastewater may be significant. Assuming that 2.5 percent of the Ra-226 in the ore is extracted, the pregnant leachate will contain about 1,520 pCi/l, yielding 1.6 Ci/yr. However, it is assumed that most of this radium will be removed by the calcium control unit.

There are no planned releases of liquid wastes to the environment at in situ solution mines. The contaminants dissolved in the liquid wastes will accumulate on the pond bottoms as the liquid evaporates. Barring dike failure and seepage through the lined pond bottoms, no impact should be imposed upon the environment by this source during operation.

Another method, other than evaporation, to remove wastewater from an in situ site is deep well injection. This is the dominant method of wastewater removal at operations in South Texas (Durler, D.L., U.S. Steel Corporation, Texas Uranium Operations, Corpus Cristi, TX, 9/79, written communication).

3.5.3 Airborne Emissions

Airborne emissions from an in situ solution mining operation will originate from three principal sources: the uranium recovery and processing unit, the waste storage evaporation ponds, and the radon released from the pregnant leach surge tanks. The primary radioactive species emitted is Rn-222. The nonradioactive species emitted are a function of the lixiviant and the uranium recovery processes employed. Fugitive dust emissions, primarily from vehicular traffic, will also occur on the site. However, because very little heavy equipment is used, the potential for adverse environmental impact from this source will not be significant and is not considered in this assessment.

Table 3.58 Estimated average concentrations and annual accumulation of some contaminants in wastewater

Contaminant	Concentration, mg/l	Annual Accumulation, kg
Calcium	64	5,380
Chlorine	2,070	173,880
Carbonate	31	2,600
Bicarbonate	36	3,020
Magnesium	24	2,020
Sodium	1,320	110,880
Uranium-238	1	84
Radium-226	21 ^(a)	1.8 ^(b)
Thorium-230	6 ^(a)	0.5 ^(b)

(a) Units are pCi/l

(b) Units are mCi.

Note.--Mass emissions estimated for the Highland Uranium Project (NRC78b), adjusted for flow rates and U_3O_8 production of the hypothetical solution mine.

Estimated average annual airborne emissions were computed for the hypothetical facility using data supplied by the Irigaray and Highland Uranium Projects and from the report of Kasper, et al. (1978) (Wy77, NRC78b). Table 3.59 gives the results, proportioned to a production rate of 227 MT/ yr.

The major sources of emissions from the uranium recovery plant are by-products of combustion from the dryers, volatilized solution residuals, and U_3O_8 fines generated during product drying. Carbon dioxide is the major combustion product emitted, although sulfur dioxide may also be significant if oil is used to fuel the dryers. Ammonium salts, used in the precipitation of uranium and resin regeneration, will volatilize as both ammonia and ammonium chloride during yellow cake drying. Airborne particulates that include uranium and some decay products are generated during the drying and packaging processes. The emission rates of U_3O_8 and daughter products were computed on the basis of an average release rate of 363 kg of U_3O_8 per year

Table 3.59 Estimated average annual airborne emissions from the hypothetical in situ leaching facility

Source	Annual Release Rate
<u>Recovery Plant</u> ^(a)	
Uranium-238	1.0×10^{-1} Ci
Uranium-234	1.0×10^{-1} Ci
Uranium-235	4.8×10^{-3} Ci
Thorium-230	1.7×10^{-3} Ci
Radium-226	1.0×10^{-4} Ci
Lead-210	1.0×10^{-4} Ci
Polonium-210	1.0×10^{-4} Ci
Ammonia	3.2×10^0 MT
Ammonium chloride	1.2×10^1 MT
Carbon dioxide	6.8×10^2 MT
<u>Surge Tank</u>	
Radon-222 ^(b)	6.5×10^2 Ci
<u>Storage Ponds</u> ^(c)	
Ammonia	1.0×10^2 MT
Ammonium chloride	3.0×10^2 MT
Carbon dioxide	7.5×10^1 MT

(a) Includes the calcium control unit.

(b) Assumes all radon formed dissolves in the lixiviant and 100 percent is released on contact with the atmosphere.

(c) Based on a release rate of 14.6 MT/yr of NH_3 , 10.6 MT/yr of CO_2 and 42.0 MT/yr of NH_4Cl per hectare of pond surface (Wy77), and an average pond surface area of 7.1 hectares (1.42 ha/yr x 5 yrs).

from a 227 MT/yr facility (Wy77, Ka78b). High efficiency filters and scrubbers are used, which significantly reduce the releases from the uranium recovery plant.

Emission rates from the wastewater storage ponds are determined by the composition of the waste solutions, evaporation rate, feed rate to the ponds, and the water temperature. The principal emissions from storage ponds servicing an alkaline leach process, as defined for the hypothetical facility, are ammonia, ammonium chloride, and carbon dioxide. Different atmospheric releases would result from waste ponds servicing an acid leach facility. The release of Rn-222 from the pond surfaces has not been measured. The emission rate of Rn-222 resulting from the decay of Ra-226 contained in the pond sediments will be inhibited by the liquid seal maintained over the entire surface area of the pond. Because of its low solubility in the unagitated pond water, it is reasonable to conclude that the rate of release for radon from the water surface will be small compared to that from the pregnant leach surge tanks. The liquid seal maintained over the pond area minimizes airborne particulate emissions from the storage ponds.

The principal source of airborne radioactive emissions is the release of Rn-222 from the pregnant leach surge tanks. Rn-222 is mobilized from the ore zone during solution mining and will be largely soluble in the lixiviant under the very high pressure (~15 atm) that exists at the ore zone depth (~500 ft). Upon reaching the atmosphere at the surge tank, nearly complete release of the absorbed radon will take place. Since nearly all Ra-226 remains underground in the leach zone--only 2.5 percent is assumed to be extracted--Rn-222 will continue to be generated in areas leached of uranium.

Consider a 2-year leach period in each of 5 sectors that is 80 percent efficient and yields an average of 227 MT of U_3O_8 per year. If U-238 and Ra-226 are initially in secular equilibrium and 97.5 percent of the Ra-226 remains underground, 156 Ci of Ra-226 will be continually available for Rn-222 production. This quantity of Ra-226 will yield a lixiviant concentration in the 252,800 m³ aquifer (Section 3.5.5) of 6.18×10^5 pCi/l, assuming a maximum emanating power of 100 percent. The latter assumption will result in a maximum Rn-222 concentration in the lixiviant. A high emanating power is probable considering the conditions that exist in the aquifer: high pressure, high permeability due to leaching, the presence of water in the rock pores, radium present on grain surfaces, and the flow rate of water through the ore zone (Ta78, Tanner, A.B., Department of Interior,

Geological Survey, Reston, Va, 11/79, personal communication). Therefore, applying these maximizing conditions with a pumping rate of 2,000 μ /min, 650 Ci/yr of Rn-222 will be released at the pregnant leachate surge tanks.

Apparently very few measurements of Rn-222 concentrations in pregnant leachates have been made at operating facilities. One investigator reports that measured concentrations range from 10,000 pCi/ μ to over 500,000 pCi/ μ and may vary with time at the same well by factors greater than ten (Waligora, S., Eberline Instrument Corp., Albuquerque, N.M., 1979, personal communication). The concentration computed above for the model facility lies above the observed range.

3.5.4 Excursion of Lixiviant

A production zone excursion refers to the event when the leach solution flows from the leach field contaminating the surrounding aquifer. Production zone excursions are usually prevented by bleeding a small fraction (2 to 7 percent) of the lixiviant before reinjection. This imposes an imbalance in the injection-recovery volumes and causes groundwater to flow into the leach field from the surrounding stratum.

Production zone excursions are detected by wells placed 60 m to 300 m from the well field. These wells are routinely monitored, generally bi-weekly, to detect concentration increases of one or more constituents of the lixiviant. Lixiviant constituents monitored may be chloride, ammonia, bicarbonate, sulfate, calcium, or uranium. In addition, conductivity and pH measurements are usually included. When one or more of the indicators exceeds a maximum limit specified in the operator's permit, the observation is verified by resampling. If positive, sampling frequency is increased, appropriate government agencies are notified and corrective actions are begun.

An excursion from the production zone may be terminated by one of the following suggested methods (Wy77):

- (1) Overpumping - increasing the flow rate of the recovery wells to increase the inward flow of native groundwater
- (2) Reordering - applying different pumping rates of the recovery wells to different areas of the well field, providing a greater inflow of native groundwater at specific points

- (a variation of overpumping)
- (3) Reducing Injection - another method of increasing the ratio of recovery flow to injection flow providing the same effect as overpumping
- (4) Ceasing to Pump - stopping both recovery and injection flows (migration is then due entirely to natural groundwater flow, which is many orders of magnitude less than with wells pumping)
- (5) Begin Restoration - initiated when all other efforts have failed to stop the migration of lixiviant from the leach field (Section 3.5.5)

Excursions are likely to occur during the operation of an in situ leach mine. Adverse consequences of an excursion will be determined by its extent, the rate of outward flow, contamination levels, aquifer hydrology, and the effectiveness of corrective measures applied.

3.5.5 Restoration and Reclamation

Restoration is the process by which the in situ leach site is returned to an environmentally acceptable state after mining is complete. Surface restoration consists of removing all structures, pipelines, and so on and sealing the evaporation ponds. Subsurface restoration, the primary area of concern, is done by discontinuing lixiviant injection and continuing pumping to sweep fresh groundwater from the surrounding area through the leached ore zone. It is anticipated that this process will flush out the remaining lixiviant and chemical compounds or elements that have adsorbed or reacted with the mineral content of the aquifer. The water recovered can be purified by chemical precipitation, ion exchange, reverse osmosis, or other processes, and then recycled. This reduces considerably the quantity of water that must be managed. Between 75 and 80 percent of the water can be reinjected while the remainder containing the contaminants is transferred to an evaporation pond (Wy77, NRC78b). During the initial restoration process, it is generally cost effective to recover the uranium from the process wastewater.

Aquifer restoration continues until the groundwater quality in the mining zone meets a criterion established on a basis of the premining water quality. In many cases, the premining groundwater quality criterion is difficult to establish because water quality can vary considerably over the ore zone region and may contain high natural levels of contaminants. Samples of water from wells monitored prior to mining in Texas contained concentrations

of Rn-222 approaching 20,000 pCi/l (Tanner, A.B., Department of Interior, Geological Survey, Reston, Va, 11/79, personal communication), and it is probably unrealistic to attempt to restore an aquifer to a better quality than existed naturally before mining. Wells and flow rates used in this process must be carefully selected and controlled to provide efficient groundwater sweeps and to insure that all affected areas of the leach zone are restored.

The affected aquifer volume that is to be restored may be estimated by the following equation:

$$\text{affected volume} = \text{area of well field} \times \text{aquifer thickness} \times \frac{(\text{porosity})}{100 \text{ percent}} \quad (3.12)$$

Assuming a porosity for sandstone of 30 percent (NRC78b), the affected volume of the hypothetical in situ solution mine defined in Section 3.5 would be:

$$\text{affected volume} = 52.6 \text{ hectares} \times 8 \text{ m} \times \frac{30 \text{ percent}}{100 \text{ percent}} = 1.26 \times 10^6 \text{ m}^3.$$

Because of mixing leach solution with the incoming sweep water and the gradual desorption of some contaminants from clays present in the ore body, more water is required to adequately flush the contaminants than one pore volume. It has been estimated that five to seven pore volumes of water would be required for adequate restoration (Wy77, NRC78b). Using the seven pore volume value and assuming that 80 percent of the sweep water is reinjected after purification, a total of $1.76 \times 10^6 \text{ m}^3$ of wastewater having high TDS would be transferred to the evaporation ponds during the restoration phase. If the aquifer is swept at a flow rate of 2,000 l/min, restoration would take 8 years (1.6 yr per sector), and wastewater will accumulate at about $2.22 \times 10^5 \text{ m}^3/\text{yr}$ during this period. With careful control, restoration can be concurrent with leaching in different areas of the well field.

Table 3.60 lists estimated average concentrations of contaminants in the restoration wastewater (NRC78b) and annual accumulation rates of the contaminants based on a flow rate of 2,000 l/min. In the last column are estimates of the total mass of substances produced by restoration that would become sediments in the evaporation ponds. Data were not provided for calcium, magnesium, chloride, and ammonium ions, even though the latter two are major constituents expected from an alkaline leach process (Wy77). These concentrations reflect average values, but concentrations in the wastewater during the initial phase of the restoration process will be much higher. For

Table 3.60 Estimated average concentrations and annual and total accumulations of some contaminants in restoration wastewater

Contaminant	Concentration mg/l	Annual Accumulation, Kg ^(a)	Total Accumulation, MT ^(b)
Arsenic	0.2	210	1.7
Calcium	NA ^(c)	NA	NA
Chloride	NA	NA	NA
Carbonate	450	473,000	3,780
Bicarbonate	550	578,000	4,620
Magnesium	NA	NA	NA
Sodium	550	578,000	4,620
Ammonium	NA	NA	NA
Selenium	0.10	100	0.8
Sulfate	150	157,000	1,250
Uranium-238	< 1 ^(d)	< 900	< 7.2
Thorium-230	100 ^(e)	0.10 ^(f)	0.8 ^(f)
Radium-226	75 ^(e)	0.08 ^(f)	0.6 ^(f)
Radon-222	618,000 ^(e)	650 ^(f)	5,200 ^(f)

(a) Produced only during the estimated 8-yr restoration period.

(b) Total accumulation during the estimated 8-yr restoration period.

(c) NA - Data not available.

(d) Concentration after uranium extraction.

(e) Units are pCi/l.

(f) Units are Ci/yr or total curies.

Source: Concentrations based on those estimated for the Highland Uranium Project (NRC78b), adjusted for a flow rate of 2,000 l/min.

Table 3.61 A comparison of contaminant concentrations in pre-mining groundwater and pre-restoration mine water (Wy77)

Contaminant	Pre-mining Water, mg/l	Pre-restoration Water, mg/l
Arsenic	<0.0025	0.021
Barium	0.12	0.069
Boron	0.16	0.283
Cadmium	<0.005	0.014
Chromium	0.0135	0.002
Copper	0.019	0.220
Manganese	0.12	0.97
Mercury	0.0028	<0.0002
Nickel	0.018	0.218
Selenium	0.013	1.75
Silver	<0.005	0.015
Zinc	0.003	0.22
Lead	0.0035	0.110
Chloride	10.75	524
Ammonia	<1.0	235
Bicarbonate	139	805
Uranium (U_3O_8)	0.098	24.4
Radium-226	27 ^(a)	371 ^(a)
Total dissolved solids	793	1324

(a) Units are pCi/l.

example, Table 3.61 compares concentrations of substances in the groundwater before mining with those after mining but before restoration. These data are from tests conducted for the Irigaray Project (Wy77) and indicate those substances whose groundwater concentrations may be elevated by in situ leaching.

Radon emission during the restoration process has not been considered (Wy77, NRC78b, Ka78b). Because essentially all Ra-226 remains in the ore zone (about 97.5 percent), it appears reasonable to expect Rn-222 emissions to continue during restoration. A leached-out sector of the model mine will contain 156 Ci of Ra-226 in an aquifer volume of $2.53 \times 10^5 \text{ m}^3$ ($1.26 \times 10^6 \text{ m}^3 + 5$). Although no measurements have been made, it would appear that the restoration wastewater will contain about the same Rn-222 concentration as the pregnant leachate during leaching, $6.18 \times 10^5 \text{ pCi/l}$ (Section 3.5.3). Assuming a pumping rate of 2,000 l/min, a maximum of 650 Ci of Rn-222 will be released during each year of restoration, resulting in a maximum total release of 5,200 Ci during the estimated 8-yr restoration.

Restoration is presently in the experimental stages. No commercial-sized facility has reached that phase of operation. Although restoration by flushing appears feasible, there have been problems when alkaline lixiviants were used, particularly those containing ammonium ions. Ammonium is the preferred cation because sodium causes the clays to swell and plug the formation, and calcium forms an insoluble sulfate that also decreases the permeability of the formation. However, ammonium ions adsorb tightly on to clays by replacing the calcium and magnesium atoms in the clays. Montmorillonite, prevalent in the Texas mining areas, has extensive surface areas that result in very large ion-exchange capacities. Once adsorbed, the ammonium ions desorb at a very slow rate and prolong the restoration. It has been reported that after sweeping a leached ore zone with 10 ore zone volumes of water, the ammonium concentration of the water was reduced to 15 to 25 mg/l (Ka78b). This concentration of ammonium may not be significant, although, under aerobic conditions, ammonium ions can be oxidized to the more toxic nitrate. In a deep aquifer, this oxidation process is not likely to occur, and, because of the very low leachability of ammonium ions from clays, any ammonium retained after restoration will move to surrounding aquifers at a very slow rate.

Several ongoing research studies are trying to solve the ammonium problem (Ka78b). Potassium is being tested as a cation replacement for ammonium

in hopes that its adsorption and swelling characteristics will be favorable. Sweep solutions enriched in calcium and magnesium are being tested to determine if they will facilitate the flushing of the ammonium ion by replacing it on the clays by ion-exchange.

Restoration of the aquifer after mining stops is in the research stage. The adequacy of the restoration process and the procedures required will depend on a number of factors: the lixiviant used, concentration of specific ions in the lixiviant, the physical character of the stratigraphic unit, and the geochemical nature of the ore deposit. Undoubtedly, research will improve the process in the next few years. If the criteria of the restoration process are met, it is unlikely that there will be any adverse environmental impact from a properly restored aquifer.

Generally, the goal of reclaiming the site surface is to return the area to a state similar to that which existed naturally before mining. This often means one suitable for livestock grazing and wildlife habitat. The following site reclamation actions have been proposed (Wy77):

- (1) Remove all structures and exposed pipes and plug all wells with concrete.

- (2) After all impounded liquids have completely evaporated, cover the remains with overburden to a depth [2 m has been suggested at the Irigaray site (Wy77)] that will support plant growth and suppress Rn-222 emissions or transport and deposit the remains in a mill tailings impoundment.

- (3) Before backfilling, dispose of the solids containing sufficient radioactivity to warrant removal by one of the methods suggested in Section 3.5.1.

- (4) Grade surfaces of the backfilled ponds and all other barren areas to create a suitable topography and then revegetate them.

- (5) Irrigate and fertilize sites to develop adequate plant cover.

- (6) Maintain fences to prevent grazing by livestock until stable vegetative cover becomes established.

- (7) Monitor reclaimed sites for radiation, verification of vegetative cover, and the absence of adverse erosion.

- (8) Sample monitoring wells one year after restoration to verify aquifer restoration.

3.6 Other Sources

3.6.1 Mineral Exploration

During early exploration, uranium was identified by its mineral color, i.e., pitchblende from the Central City District in Colorado and carnotite in the Uravan Mineral Belt in Utah and Colorado. It was usually mined in conjunction with other metals and minerals. Later, when portable radiation survey meters became available, a substantial portion of the uranium findings (generally outcrops) were made by non-geologic prospectors (UGS54). Current uranium exploration uses extensive geological studies to locate formations with a strong potential for uranium ore content. These formations are then explored and field surveyed to verify the presence of ore. Much of the current exploratory activity is directed at expanding known deposits and mining areas.

As the surface and near-surface uranium deposits are found, mined, and depleted, exploration for reserves must be conducted at greater depths. The deeper uranium deposits, however, offer few radiometric clues on the surface regarding their location. In these cases, geologic studies and field work postulate the existence of promising geological formations. Actual exploration must be done by drilling. Drilling is also used to extend and explore known uranium producing areas.

There are two categories of drilling: exploratory and developmental. Exploratory drilling is used to sample a promising formation to determine if uranium ore is present. The drilling is generally done on a grid with the drill holes spaced 60 m to 1.6 km or more apart. Development drilling, to define the size and uranium content of the ore body, occurs when ore is struck in an exploratory hole. The development hole spacing ranges from 8 m to 100 m, depending on the characteristics and depth of the ore body. Usually, the same drilling equipment is used for both the exploratory and development drilling.

Ordinarily, there are three vehicles in a drilling unit. One vehicle carries and operates the drill rig, the second carries the drill rods, and the third carries water. Although the drill rig is a well-engineered, compact design, its physical size is increasing to meet the demands of deeper drilling (Personal communication with G. C. Ritter, 1979, Bendix Field Engineering Corp., Grand Junction, CO).

Early drilling (1948-1956) was predominantly done with percussion drills. These drills could drill to depths of about 76 m using 2.8 cm diameter drill steel. The drill bit was cooled and cuttings were removed from the drill hole by forcing air down the center of the drill stem. The cuttings (chips, sands, and dusts) were carried up and out of the drill hole by the air stream with velocities of 914-1520 m per minute (Ni76). The chips and coarse sands collected near the bore hole while the fine sands drifted and deposited around the drill site. Dusts, however, were free to drift with the winds.

Rotary drilling, used for boring deep holes, generally has replaced percussion drilling. Drill stems of 7.3 cm diameter are used to bore holes to depths of about 1300 m. Stems with diameters of 11.4 cm and larger are used for drilling holes in excess of 1300 m. The rotary drill bits are cooled generally in the same manner as percussion drills. When groundwater is encountered, water is used as a drilling medium and for removing cuttings. The cuttings are removed from the drill hole in the form of a slurry or drilling mud. They are usually stored in basins, either fabricated or dug in the ground. If unavailable, water is hauled to the drill site by truck. The drilling muds and water are stored in portable tanks or an earth impoundment for recirculation. After the drilling is completed, very often the cuttings are scattered and the drilling mud left at the site. This practice has been discouraged over the past 10 years in the Uravan area (Personal communication with G.C. Ritter, 1979, Bendix Field Engineering Corp., Grand Junction, CO). In some cases, the cuttings are disposed of in a trench and covered up with earth. Drilling muds are also sometimes covered. In either case, containment of the drilling wastes does not appear to be a prevalent practice.

Development drilling is conducted if ore is struck in an exploratory hole. The offset distance (i.e., the distance between development drill holes) is dependent on the previous history of the ore body sizes in the area. Offsetting may occur as soon as ore is struck, or it may be delayed until the exploratory drilling is completed.

The ore body may be evaluated by bore hole logging or by examining and analyzing cores. Core drilling, if used, usually begins at the top of the ore horizon. Ore (cores and cuttings) removed from the bore hole are sometimes removed from the drill site. In cases where the ore is not removed from the drill site, it remains with the dry cuttings or in the drilling muds. The drill hole collar is sometimes plugged with 0.9 - 1.5 m of concrete after the bore hole has been evaluated. In some states, the drill hole must be plugged to seal off aquifers in order to minimize groundwater contamination.

3.6.1.1 Environmental Considerations

By 1977, the uranium industry had completed 101×10^6 meters of surface drilling, with an all-time yearly high of 12×10^6 meters (DOE79). From 1958-1977, about 821,900 surface holes were drilled, resulting in 87.8×10^6 meters of bore holes. No statistics are available on the number of holes drilled from 1948-1958, but the annual and cumulative meters drilled for that period is known (DOE79). In order to estimate the number of drill rig placements for that period, the total annual meters of drilling was divided by the annual average bore hole depth. The average depth per bore hole was estimated by plotting the average annual bore hole depths for 1958-1977 then using that data to estimate the annual bore hole depths for 1948-1957 by linear regression analysis (Fig. 3.22).

The data points in Fig. 3.22 appear to fall into two groups: 1958-1966 and 1966-1977. The average drilling depth of the 1966-1977 group of data points probably reflects the deep drilling in the Grants, New Mexico area that became significant in 1969. Using this information, the 1948-1958 average drilling depths were estimated from regression analysis using the 1958-1966 data points only. Table 3.62 is a summary of the DOE drilling data and the number of estimated bore holes by type and year.

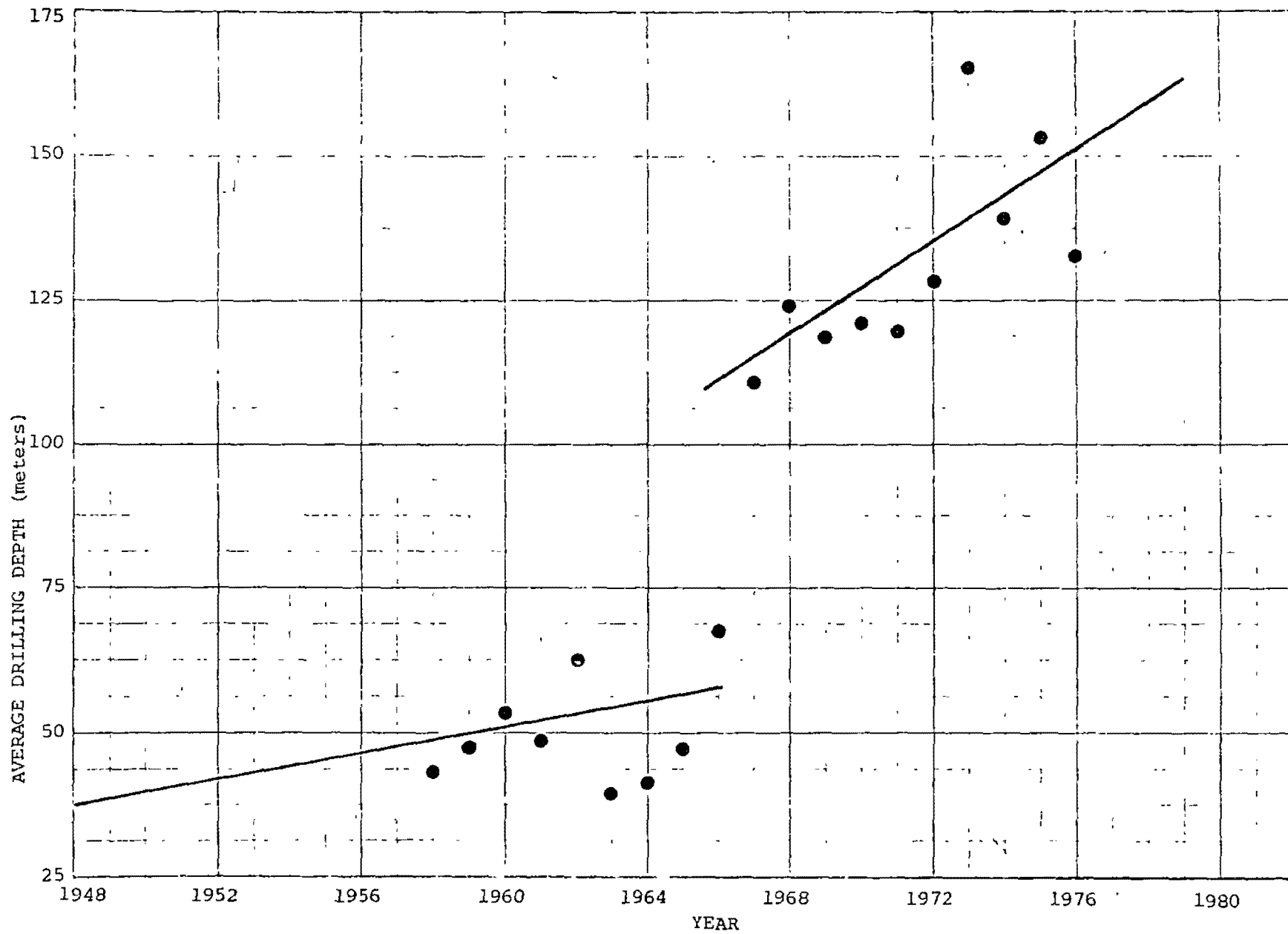


Figure 3.22 Average depth of exploratory drilling in the U.S. uranium industry from 1948 to present.

Table 3.62 Estimates of exploratory and development drill holes (1948-1979)

Year	Surface Drilling (10^6 Meters)		Average Hole Depth(Meters)	Number of Holes	
	Exploration	Development		Exploration	Development
1948	0.052	0.012	38.1 ^(a)	1,360 ^(b)	320 ^(b)
1949	0.110	0.016	38.1 ^(a)	2,880 ^(b)	424 ^(b)
1950	0.174	0.063	39.6 ^(a)	4,380 ^(b)	1,600 ^(b)
1951	0.329	0.106	41.1 ^(a)	8,000 ^(b)	2,580 ^(b)
1952	0.415	0.091	41.1 ^(a)	10,100 ^(b)	2,220 ^(b)
1953	1.11	0.112	42.7 ^(a)	26,100 ^(b)	2,620 ^(b)
1954	1.24	0.169	42.7 ^(a)	29,000 ^(b)	3,950 ^(b)
1955	1.61	0.232	44.2 ^(a)	36,300 ^(b)	5,260 ^(b)
1956	2.22	0.457	45.7 ^(a)	48,600 ^(b)	10,000 ^(b)
1957	2.24	0.564	45.7 ^(a)	49,000 ^(b)	12,300 ^(b)
1958	1.15	1.06	45.7	25,300	22,900
1959	0.722	1.00	48.2	16,300	19,600
1960	0.427	1.28	53.9	7,340	24,400
1961	0.402	0.972	50.0	8,260	19,300
1962	0.451	0.741	61.9	6,440	12,900
1963	0.268	0.604	39.6	8,470	13,500
1964	0.294	0.381	42.4	5,970	9,910
1965	0.354	0.289	47.5	6,230	7,330
1966	0.549	0.731	67.7	5,750	13,200
1967	1.67	1.62	110	12,800	16,900
1968	4.97	2.30	125	38,500	19,500
1969	6.25	2.86	120	47,900	28,000
1970	5.49	1.69	122	44,000	14,900
1971	3.47	1.23	121	28,400	10,400
1972	3.60	1.10	128	26,900	9,710
1973	3.29	1.70	146	22,600	11,700
1974	4.88	1.83	168	27,400	12,300
1975	5.03	2.74	139	34,300	21,600
1976	5.94	4.48	154	40,400	27,200
1977	7.89	4.45	132	62,600	30,900
1978	10.8	5.24	155 ^(a)	69,200 ^(b)	33,700 ^(b)
1979	9.94	5.18	158 ^(a)	62,700 ^(b)	32,700 ^(b)
TOTAL	286	149		823,000	454,000

(a) Indicates estimated average depth from Fig. 3.22.

(b) Indicates number of drill holes estimated by dividing the annual exploration and surface drilling depths by the average hole depth.

Cuttings produced by drilling can degrade the drill site area and the local air quality. For convenience of evaluation, the cuttings are divided into two general categories--dusts and wastes. The dusts are drilling fines that become airborne, and wastes are drilling chips and sands deposited around the borehole. The maximum dust production occurs when compressed air is used solely for cleaning the boreholes. Generally the drilling industry uses foaming agents injected into the compressed air stream to help remove drill cuttings. The foam traps and contains the fine particulates and substantially reduces the airborne dust. In practice, the drillers minimize airborne dust, because it causes excessive wear on engines and compressors. Dust production also indicates improper drilling energy being used to grind up cuttings in the borehole rather than bore. Occasionally some water may also be injected into the air stream to remove cuttings and to keep the drill hole from collapsing when loose materials are encountered.

There are some estimates of airborne dust production and general assumptions concerning drilling practices (Private communication with Mr. T. Price, Bendix Corp., Grand Junction, CO and E. Borgerding, Borgerding Drilling Co. Inc., Montrose, CO). They are as follows:

(1) The ratio by weight of the chips, sands, and dusts produced by drilling is approximately 60:37:3, respectively (i.e., 3 Kg of every 100 Kg of cuttings removed from a borehole is available as airborne dust).

(2) Fifty percent of all drill holes are wet (mud) drilled and 50 percent are air drilled; ninety-five percent of the latter are drilled using mist or foam (i.e., 2.5 percent are dry-drilled).

(3) The first 6.6 m of all drill holes are drilled dry (i.e., no mist or foam is used).

We estimated dust production from contemporary drilling by averaging drilling data from Table 3.62 for the years 1975 through 1979. The average depth of the holes for this period is 148 m. The annual average numbers of exploration and development holes are 53,800 and 29,200, respectively. Airborne dust production from those holes that are drilled with mud (wet), foam, or mists (97.5 percent of both the exploratory and development holes) will originate only from the first 6.6 m depth. The weight of dust generated per hole will be as follows:

$$\text{Airborne dust (kg)} = \text{Volume of borehole (m}^3\text{)} \times \text{density (kg/m}^3\text{)} \times \text{airborne dust fraction (.03) per drill hole}$$

$$\begin{aligned}
 &= (\pi r^2 h) \left(\frac{2000 \text{ kg}}{\text{m}^3} \right) (0.03) \quad \text{where } h = 6.6 \text{ m} \\
 &\quad \quad \quad r = 0.0865 \text{ m (assumed average radius of 2 bit sizes } r = 7.3 \text{ cm and } 10 \text{ cm) (Pe79)} \\
 &= (3.14)(7.48 \times 10^{-3}) \text{ m}^2 \times 6.6 \text{ m} \times 2000 \frac{\text{kg}}{\text{m}^3} \times 0.03 \\
 &= 9.3 \text{ kg}
 \end{aligned}$$

The average weight of airborne dust (kg) produced from all contemporary annual drilling (first 6.6 m) is

$$83,000 \text{ drill holes} \times \frac{9.3 \text{ kg}}{\text{drill hole}} = 7.7 \times 10^5 \text{ kg.}$$

The annual total weight (kg) of airborne dust produced from 2.5 percent of the annual number of drill holes bored (dry) where no mud, mists, or foams are used

$$= 83,000 \text{ drill holes} \times \frac{148 \text{ m}}{\text{drill hole}} \times 0.025 \times 47 \frac{\text{kg}}{\text{m}} \text{ cuttings} \times$$

$$0.03 \text{ kg dust/kg cuttings} = 4.3 \times 10^5 \text{ kg/yr.} \quad (3.13)$$

The total weight of airborne dust produced annually from each dry-drilled borehole is 209 kg.

Assuming that each development hole penetrates the 3.6 m ore body, the total amount of airborne ore and sub-ore dust produced from development drilling annually is

$$29,200 \frac{\text{drill holes}}{\text{yr}} \times \frac{3.6 \text{ m (ore and sub-ore)}}{\text{drill hole}} \times \frac{47 \text{ kg cuttings}}{\text{m}} \times$$

$$0.03 \text{ kg dust/kg cutting} \times 0.025 = 3.7 \times 10^3 \text{ kg.} \quad (3.14)$$

The total weight of airborne ore and sub-ore dust produced from each development drill hole (no mud, mists, or foams used) is 5.1 kg.

The estimated annual quantity of ore and sub-ore brought to the surface by contemporary drilling equals:

$$\begin{aligned}
 &29,200 \frac{\text{drill holes}}{\text{yr}} \times \frac{3.6 \text{ m}}{\text{drill hole}} \times \frac{47 \text{ kg cuttings}}{\text{m}} \quad (3.15) \\
 &= 4.9 \times 10^6 \frac{\text{kg}}{\text{yr}} \text{ or } 4.9 \times 10^3 \frac{\text{MT}}{\text{yr}}
 \end{aligned}$$

Most of the ore will remain at the drill site with drilling muds or with the drilling wastes around the drill holes. Since the ore most usually will be the last material removed from the boreholes, it will be deposited on the

surface of the cuttings and drilling muds. This will expose the ore to the elements and subject it to erosion.

3.6.1.2 Radon Losses from Drill Holes

When the development drill penetrates an ore body, some of the ore and sub-ore bearing formations will be exposed to air in the drill hole. Some of the radon gas produced in the ore can enter into the air in the drill hole and escape to the atmosphere. The mechanisms affecting the release rate of radon from boreholes are poorly understood. Tanner observed a wide variation in radon concentrations as a function of depth in an open borehole as compared to a closed borehole (Ta58). Tanner also noted that strong winds could significantly reduce the total radon content of an uncovered borehole. Since so little is known about radon discharges from development boreholes, radon losses in this report are assessed on a "worst case" basis using the following assumptions:

1. The drill hole is not plugged.
2. About 3.6 m of ore and sub-ore were drilled.
3. All radon released into the borehole escapes to the atmosphere.
4. The average grade of the ore and sub-ore is 0.17 percent.
5. No water accumulates in the borehole.

The surface area of the borehole passing through the ore and sub-ore body is

$$2 \pi r h = 2 \times 3.14 \times 0.0865 \text{ m} \times 3.6 \text{ m} = 2.0 \text{ m}^2. \quad (3.16)$$

The radon release rate is estimated for ore and sub-ore in the borehole using an exhalation rate of 0.092 Ci/m^2 per year per percent of U_3O_8 (Ni79). The quantity of radon (Q) per development hole escaping per unit time is

$$\frac{0.092 \text{ Ci}}{\text{m}^2 \text{ yr \%}} \times 0.17\% \times 2.0 \text{ m}^2 \times \frac{1}{3.15 \times 10^7 \text{ sec/yr}} \times 10^{12} \frac{\text{pCi}}{\text{Ci}} = 990 \text{ pCi/sec} \quad (3.17)$$

The total quantity of radon per annum escaping from all development holes drilled through 1979

$$\begin{aligned}
 &= 4.5 \times 10^5 \text{ drill holes} \times \frac{990 \text{ pCi}}{\text{sec-drill hole}} \times 3.15 \times 10^7 \text{ sec/yr} \\
 &\quad \times \frac{1}{10^{12} \frac{\text{pCi}}{\text{Ci}}} \\
 &= 14,000 \text{ Ci/yr} \quad (3.18)
 \end{aligned}$$

The "worst case" estimate can be modified by assuming 50 percent of the holes are wet and 30 percent of the remaining holes are plugged or have collapsed. In this case, the total source term would be about 4,900 Ci/yr. Since about 31 percent of the development drill holes are at surface mines and are consumed by the pits, the annual Rn-222 release from the remaining holes will be 3,400 Ci/yr.

3.6.1.3 Groundwater

Progressively deeper holes are being drilled as the ore bodies near the surface become depleted. As the drilling depths increase, one or more aquifers may be intercepted by a drill hole, and an aquifer with poor water quality may be connected with an aquifer with good water quality. Depending on the direction of flow, the quality of water may be downgraded in a good aquifer. Most states require some plugging of the drill holes to seal the aquifer in order to maintain water quality. Adequate plugging of the drill holes requires a conscientious effort on the part of the driller and the regulatory agency. Since the movement of groundwater is relatively slow, the change in the quality of water in an aquifer will not be apparent for some time. Thus, it may take a long time to correct the quality of water in a downgraded aquifer.

3.6.1.4 Fumes

It is estimated (Pe79) that 11.2 liters of diesel fuel are needed to drill 1.0 m. In 1979, the average borehole depth was estimated to be 158 m and would require about 1770 liters of diesel fuel. This fuel would be burned at a rate of approximately 173 liters per hour. Some individual holes, however, are drilled in excess of 914 m and require 10,200 liters of diesel fuel. It is estimated that about 170 million liters of diesel fuel were consumed for all 1979 drilling.

The principal emissions from the drilling power sources are particulates: sulfur oxides, carbon monoxide, nitrogen oxides, and hydrocarbons. Because of the transient nature of the drilling, these releases are not expected to substantially lower air quality over time.

3.6.1.5 Model Drilling

About 1.3×10^6 holes have been drilled and bored for all uranium mining from 1948 through 1979 for approximately 3000 mines. This would amount to about 430 holes per mine. Thirty-six percent of the holes were for development drilling, and 64 percent were for exploratory drilling. Assuming that 50 percent of the exploratory and development holes are air drilled (see Section 3.6.1.1), the airborne dust production for an average mine may be estimated as follows:

$$\begin{aligned} \text{Airborne dust from all drill holes (first 6.6 m of depth air drilled dry)} \\ = 430 \text{ drill holes} \times \frac{9.3 \text{ kg}}{\text{drill hole}} = 4000 \text{ kg.} \end{aligned} \quad (3.19)$$

$$\begin{aligned} \text{Airborne dust from all dry air drilling, less the first 6.6 m,} \\ = (430 \text{ drill holes} \times \frac{209 \text{ kg dust}}{\text{drill hole}} \times 0.5 \times 0.05) - 100 \text{ kg} = 2100 \text{ kg.} \end{aligned} \quad (3.20)$$

$$\begin{aligned} \text{Airborne ore and sub-ore dust produced by dry air drilling} \\ = 430 \text{ drill holes} \times 0.36 \times 0.5 \times 0.05 \times \frac{5.1 \text{ kg dust}}{\text{drill hole}} = 20 \text{ kg.} \end{aligned} \quad (3.21)$$

$$\begin{aligned} \text{Total airborne dust produced from all drilling at an average mine site} \\ = 4000 \text{ kg} + 2100 \text{ kg} = 6100 \text{ kg} = 6.1 \text{ MT.} \end{aligned} \quad (3.22)$$

Twenty kilograms of the total dust produced will be ore and sub-ore dusts. The Rn-222 emissions from the bore holes at an average mine site would be

$$430 \text{ drill holes} (0.5)(0.36) \frac{(990 \text{ pCi})}{\text{sec-drill hole}} = 7.7 \times 10^4 \frac{\text{pCi}}{\text{sec}}, \quad (3.23)$$

or 2.4 Ci/yr.

Development drill holes at a surface mine would be consumed by the pit.

Tables 3.63--3.65 show airborne particulate source terms for uranium drilling for individual drill holes and for an average uranium mine. Table 3.63 lists the airborne dust produced for each type exploratory and development borehole; Table 3.64 summarizes the quantity of airborne dust produced by all types of drilling at an average mine site; and Table 3.65 lists the pollutants emitted from a drill rig power source.

3.6.2 Precipitation Runoff from Uranium Mines

Unquestionably, overland flow or surface runoff from precipitation transports dissolved and suspended contaminants from mining areas to the offsite environment. Unfortunately, the significance of this pathway rela-

Table 3.63 Estimated source terms per borehole for contemporary surface drilling for uranium

Type of Drilling	Thickness of Ore and Sub-ore Bodies (m)	Airborne Dust Production		Airborne Ore and Sub-Ore Dust Production	
		Total(kg)	Rate(kg/min) ^(a)	Total(kg)	Rate(kg/min) ^(a)
<u>Exploratory</u>					
Air (dry)	NA ^(b)	209	0.27	NA	NA
Air (mist or foam)	NA	9.3	0.27	NA	NA
Wet (mud)	NA	9.3	0.27	NA	NA
<u>Development</u>					
Air (dry)	3.6	209	0.27	5.1	0.27
Air (mist or foam)	3.6	9.3	0.27	NA	NA
Wet (mud)	3.6	9.3	0.27	NA	NA

(a) Based on an air drilling rate of 11.5 m/hr.

(b) NA - not applicable.

Table 3.64 Airborne dusts produced at an average mine site
from exploratory and development drilling

Type of Drilling	Quantity of Airborne Dust (kg)
All types (first 6.6 m depth)	4,000
Air drilling (dry)	<u>2,100</u>
Total	6,100 kg ^(a)

^(a) Twenty kg of the total will be ore and sub-ore dusts.

Table 3.65 Estimates of emissions from drill rig
diesel power source

Pollutant	Production Rate (kg/10 ³ liters fuel)	Quantity ^(a) (kg/drill hole)	Rate ^(a) (kg/hr)
Carbon monoxide	12.2	20.2	1.5
Hydrocarbons	4.49	7.4	0.55
Nitrogen oxides	56.2	93	6.9
Aldehydes	0.84	1.39	0.10
Sulfur oxides	3.74	6.2	0.46
Particulates	4.01	6.6	0.49

^(a) Based on a drilling rate of 11m/hr.

Source: EPA77b.

tive to uranium mines is highly site specific and poorly understood. Very few field studies of runoff from uranium mining areas have been conducted, and what field data do exist frequently relate to the combined and probably greater influences of mine water discharge and milling. Most of the NRC regulations apply to mill operations, since mining is generally exempt from the agency's charter. The EPA regulations (Environmental Radiation Protection Standards for Nuclear Power Operations; 40 CFR Part 190) applicable to the uranium fuel cycle establish dose limits for individuals to provide protection for populations living in the vicinity of uranium mills. Uranium mines are excluded, and so are liquid effluent guidelines for ore mining and dressing (40 CFR 440, Subpart E). Regulations being developed under the Resource Conservation and Recovery Act (RCRA) of 1976 apply to radioactive wastes not covered by the Atomic Energy Act of 1954, as amended. Solid and liquid waste categories will be defined in forthcoming EPA regulations developed under RCRA, but it is not anticipated that runoff from mined lands will meet the waste characteristics in the regulations. Similarly, the Federal Water Pollution Control Act Amendment of 1972, the Clean Water Act of 1977, the Safe Drinking Water Act, and State regulations in general do not address surface runoff effects of mining. Without the regulatory base, studies and field data are, not surprisingly, rather scarce. In New Mexico, the State's 208 Water Quality Management Plan calls for, among other things, improved data collection on runoff from active and inactive tailings piles and from drilling, exploration, and development activities such as access road and drill site construction (So79).

We have not estimated chemical transport by overland flow because of the limited time for the study. But, it is reasonable to expect that such transport may be quite significant in an arid and semiarid climate where much of the precipitation that does infiltrate is discharged back into the atmosphere as water vapor. This has been well demonstrated in the case of uranium mill tailings (K178). Water moving back out of the soil transports dissolved salts that are deposited on the soil surface when the carrier (water) evaporates. Subsequent precipitation further transports these salts downward into the soil and laterally to offsite areas. So-called "blooms" of salt crystals, composed mainly of sulfate and chloride compounds, characterize uranium ore bodies, mill tailings piles, and mine wastes in a number of Western States, and we must presume that such salts solubilize in runoff.

This also indicates that there may be large concentrations of contaminants available for plant uptake. Molybdenum, in particular, is one of the toxic elements on such blooms, and uranium is also highly suspect. Selenium, arsenic, and vanadium may also be present, since their anions are mobile under oxidizing conditions characteristic of the near-surface, unsaturated zone (Fu78).

Overburden has been used extensively to backfill surface mines operating since the early to mid 1970's, but this is not true at many if not most older and now inactive mines. Erosion of these piles by water and wind may present the greatest problem (Ka75). Using overburden to construct access roads and dikes distributes contaminants in the local environment and may aggravate air and water pollution. Considering that 75 percent of the overburden has a grain size exceeding 2000 μm (see Table 3.12), it is unlikely that widespread physical transport will result from overburden piles. However, using overburden for roads decreases the grain size. The association of uranium and progeny with the smaller sediment-size fractions, by a factor of 2.5, increases the potential for transport by overland flow.

Tables 3.15, 3.16, and 3.19 show stable and radioactive trace elements in ores, sub-ore, and overburden from uranium mines. Understandably, uranium, thorium, and radium are high. Arsenic, selenium, vanadium, and molybdenum are almost always closely associated with uranium. Barium, zinc, manganese, copper, iron, and potassium may also be associated in certain mineral provinces and districts. Mercury and cadmium are occasionally present (Th78). There is no consistent relationship between ore grade and trace metal content in selected New Mexico and Wyoming study areas (Wo79).

Particularly in the case of active or recently active mines, surface runoff is collected with dikes and ditches that route water to settling ponds. Water spray or chemical additives can control road dust. They are commonly used in the active mining stage, but almost never used during exploratory drilling. Grading piles to a slope of 3:1 or less also helps to reduce runoff (St78), and this practice is becoming common in Texas and Wyoming. Proper planting techniques further reduce runoff by increasing infiltration and decreasing sediment transport.

The significance of surface runoff from mining areas as a dispersal mechanism was investigated as part of this study (Wo79) (see also Section 3.2.3.2). We examined stable and radioactive trace elements in soils

affected by runoff from ore, sub-ore, and mine waste/overburden piles from one active surface mining area in Wyoming and two inactive areas (surface and underground mines) in New Mexico. Although there was evidence of offsite movement of uranium and radium at all sites, transport is limited and decreases with distance from the site. In Wyoming, pollutant releases from the mine studied do not reach nearby water courses although onsite transport of stockpiled ore as a result of precipitation runoff does occur.

A U.S. Bureau of Mines (BOM, no date) study of strip and surface mining operations and their effects in the United States involved questionnaires, literature survey, and onsite examinations of 693 selected sites, among which were uranium mines in New Mexico and Wyoming. At 60 percent of the sites, on a national basis, there were no serious problems because vegetation was reestablished and the slope of the land was gentle both before and after mining. Thirty percent of the sites had eroded to depths of 0.3 m or less, and the remainder were gullied to greater depths. There were sediments from mined lands in 56 percent of the ponds and 52 percent of the streams on or adjacent to the sample sites. Spoil bank materials ranged in pH from 3 to 5 at 47 percent of the sites and are thus not amenable to plant growth. Field observations substantiate that mined land areas, be they former forests or grasslands, did not return to the pre-mining condition. Idle land increased almost fourfold because of mining. The study concluded that natural processes need to be strongly supplemented if mined sites are to revert to former uses. Since only 6.3 percent of lands mined for uranium were reclaimed from 1930 through 1971 (Pa74), it seems reasonable to conclude that there are increased sediment loads, gullyng, and poor revegetation at most older inactive mines that were poorly stabilized, if at all.

The Bureau of Mines study concluded that peak sediment loads in runoff are characteristic of areas with high intensity storms and steep slopes, particularly during and shortly after mining. Such problems are less severe in arid regions, but large quantities of sediment are discharged from mine workings, spoil heaps, and access roads. In some instances, effects of wind and water erosion on steep spoil banks in arid lands are evident many years after abandonment. In areas outside Appalachia, 86 percent of the areas investigated had sufficient runoff control, and those areas where there was a problem almost exclusively involved coal, phosphate, manganese, clay, and gold.

Incidences of radioactive contamination of local surface water have been documented for the Shirley Basin uranium mine (Utah International, Inc.) in Wyoming (Ha78). The most pronounced changes in water and stream sediment quality coincided with initial strip mining and mill processing operations. Early acid-leach solution mining also had a decided impact. Pollutant loadings from overland flow, per se, were not determined but are presumed to be minor compared to aqueous discharges from mines and mills. These findings contradict those of an earlier study (Wh76) of the same mine. Soil and vegetation collected from 1971 through 1975 at 28 stations in the vicinity of the mine were analyzed for gross alpha and gross beta (1971 to 1974) and total uranium, Ra-226 and Pb-210 (1975). The study (Wh76) concluded that--

1. concentrations of the foregoing parameters were extremely variable but reasonably consistent with previously reported information;
2. there is no evidence that radionuclide concentration in soil or vegetation collected from routine monitoring stations are changing with time;
3. concentrations of radioactivity in soil and vegetation correlate with distance from the mill area to a distance of 1.2 miles; and
4. measurable ecological effects from radiation in the environs of the Shirley Basin mine cannot be demonstrated.

The absence of statistically significant soil and vegetation contamination from the mine versus the mill is noteworthy. Overall, vegetation tends toward higher alpha and beta concentrations than soil, except at the close-in, upwind sampling areas. This selective concentration in vegetation suggests aerial deposition of contaminated dust particles on vegetation, with some additional possibility for root uptake.

Estimates of surface drilling for uranium reveal that relatively large land areas are involved. The volume of cuttings removed from borings in the period 1948 through 1979 is calculated using 286×10^6 m of exploratory drilling and 104×10^6 m of development drilling (from Table 3.62). We assumed that 30 percent of the mines are surface mines, which eliminates the borings and related debris. Thus the value of 149×10^6 (in Table 3.62) is reduced by 30 percent. Average diameter for 8.5×10^6 m of borings in the period 1948 through 1956 is 2.8 cm versus 7.3 cm for the period 1957 through 1979 (see Section 3.6.1) when 426×10^6 m of drilling took place. A sample calculation for the volume removed from borings made in the period 1975-1979 follows:

$$\begin{aligned}
 V &= \pi r^2 h & (3.24) \\
 &= (3.14) \left(\frac{7.43 \text{ cm}}{2} \right)^2 (146\text{m}) \\
 &= 0.632 \text{ m}^3
 \end{aligned}$$

Assuming a bulk density of 2000 Kg per m^3 , each boring results in 1265 Kg of cuttings at land surface. There were 415,300 borings, resulting in 263,000 m^3 of cuttings. Assuming that the average thickness of cuttings is 0.5 m, 526,000 m^2 or 0.53 Km^2 is affected. The inclusive area affected by drilling from 1948 through 1979 is 3.6 Km^2 .

Table 3.66 summarizes the surface areas affected by mine wastes, ore piles, and exploration and development activities. Maximum use was made of data developed elsewhere in this report on the number of mines, waste pile dimensions and surface areas, and the summary of exploration and development. The estimate is, at best, a first approximation and needs considerable refinement.

For example, grain size, degree of consolidation, slope, vegetative cover, and other characteristics may vary considerably between ambient soil and rock materials versus mine wastes. The latter very often occur in steep, unvegetated piles and are composed of easily-eroded, friable sandstone, boulders, and fines. It is likely, therefore, that the sediment yield on a mass per time per area basis exceeds that of the surrounding areas; thus the estimate developed below may well be on the low side.

Sediment yields from areas affected by various mining operations are roughly estimated from consideration of land areas affected and unit soil loss values for the surrounding regions. Actual values for individual tailings or waste piles may be considerably different, but refining the values given will require additional analysis beyond the scope of the present study.

Potential coal mining lands in the Northeastern Wyoming range lose soil at rates of 4.8 to 167 $\text{m}^3/\text{Km}^2/\text{yr}$ (Ke76). Upland erosion and stream channel erosion in the Gillette study area are not generally serious problems, since land dissection is presently minimal and vegetative cover is well established. The potential for increased sediment yield is large, if vegetative cover were to be reduced or eliminated and slopes steepened because of mining. Certainly, during active mining, these conditions will be at least locally present. Erosion rates of 600 to 1,100 $\text{m}^3/\text{Km}^2/\text{yr}$ from mined lands in the South Powder River Basin are expected, and they are reasonably close to

Table 3.66 Sediment yields in overland flow from uranium mining areas

Source Term	Factor	No. Installations	Cumulative Source, Km ²	Annual Sediment Loading, m ^{3(a)}
<u>Active Mines</u>				
Underground				
Ore piles	603 m ² /mine	251 mines	0.15	143
Sub-ore piles	26,700 m ² /mine	251 mines	6.7	6385
Waste rock piles	26,700 m ² /mine	251 mines	6.7	6385
Surface				
Ore piles	4.15 x 10 ³ m ² /mine	36 mines	0.15	143
Sub-ore piles	67 x 10 ³ m ² /mine	36 mines	2.4	2287
Overburden piles	380 x 10 ³ m ² /mine	36 mines	13.7	13056
<u>Inactive Mines</u>				
Underground				
Waste piles and sub-ore	4.07 x 10 ³ m ² /mine	2108 mines	0.86	820
Surface				
Overburden and sub-ore	6.73 x 10 ⁴ m ² /mine	944 mines	64	61000

Table 3.66 (Continued)

Source Term	Factor	No. Installations	Cumulative Source, Km ²	Annual Sediment Loading, m ^{3(a)}
<u>Exploration and Development</u>				
Drilling				
1948-1979	435 x 10 ⁶ m	1.28 x 10 ⁶ borings	3.6	3431
1975-1979	1265 kg/boring	415,300	0.53	506
Access roads and pads				
	1.25 acres or 0.5 ha/boring	1.28 x 10 ⁶	6500	6.2 x 10 ⁶

(a) Assumes average sediment yield of 953 m³/Km².

Note.--Data in this table are based on average mine vs. average large mine as defined in Section 3 of report.

natural, pre-mining conditions (R. Loeper, Soil Conservation Service, 1979, personal communication). At the Bear Creek mine, the reclamation design calls for maximum losses from overburden piles of $1,100 \text{ m}^3/\text{Km}^2/\text{yr}$ initially and $600 \text{ m}^3/\text{Km}^2/\text{yr}$ after the first 3 years. In general, erosion and soil loss from uranium mining in this part of Wyoming is not a significant problem, mainly because of reclamation by industry. Sediment yields in the Grants Mineral Belt range from 95 to $240 \text{ m}^3/\text{Km}^2/\text{yr}$ in the area of the large Jackpile-Paguate surface mine to 500 to $1,400 \text{ m}^3/\text{Km}^2/\text{yr}$ near the underground mining centers around Smith Lake, Ambrosia Lake, and Churchrock (P. Boden, Soil Conservation Service, 1979, personal communication). Considering both the Wyoming and New Mexico model mine areas, this study used an overall average annual soil loss rate of $953 \text{ m}^3/\text{Km}^2$. This average sediment yield rate is based on studies by the Soil Conservation Service of large areas in New Mexico, Wyoming, and other Western States.

In summary, the total land area directly affected by uranium mining is about 6600 Km^2 . Assuming an overall average sediment yield of $953 \text{ m}^3/\text{Km}^2$, annual sediment transported by overland flow is approximately $6.3 \times 10^6 \text{ m}^3$. Obviously exploration and development activities affect the greatest area (6500 Km^2), but they do not necessarily have the greatest impact. Exploration and development, for example, affect large areas, but most of the area affected is a result of constructing access roads and drill pads. Whereas sediment yields from ore, sub-ore, overburden, and waste rock is estimated at $90,000 \text{ m}^3$ per year. Surface mining, although it supplies only about 30 percent of U.S. production, affects the second greatest area (80 Km^2). We have not attempted to characterize the quality of sediment runoff. The fate of these sediments is very poorly understood and has not been the subject of intensive investigation. Further study in the area of intensive surface mining such as in Texas and Wyoming is needed to determine changes in erosion rates resulting from mining and to quantify the contaminant flux and fate.

3.7 Inactive Mines

3.7.1 Inactive Surface Mines

For generic purposes, a model inactive open pit or surface uranium mine must be defined in order to estimate the environmental impact from this type

of mining. We have assumed that an inactive surface mine has a single hole or pit in the ground, with all of the materials (wastes) stacked into piles adjacent to the pit area. The size or volume of the pit would be approximately equal to the volume of the ore and wastes removed from it. Since only 6.3 percent of all of the land used for uranium mining has been reclaimed from 1930 through 1971 (Pa74), no credit for reclamation is given to the model mine.

Ideally, the model mine size could be established by averaging the ore and waste production for each inactive surface mine. Unfortunately, these statistics are either not thoroughly documented or they are retained as company confidential information. In lieu of specific information, the model surface mine size was established from annual ore and waste production statistics for all surface mines, divided by the number of inactive surface mines.

Table 3.67 is a summary of inactive mines, obtained from the Department of Energy mine listing. The mines are listed by type, surface and underground. Most of the inactive surface mines are in Colorado, Utah, Arizona and New Mexico. For model derivation purposes, we assumed that there are presently 1250 inactive surface uranium mines.

Table 3.68 lists mine waste and ore production information from 1932 to 1977. Uranium mine waste and ore production statistics, on an annual basis, were available from both surface and underground uranium producers from 1959 to 1976 (DOI59-76). Annual uranium ore production statistics for each uranium mining type (surface and underground) are available for 1948 to 1959 (DOE79) and for combined uranium production from 1932 to 1942 (DOI32-42). In order to estimate waste production for the years prior to 1956, the annual mine type ore production records were multiplied by waste-to-ore ratios. These ratios were estimated from published 1959 to 1976 ore and waste production statistics (DOI59-76). Very little uranium ore was mined from 1942 to 1948, since most of the uranium was obtained by reprocessing vanadium and radium tailings (personal communication with G. Ritter, Bendix Field Engineering Corp., Grand Junction, CO, 1979). The annual waste production for surface mining from 1948 to 1959 was estimated by extrapolating known waste-to-ore ratios (1959 to 1976) through the 1948 to 1959 time period using a "best fit" regression analysis (Fig. 3.23). This method cannot be used to estimate waste-to-ore ratios because the waste production is finite and will always occur, and also surface mining for uranium essentially began in 1950.

Table 3.67 Consolidated list of inactive uranium producers by
State and type of mining

State	Surface	Underground	Percent of Total Surface Mines	Percent of Total Underground Mines
AL	0	1	0.0	< 0.1
AZ	135	189	11	9.3
CA	13	10	1.0	0.49
CO	263	902	21	44
ID	2	4	0.16	0.20
MT	9	9	0.72	0.44
NV	9	12	0.72	0.59
NJ	0	1	0.0	< 0.1
NM	34	142	2.7	7.0
ND	13	0	1.0	0.0
OK	3	0	0.24	0.0
OR	2	1	0.16	< 0.1
SD	111	30	8.9	1.5
TX	38	0	3.0	0.0
UT	378	698	30	34
WA	13	0	1.0	0.0
WY	223	32	18	1.6
Total	1246	2031		

Table 3.68 Uranium mine waste and ore production (MT x 1000)

Year	Surface Mining		Underground Mining		Surface Mining Waste/Ore	Underground Mining Waste/Ore	Total Ore Produced By Surface and/or Underground Mines
	Crude Ore	Waste	Crude Ore	Waste			
1977	5059	237800	4305	3487	47	0.81	9364
1976	4238	190700	3569	2605	45	0.73	7807
1975	3809	139700	2485	2195	37	0.88	6295
1974	3510	129700	2222	1424	37	0.64	5732
1973	3800	182300	1614	934	48	0.58	5414
1972	3447	155100	2439	593	45	0.24	5886
1971	2656	120200	2836	858	45	0.30	5492
1970	2490	76870	3304	962	31	0.29	5794
1969	1653	81000	3171	1184	49	0.37	4824
1968	1989	31360	3382	1163	16	0.34	5371
1967	1393	32510	2897	1024	23	0.35	4290
1966	905	24400	2777	863	27	0.31	1768
1965	1630	17710	3055	809	11	0.26	4685
1964	2344	26680	3227	941	11	0.29	5571
1963	3578	33120	3575	946	9.0	0.26	7153
1962	2895	44640	4892	1087	15	0.22	7787
1961	3051	42500	5017	1117	14	0.22	8068
1960	2691	73570	5104	1868	27	0.37	7795
1959	2494	46790	3796	941	19	0.25	6290
1958	2139	19240	2558	690	9.0 (a)	0.27 (b)	4697
1957	1462	11700	1888	510	8.0	0.27	3350
1956	1131	9048	1595	414	8.0	0.26	2726
1955	339	2650	1043	271	8.0	0.26	1382
1954	241	1930	762	198	8.0	0.26	1003

Table 3.68 (continued)

Year	Surface Mining		Underground Mining		Surface Mining Waste/Ore	Underground Mining Waste/Ore	Total Ore Produced By Surface and/or Underground Mines
	Crude Ore	Waste	Crude Ore	Waste			
1953	162	1300	503	126	8.0	0.25	665
1952	59	472	341	85	8.0	0.25	400
1951	25	203	289	73	8.0	0.25	314
1950	21	167	207	50	8.0	0.24	228
1949			156	37		0.24	156
1948			34	8.3		0.24	34
1947			0			0.24	
1946			0			0.23	
1945			0			0.23	
1944			0			0.23	
1943			0			0.22	
1942			0			0.22	
1941			0.824	0.181		0.22	0.824
1940			0.7221	0.151		0.21	0.722
1939			5.68	1.19		0.21	5.68
1938			3.89	0.817		0.21	3.89
1937			1.55	0.310		0.20	1.55
1936			1.31	0.261		0.20	1.31
1935			1.03	0.207		0.20	1.03
1934			0.230	0.0461		0.20	0.230
1933			0.047	0.00896		0.19	0.047
1932			0.0553	0.0105		0.19	0.0553

(a) Waste to ore ratios from 1950 - 1958 estimated from 1959 - 1972 ratios.

(b) Waste to ore ratios from 1932 - 1958 estimated from 1959 - 1972 ratios.

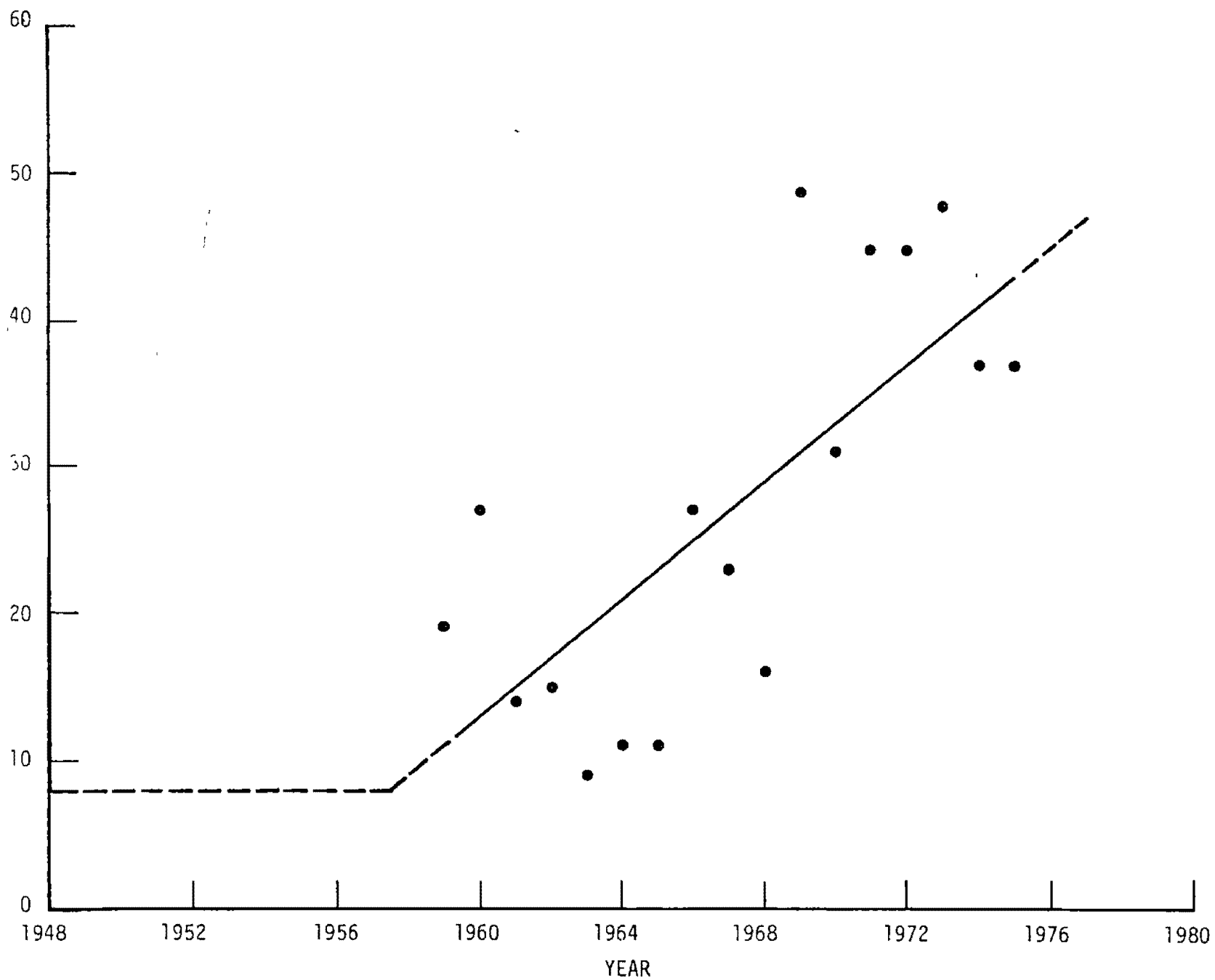


Figure 3.23 Annual waste to ore ratios for surface mining of uranium (1948 to 1979).

Since early surface mines recovered ore bodies very close to the surface, the ore-to-waste ratio would be expected to be relatively small. A range of waste to ore ratios of 8:1 to 35:1 for surface mining has been estimated (C174). The lower ratio was selected to be typical for surface mining from 1948 to 1957 and was used to estimate the waste production for that period. The increase in waste-to-ore ratios from 1959 to 1976 was probably due to several reasons. The gradual depletion of near surface ore deposits required mining deposits at increasing depths, and the development of surface mining equipment now permits economical recovery of ore at greater depths below grade. The waste-to-ore ratios for 1976 to 1977 were projected with the previous regression analysis line fit.

The estimated annual cumulative waste production from uranium surface mining for 1950 to 1978 (Table 3.69) is 1.73×10^9 MT. A crude estimate of the waste production for the model inactive surface mine can be made by dividing the total waste produced to 1978 by the number of inactive mines. But, this overestimates waste production because some of the contemporary wastes are being produced by active mines, and the waste production per mine has increased with increasing contemporary waste-to-ore ratios. To adjust the contemporary waste production for the active mines and the increasing waste-to-ore ratios, we assumed a cutoff date of 1970, based on the description of a contemporary active surface mine (N179). The model mine age is about 1 year as of June 1978, and has an expected life of approximately 17 years. Those mines that were active in 1970 are all assumed to have become inactive between 1970 and 1978. Their percentage of the annual waste of about 12.5 percent was assumed to decrease linearly with time from 1970-1978. For example, all of the wastes produced by surface mines in 1970 (i.e., 7.69×10^7 MT) were produced by surface mines that would be inactive by 1978. The waste production for the following years (1971-1977) was: 1.05×10^8 MT in 1971; 1.16×10^8 MT in 1972; 1.14×10^8 MT in 1973; 6.49×10^7 MT in 1974; 5.24×10^7 MT in 1975; 4.77×10^7 MT in 1976; 2.97×10^7 MT in 1977. The ore production was calculated in the same manner as for the wastes and was 3.27×10^7 MT in 1970. The ore production for the following years was: 2.32×10^6 MT in 1971; 2.58×10^6 MT in 1972; 2.38×10^6 MT in 1973; 1.76×10^6 MT in 1974; 1.43×10^6 MT in 1975; 1.06×10^6 MT in 1976, and 6.32×10^5 MT in 1977. The adjusted cumulative wastes from surface mining from 1950-1978 was 1.11×10^9 MT, and the adjusted cumulative ore production was 4.49×10^7 MT.

Table 3.69 Cumulative uranium mine waste and ore production

Year	Waste (10^3 MT)		Ore (10^3 MT)	
	Surface	Underground	Surface	Underground
1977	1733000	29250	59220	73100
1976	1496000	24950	54160	68840
1975	1305000	21380	49920	65210
1974	1165000	19180	46110	62760
1973	1036000	17760	42600	60500
1972	853200	16820	38800	58960
1971	698100	16240	35350	56510
1970	577800	15370	32700	53600
1969	501000	14410	30200	50330
1968	420000	13220	28550	47160
1967	388600	12060	26560	43810
1966	356200	11040	25170	40910
1965	331800	10180	24260	38090
1964	314000	9369	22640	35000
1963	287300	8425	20290	31750
1962	254200	7479	16720	28210
1961	209600	6391	13810	23310
1960	167100	5273	10770	18320
1959	93510	3406	8075	13150
1958	46720	2466	5580	9433
1957	27470	1776	3442	6839
1956	15770	1266	1979	4943
1955	6720	852	848	3356

Table 3.69 (Continued)

Year	Waste (10^3 MT)		Ore (10^3 MT)	
	Surface	Underground	Surface	Underground
1954	4071	580	509	2313
1953	370	171	46.3	702
1952	842	257	105	1043
1951	370	171	46.3	702
1950	167	98.9	20.9	413
1949		48.6		206
1948		11.4		49.8
1947		3.18		15.3
1946		3.18		15.3
1945		3.18		15.3
1944		3.18		15.3
1943		3.18		15.3
1942		3.18		15.3
1941		3.18		15.3
1940		3.00		14.5
1939		2.85		13.8
1938		1.67		8.12
1937		0.844		4.24
1936		0.533		2.68
1935		0.272		1.37
1934		0.0656		0.333
1933		0.0195		0.102
1932		0.0105		0.0553

Using these adjusted waste and ore values, the model inactive uranium surface mine produced 8.88×10^5 MT of waste and 3.59×10^4 MT of ore.

The volume of the remaining pit of the model surface mine would be equal to the total of the volume of wastes and ore that were removed from the mine. Assuming a density of 2.00 MT/m^3 , the volume of wastes and ore removed from the mine pit would be 4.44×10^5 and $1.80 \times 10^4 \text{ m}^3$, respectively. The pit was assumed to have the shape of an inverted truncated cone with a wall angle of 45° (Fig. 3.24). The ore body was assumed to be a solid right cylinder with a radius of 43.7 m and height of 3.0 m. The pit depth (ground surface to bottom of ore bed) was 36.7 m, and the ground surface area of the pit opening was calculated to be $2.03 \times 10^4 \text{ m}^2$.

3.7.1.1 Waste Rock Piles

Overburden and sub-ore wastes from surface mines have been handled in several ways in the past. In one case, the sub-ore (generally the last material removed from the pit) was piled on top of the overburden. In another case the sub-ore was piled separately and blended with higher grade ore for shipment to the ore buying stations or mills. If the quantity of sub-ore was in excess of that required for blending, it was also dumped on top of the overburden (personal communication with G. Ritter, Bendix Field Engineering Corp., Grand Junction, CO, 1979). The earlier surface mining practices, therefore, generally produced waste piles with their cores containing overburden and their outer surface containing a mixture of overburden and sub-ore.

The actual method of removing and stacking overburden and sub-ore varies from mine to mine. In many cases the wastes were dumped in depressions or washes or stacked in more than one pile. For calculation purposes, we assume that wastes are stacked on a single pile in the shape of a solid truncated cone 10 m high with a 45 degree slope. It is further assumed that the sub-ore removed from the pit is placed evenly on top of the stacked overburden. The area and depth of the sub-ore placed on the waste pile is estimated by determining the areas of the base and top of the pile by iteration, computing the exposed surface area of the pile, computing the volume of the sub-ore, and calculating the depth of the sub-ore.

The areas of the base and top of the waste pile (truncated cone) were determined from the following equation:

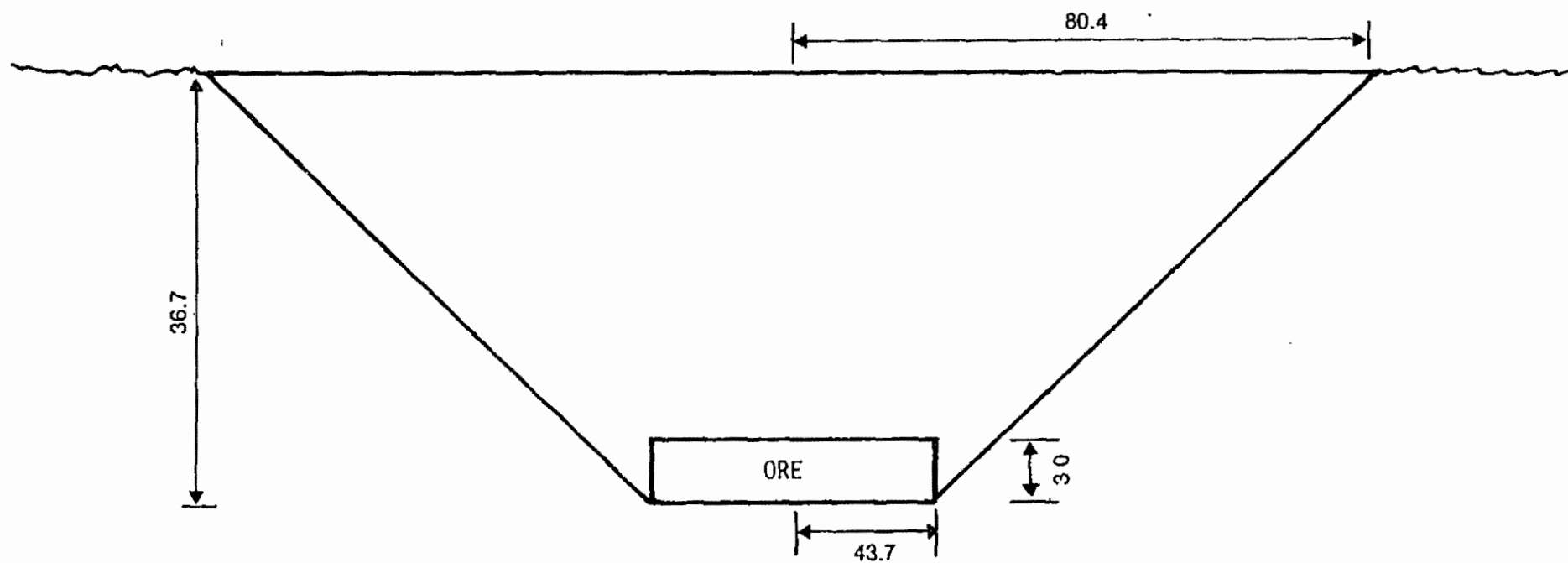


Figure 3.24 Cross section of model inactive surface mine (meters).

$$V = \frac{h}{3} (A_B + A_T + \sqrt{A_B A_T}) \quad \text{where } V = \text{volume of wastes (overburden and sub-ore)} \text{ (m}^3\text{)} \quad (3.25)$$

A_B = area of the base (m²)

A_T = area of the top (m²)

h = perpendicular distance between the base and top (10 m)

Different values of A_B were substituted into the equation until the value of V was equal to the combined volumes of the overburden and sub-ore (i.e; $5.55 \times 10^5 \text{ m}^3$) using a bulking factor of 25%. The area of the cone top was computed (assuming a 45 degree slope) from the diameter of the top (D_T), which is equal to the diameter of the base (D_B), minus 20 meters or $D_T = D_B - 20$. The calculated diameters, D_T and D_B , are 256 m and 276 m, respectively.

The exposed surface area of the waste pile was calculated using the following equation:

$$S = S_L + S_T \quad \text{where } S_L = \text{lateral surface area (m}^2\text{)} \quad (3.26)$$

$$S_L = \frac{L}{2} (C_B + C_T)$$

$$\text{and } S_T = \text{area of the top (m}^2\text{)}$$

$$= \pi r_T^2$$

$$S = \frac{L}{2} (C_B + C_T) + \pi r_T^2 \quad \text{where } C_B = \text{circumference of the base (m)}$$

$$C_T = \text{circumference of the top (m)}$$

L = slant height (m)

r_T = radius of the top (m)

$$S = \frac{14.1}{2} (\pi D_T + \pi D_B) + \pi r_T^2 \quad \text{where } D_T = \text{diameter of the top (m)}$$

$$D_B = \text{diameter of the base (m)}$$

$$S = \frac{14.1}{2} (3.14) (256 + 276) + 3.14 (16384)$$

$$S = 6.33 \times 10^4 \text{ m}^2 \text{ (exposed surface area of waste pile)}$$

The volume of sub-ore removed from the pit is assumed to be equal to the volume of ore removed from the pit. The thickness (T) of the sub-ore plate on the overburden is--

$$T = \frac{V_o}{S} = \frac{2.25 \times 10^4 \text{ m}^3}{6.33 \times 10^4 \text{ m}^2} = 0.36 \text{ m.} \quad (3.27)$$

In summary, the waste pile produced at an inactive uranium surface mine is to be in the shape of a truncated cone having a surface area of $6.33 \times 10^4 \text{ m}^2$. The pile is assumed to have an inner-core of overburden plated with 0.36 m of sub-ore on its exposed surface. In practice, the plate would be a mixture of overburden and sub-ore with the sub-ore concentrations increasing towards the pile surface.

Table 3.70 lists average annual emissions of contaminants due to wind erosion of the overburden pile. To compute these values, an emission factor of 0.850 MT/hectare-yr, computed in Appendix I, was multiplied by the pile surface area, 6.33 hectares, and the stable element concentrations listed in Table 3.19. Uranium and thorium concentrations were assumed to be 110 pCi/g and 2 pCi/g, respectively.

3.7.1.2 Radon-222 from the Mine Area

After the termination of active mining, Rn-222 will continue to exhale from the wall and floor of the pit. Since all of the ore has been removed, the Rn-222 will originate from the overburden and sub-ore surfaces. The surface area of the sub-ore region of the pit is estimated from the volume of ore and sub-ore ($3.6 \times 10^4 \text{ m}^3$) and the shape and size of the pit using the following equations:

$$V = 1/3 h (A_T + A_B + \sqrt{A_T A_B}) \text{ where: } A_T = \pi r_T^2 \quad (3.28)$$

$$S = 1/2 L (C_B + C_b) + A_B \quad A_B = \pi r_B^2 \quad (3.29)$$

The terms in the equations are defined in the previous Section. By substituting the terms $r_B + h$ for r_T in Equation 3.28, h can be solved by iteration.

$$V = 3.6 \times 10^4 \text{ m}^3 = 1/3 h [\pi (r_B + h)^2 + \pi r_B^2 +$$

$$\sqrt{\pi (r_B + h)^2 (\pi r_B^2)}] \text{ when } h = 5.3 \text{ m}$$

The exposed surface-area of the pit that contains the sub-ore is

$$S_s = 1/2 (7.50) \pi (87.4 + 98.0) + 6000 = 8.18 \times 10^3 \text{ m}^2,$$

and the surface area of the overburden section of the pit is

$$S_o = 1/2 (44.4) (\pi) (98.0 + 161.0) = 1.81 \times 10^4 \text{ m}^2.$$

Table 3.71 shows the results of radon flux measurements made at 20 of the tailings piles at inactive uranium mill sites. Also shown is the estimated average Ra-226 content of the tailings and the average Ra-226 content

Table 3.70 Average annual emissions of radionuclides (μCi) and stable elements (kg) in wind suspended dust at the model inactive surface mine

Contaminant	Overburden Pile ^(a)	Contaminant	Overburden Pile ^(a)
Arsenic	0.46	Molybdenum	0.62
Barium	4.9	Nickel	0.11
Cadmium	ND ^(b)	Lead	0.42
Cobalt	0.09	Ruthenium	ND
Copper	0.33	Selenium	0.59
Chromium	0.11	Strontium	0.70
Iron	84	Vanadium	7.6
Mercury	ND	Zinc	0.16
Potassium	135	Uranium-238 and each daughter	1480
Magnesium	19	Thorium-232 and each daughter	11
Manganese	5.2		

(a) Mass Emissions = 5.38×10^6 g/yr.

(b) ND - Not detected.

Table 3.71 Average radon flux of inactive uranium mill tailings piles

Location	Average Radon Flux ^(a) (pCi/m ² -sec)	Estimated Ra-226 ^(b) Tailings Content (pCi/g)	Average Ra-226 Background Soils ^(a) (pCi/g)	Reference ^(a)
<u>ARIZONA</u>				
Monument Valley	20	50	0.95	FBD-GJT-4 (1977)
Tuba City	193	924	0.95	FBD-GJT-5 (1977)
<u>COLORADO</u>				
Durango	197	840	1.48	FBD-GJT-9 (1977)
Grand Junction	359	784	1.52	FBD-GJT-9 (1977)
Gunnison	470	420	1.48	FBD-GJT-12 (1977)
Maybell	86	252	1.52	FBD-GJT-11 (1977)
Naturita	1446	756	1.48	FBD-GJT-8 (1977)
New Rifle	458	504	1.52	FBD-GJT-10 (1977)
Old Rifle	553	980	1.52	FBD-GJT-10 (1977)
Slick Rock	70	171	1.48	FBD-GJT-7 (1977)
<u>IDAHO</u>				
Lowman	125	---	1.12	FBD-GJT-17 (1977)
<u>NEW MEXICO</u>				
Ambrosia Lake	173	760	1.02	FBD-GJT-13 (1977)
Shiprock	340	700	1.7	Bernhardt et al. (1975)
<u>OREGON</u>				
Lakeview	660	420	0.81	FBD-GJT-18 (1977)
<u>SOUTH DAKOTA</u>				
Edgemont	143	---	1.33	FBD-211 (1978)
<u>TEXAS</u>				
Falls City	65	448	0.93	FBD-GJT-16 (1977)
Ray Point	430	518	0.93	FBD-GJT-20 (1977)
<u>UTAH</u>				
Green River	77	140	1.43	FBD-GJT-14 (1977)
Mexican Hat	290	784	0.83	FBD-GJT-3 (1977)
Salt Lake City	1200	896	1.4	Bernhardt et al. (1975)
<u>WYOMING</u>				
Spook Site	1770	356	0.99	FBD-GJT-15 (1977)
Average All Sites	466	563	1.26	

(a) FBD77.
(b) Sw76.

measured in representative background soils for each site. The average radon exhalation rate per average Ra-226 content of tailings material from these data is 0.83 pCi of Rn/m²-sec per pCi of Ra/g.

Data analysis by Schiager (Sc74) indicates a radon exhalation rate of 1.6 pCi of Rn/m²-sec per pCi of Ra/g. This value has often been used in the environmental impact statements to assess the radon flux from tailings materials.

Table 3.72 summarizes data obtained during radiological surveys of inactive uranium mine sites in New Mexico and Wyoming during the spring of 1979 (Wo79). Radon exhalation rates were measured with charcoal cannisters and the radium-226 concentrations were determined for composite surface samples taken from overburden, sub-ore, and waste rock piles. The average radon-222 exhalation rate per average radium-226 content of the overburden, sub-ore, and waste rock piles was 0.27, 0.11, and 0.12 pCi of Rn/m²-sec per pCi of Ra-226/g, respectively.

Measurements of the background flux and Ra-226 content of typical background soils were reported for the Edgemont, South Dakota site (FBD78). These data indicate a value of 1.05 pCi of Rn/m²-sec per pCi of Ra/g. Table 3.73 summarizes background radon flux estimates for several regions of the United States. Considering the average U.S. background flux to be 0.82 pCi of Rn/m²-sec (Tr79) and the average U.S. background soil Ra-226 content to be 1.26 pCi of Ra/g (Oa72), the average U.S. background radon exhalation rate is estimated to be 0.65 pCi of Rn/m²-sec per pCi of Ra/g. The average background radon exhalation rate for New Mexico and Wyoming (Table 3.72) was 0.33 pCi of Rn/m²-sec per pCi of Ra/g. Therefore, the grand average U.S. background radon exhalation rate has been estimated to be 0.68 pCi of Rn/m²-sec per pCi of Ra/g, and the grand average U.S. background soil Ra-226 content has been estimated to be 1.6 pCi/g.

We estimated the total radon released from the model abandoned surface mine area from the following parameters:

1. Radon exhalation from the sub-ore surface area of the pit--
 - the exposed sub-ore surface area (S_s) = 8.18×10^3 m²;
 - the average radium-226 content of the sub-ore = 110 pCi/g; and
 - the radon flux rate for sub-ore = 12 pCi of Rn/m²-sec.

Table 3.72 Average radon flux measured at inactive uranium mine sites

Location	Area	Average Radon Flux (pCi/m ² -sec)	Number of Flux Measurements	Average Radium-226 Content of Surface Sample (pCi/g)
<u>Underground Mines</u>				
San Mateo Mine, New Mexico	Waste pile	18	11	117
	Heap leach pond	38	3	81
	Background	0.29	1	0.77
Barbara J # 1 Mine, New Mexico	Waste pile	7.9	6	110
	Background	0.41	1	3
<u>Surface Mines</u>				
Poison Canyon 1, New Mexico	Sub-ore	7.0	1	43
	Overburden piles	6.7	5	62
	Background	0.33	1	2.1
Poison Canyon 2, New Mexico	Sub-ore	5.3	3	---
	Overburden pile	9.8	6	---
Poison Canyon 3, New Mexico	Sub-ore	11	2	---
Morton Ranch (Pit 1601), Wyoming	Sub-ore	24	12	170
	Overburden	9.7	4	23
	Background	2.3	2	3
Grand Averages	Sub-ore	12		110
	Overburden	8.7		32
	Waste Rock	13		110
	Background	0.83		2.2

Source: Wo79.

Table 3.73 Background radon flux estimates

Location	Radon Flux pCi/m ² -sec
<u>Background Soils of the U.S.</u>	
Champaign County, Illinois	1.4
Argonne, Illinois	0.56
Lincoln, Massachusetts	1.3
Socorro, New Mexico	0.90
Socorro, New Mexico	1.0
Socorro, New Mexico	0.64
Yucca Flat, Nevada	0.47
Texas	0.27

Average U.S. Background Radon Flux = 0.82 pCi/m²-sec.

Source: Tr79.

Therefore, the radon released from the sub-ore surface area of the pit is $8.18 \times 10^3 \text{ m}^2 \times 12 \text{ pCi of Rn/m}^2\text{-sec} \times 86400 \text{ sec/day} = 8.48 \text{ mCi of Rn/day}$.

2. Radon exhalation from the overburden surface area of the pit--
 - . the exposed overburden surface area (S_o) = $1.81 \times 10^4 \text{ m}^2$;
 - . the average radium-226 content of the overburden = 32 pCi/g; and
 - . the radon flux rate for overburden is 8.7 pCi of Rn/m²-sec.

Therefore, the radon released from the overburden surface area of the pit is $1.81 \times 10^4 \text{ m}^2 \times 8.7 \text{ pCi of Rn/m}^2\text{-sec} \times 86400 \text{ sec/day} = 13.6 \text{ mCi of Rn/day}$.

3. Radon exhalation from the overburden pile remaining at the pit--
 - . the exposed surface area of the waste pile (S_w) = $6.33 \times 10^4 \text{ m}^2$;
 - . the Ra-226 content of the surface of the overburden pile is the same as the sub-ore content = 32 pCi/g; and
 - . the radon flux rate for the overburden pile is 8.7 pCi of Rn/ m^2 -sec.

Therefore, the radon exhalation rate from the overburden pile is $6.33 \times 10^4 \text{ m}^2 \times 8.7 \text{ pCi of Rn/m}^2\text{-sec} \times 86400 \text{ sec/day} = 47.6 \text{ mCi of Rn/day}$.

The total radon release rate at the abandoned surface mine site is the sum of the above three source terms, 69.7 mCi/day. The estimated radon release rate for background soils for an undisturbed area equivalent to the surface mine area uses the following parameters:

- . the ground surface area equivalent to the area of the pit opening ($2.03 \times 10^4 \text{ m}^2$) and the overburden pad area ($5.98 \times 10^4 \text{ m}^2$) = $8.01 \times 10^4 \text{ m}^2$, and
- . the radon flux rate for background soils in uranium mining areas = 0.83 pCi of Rn/ m^2 -sec (Table 3.72).

Therefore, the radon exhalation rate from an undisturbed area equivalent to the model surface mine is

$$8.01 \times 10^4 \text{ m}^2 \times 0.83 \text{ pCi of Rn/m}^2\text{-sec} \times 86400 \text{ sec/day} = 5.7 \text{ mCi of Rn/day}.$$

Table 3.74 summarizes the annual radon-222 release from the model inactive uranium surface mine and all inactive uranium surface mines.

3.7.1.3 Land Surface Gamma Radiation

The surface mine uranium overlying strata must be removed in order to gain access to the uranium-bearing host materials and the ore body. The ore body consists of ore and sub-ore, and the sub-ore is simply that fraction of the ore body that contains ore uneconomical to recover. The end result of the mining is that the residues (sub-ore) enhance natural radioactive materials. That is, they are exposed or brought to the earth's surface. The enhancement will cause, in most cases, increased aboveground radiation

Table 3.74 Summary of estimated radon-222 releases from
inactive surface mines

Source	Estimation Method	Annual Release, Ci
Mine Pit		
Sub-ore area	Model mine and limited field measurements	3.1
Overburden area	Model mine and limited field measurements	<u>5.0</u>
Total		8.1
Overburden Pile	Model mine and limited field measurements	17.4
Background	Rn-222 flux measurements and projected surface areas of model mine pit and overburden pile	2.1
Model Mine	Net Rn-222 release	23.4
All Inactive Mines	Annual net Rn-222 release from model times 1250 mines	29,000

exposure rates around the mining area. Ore and sub-ore lost through handling are subject to wind and water erosion. This effectively increases the mine site area in a radiological sense. The gamma radiation exposure levels on and around a mine site can be high enough to restrict use of the area after mining.

Gamma radiation surveys were conducted at some inactive uranium surface mining areas. Table 3.75 lists the ranges of exposure rates found. Appendix G contains more specific information concerning the surveys. The residual exposure rate levels would probably preclude unrestricted use of the pits, waste piles, and overburden.

Figure 3.25 depicts gamma radiation measurements made on radials extending outward from an inactive surface mine pit. The measurements were made with a pressurized ion chamber (PIC) at approximately 61 m intervals on each radial. As expected, the exposure rate decreases with distance away from the pit, indicating surface contamination from wind and water erosion of the spoils and ore piles. Some of the contamination may also have originated from ore and sub-ore dust losses during mining.

Since the pit resides over a former ore body and connecting or adjacent ore bodies may be located near the mine, some caution is necessary when interpreting the gamma exposure rates as indicative of surface contamination. Development drilling, indicating the presence of ore bodies, is prevalent throughout the north, west, and south areas around the pit. The northeast, east, and southeast areas around the pit have exploratory drill holes only. They indicate the probable absence of ore bodies. Although the north, northwest, west, and southwest radials cross below grade ore bodies, it is not reflected by the gamma measurements. Unless the ore body is very close to the surface, its gamma radiation will not be measured (i.e., the 1/10 value layer for earth shielding is about 0.3 m). The south radial, however, did cross an ore outcropping.

If the exposure rate measurements made at the end points of the radials (south radial excepted) are assumed to be near background, their mean value is $14.4 \mu\text{R/hr}$ with a 2 sigma error of $1.6 \mu\text{R/hr}$.

Assuming all measurements in excess of $14.4 + 1.6 \mu\text{R/hr}$ or $16.0 \mu\text{R/hr}$ are a result of eroded ore and sub-ore from the mining activities, an iso-exposure rate line enclosing the eroded materials can be constructed around the mine site. The line is constructed on Fig. 3.25 and is qualitatively

Table 3.75 Summary of land surface gamma radiation surveys
in New Mexico, Texas and Wyoming

Location	Area	Gamma Radiation Exposure Rate (μ R/hr)
Poison Canyon, New Mexico	Pits	40 to 190
	Waste piles	65 to 250
	Overburden	25 to 65
Texas	Pits	5 to 400
Morton Ranch, Wyoming (1601 Pit)	Pit	16 to 63
	Ore piles	200
	Overburden	59 to 138

Source: Wo79 for New Mexico and Wyoming and Co77 for Texas.

adjusted on the south radial to compensate for the ore outcropping. The line bulges into the southeast quadrant indicating erosion by the predominant northwest winds and contamination of about 0.3 km^2 .

In summary, it appears that the residual gamma radiation exposure levels at surface mining pits and overburden piles would preclude these areas from unrestricted use. It also appears that wind and water erosions of the spoils, ore, and sub-ore are occurring and causing land contamination far removed from the mining area. Several surface mines were gamma surveyed in New Mexico. The mines could not be individually gamma radiation surveyed because of their close proximity, cross contamination from eroded ore and sub-ore, and possible ore outcrops.

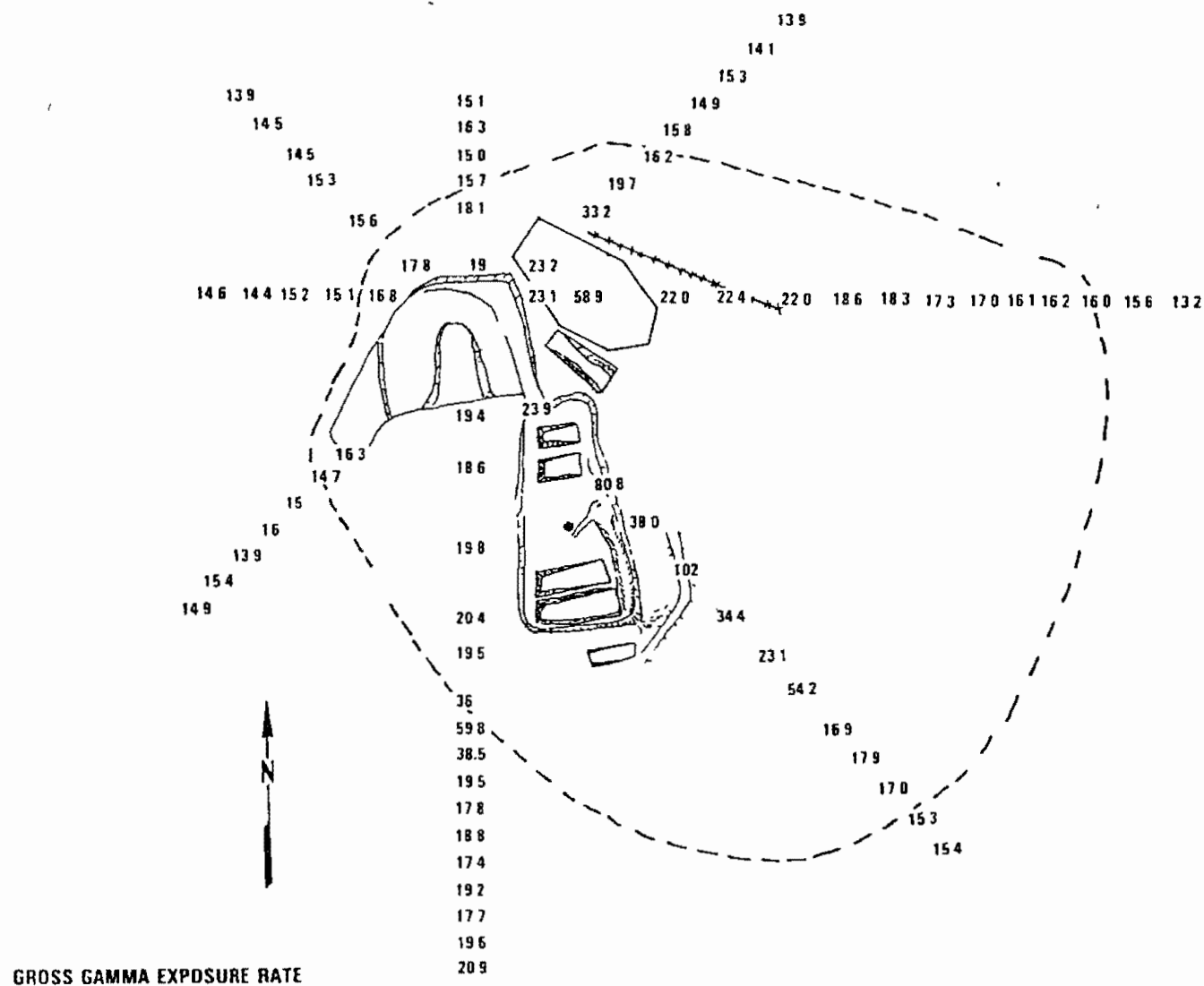


Figure 3.25 Results of gamma exposure rate survey at the 1601 pit and environs, Morton Ranch uranium mine, Converse County, Wyoming ($\mu\text{R/hr}$)

3.7.2 Inactive Underground Mines

The model inactive underground mine is basically defined by dividing the total reported volumes of ore and waste removed by inactive underground mining by the number of inactive underground mines. The number of inactive underground mines has been obtained from the U.S. Department of Energy mine listing in Table 3.67. Table 3.67 lists the mines by state and type of mine. Forty-four percent of the inactive underground mines are located in Colorado, 34 percent in Utah, 9.3 percent in Arizona, and 7.0 percent in New Mexico.

For modeling purposes, we assume that there are presently 2030 inactive underground uranium mines. Table 3.69 lists the estimated underground mine waste and ore production for 1932 to 1977. Uranium mine waste and ore production statistics, on an annual basis, were available for underground producers from 1959 to 1977 (DOI59-76). Annual uranium ore production statistics for underground mining are available from 1948 to 1959 (DOE79) and from 1932 to 1942 (DOI32-42). We estimated the mine waste production for the period of 1932 to 1960 from underground mining waste-to-ore ratios and established waste-to-ore ratios using the published ore and wastes production statistics from 1959 to 1976 (DOI59-76). These ratios were fitted with a line by regression analysis in order to estimate the waste-to-ore ratios from 1932 to 1959 (Fig. 3.26). Two lines were fitted to the known waste-to-ore ratios because of the abrupt change in the ratios from 1972 to 1976. We assumed that the steeper slope was caused by increased waste production from the larger and deeper underground mines operated during this time. The estimated annual waste-to-ore ratios were multiplied by the published annual ore production values to estimate the annual waste production from 1932 to 1959. We assumed that no ore was produced from 1942 to 1948 because most of the uranium was obtained by reprocessing vanadium and radium tailings during that period (Private communication with G. C. Ritter, 1979, Bendix Field Engineering Corporation, Grand Junction, Colorado). Table 3.69 lists the cumulative annual waste production from underground mining from 1932 through 1977. The total waste produced for this period was 2.92×10^7 MT, and the total ore produced was 7.31×10^7 MT.

A simplistic way to identify a model inactive underground mine would be to divide the cumulative tonnage of ore and wastes by the number of inactive mines. We estimated the number of inactive mines from the U.S. Department of Energy mine listing (Section 2.0 and Table 3.67). The model inactive under-

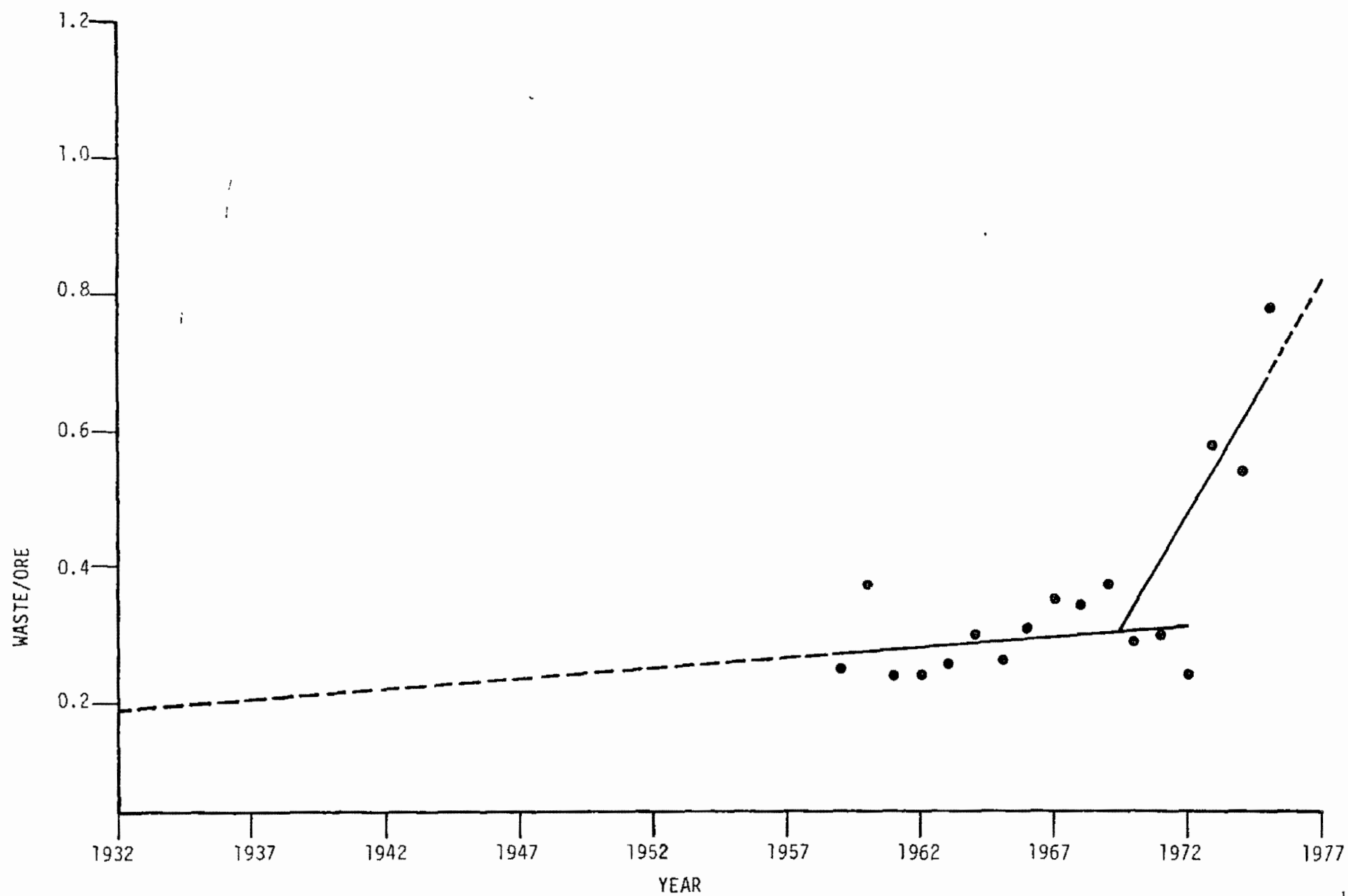


Figure 3 26 Waste to ore ratios for inactive underground uranium mines from 1932 to 1977.

ground mine produced 3.60×10^4 MT of ore and 1.44×10^4 MT of waste. Unfortunately, some of the contemporary waste and ore production has been produced by both active and inactive mines. In order to adjust the contemporary ore and waste production for that portion of the ore and wastes generated by active mining, we assumed a model active mine having a mining life of 15 years (St79). The mid-life of the mine was assumed to have occurred in 1978, with production beginning in 1971.

We also assumed that some of the mines became inactive during the 1971-1978 period and that their numbers decreased linearly. For example, 2.44×10^6 MT of ore was produced in 1972 and 85.7 percent of that ore produced was from mines that were inactive by 1978. Therefore, adjusted ore production was 2.09×10^6 MT for 1971. The ore production for 1973 was 1.15×10^6 MT and 1.27×10^6 MT in 1974; 1.06×10^6 MT in 1975; 1.02×10^6 MT in 1976; and 6.16×10^5 MT in 1977. The adjusted waste production was: 5.08×10^5 MT in 1972; 6.67×10^5 MT in 1973; 8.13×10^5 MT in 1974; 9.43×10^5 MT in 1975; 7.43×10^5 MT in 1976; and 4.99×10^5 MT in 1977.

Through 1978, the cumulative adjusted ore production from inactive underground mines was 6.37×10^7 MT, and the cumulative adjusted waste production was 2.04×10^7 MT. The model inactive underground mine was assumed to have produced 3.14×10^4 MT of ore and 1.00×10^4 MT of waste. Assuming a density of 2.0 MT per m^3 , the volume of ore and waste removed were 1.6×10^4 and $5.0 \times 10^3 \text{ m}^3$, respectively.

Fifty percent of the waste volume mined we assumed to be sub-ore. The volume of waste rock (i.e., containing no sub-ore) removed during the mining is $2.5 \times 10^3 \text{ m}^3$. Assuming an entry dimension of 1.83 m x 2.13 m, about 615 m of shafts and haulways are in the model mine. The ore body we assumed to have an average thickness of 1.8 m with a length and width of 91.2 m each. The surface area of the passages would be $4.83 \times 10^3 \text{ m}^2$. The surface area of the mined-out ore body would be $1.71 \times 10^4 \text{ m}^2$.

3.7.2.1 Waste Rock Piles

Wastes produced from underground uranium mining were generally cast or dumped near the mine entries. Those wastes that were dumped on relatively flat terrain formed dome-shaped piles. Wastes cast from rim mines generally formed long, thin sheets down the canyon slopes. Since most of the inactive underground mines are in the Uravan Mineral Belt, the waste pile shape (dome)

is assumed to be predominant (see Appendix G.1.2) and is used for the calculations of the waste pile dimensions.

The waste produced at a typical underground mine consists of waste rock and sub-ore. The waste rock is assumed to be on the bottom of the waste pile since it was generally removed first. Sub-ore, which was removed later, is assumed to cover or plate the waste pile. The waste piles are assumed to be dome shaped, covering a circular area of 0.40 hectares. The dome is assumed to be a spherical segment with a height (b) and base (c) of 71.8 m. The volume (V) of the spherical segment, $6.3 \times 10^3 \text{ m}^3$ when corrected for bulking, is equal to the volume of wastes and is expressed as

$$V = \frac{1}{24} \pi b (3c^2 + 4b^2). \quad (3.30)$$

The surface area of the spherical segment is given by the expression

$$S = \frac{1}{4} \pi (4b^2 + c^2) \text{ where } S = \text{Surface area (m}^2\text{)}. \quad (3.31)$$

The term b is solved by substitution and iteration in the former equation and is substituted in the latter equation to determine the surface area of the wastes:

$$V = 6.3 \times 10^3 \text{ m}^3 = \frac{1}{24} \pi b (15465 + 4b^2) \text{ where: } b = 3.1 \text{ m}. \quad (3.32)$$

The surface area of the waste pile is

$$\begin{aligned} S &= \frac{1}{4} \pi (4b^2 + c^2) \\ &= \frac{1}{4} (3.14) (38 + 5155) \\ &= 4.08 \times 10^3 \text{ m}^2. \end{aligned} \quad (3.33)$$

The thickness (T) of sub-ore on the surface of the waste pile is

$$\frac{\text{volume of sub-ore}}{\text{area of waste pile}} = \frac{3.2 \times 10^3 \text{ m}^3}{4.08 \times 10^3 \text{ m}^2} = 0.78 \text{ m}. \quad (3.34)$$

In summary, the waste pile at an inactive underground uranium mine is assumed to have the shape of a spherical segment with a surface area $4.08 \times 10^3 \text{ m}^2$. The pile is assumed to have an inner core of waste rock covered or plated with 0.78 m of sub-ore on its exposed surface. It is expected that the plate of sub-ore on the waste pile would be more pronounced than the sub-ore plates on overburden piles at surface mines because of diminished

blending, mining practices, and the lower waste-to-ore ratio. The grand average of the radium-226 concentrations in the waste rock and overburden piles (Table 3.72) appear to confirm this expectation.

Table 3.76 lists average annual emissions of contaminants due to wind erosion of the waste rock pile. These values were estimated by multiplying an emission factor of 2.12 MT/hectare-yr, derived in Appendix I, by the waste pile surface area, 0.408 hectares, and the stable element concentrations given in Table 3.19. We assumed uranium and thorium concentrations to be 110 pCi/g and 2 pCi/g, respectively.

Table 3.76 Average annual emissions of radionuclides (μCi) and stable elements (kg) in wind suspended dust at the model inactive underground mine

Contaminant	Waste Rock Pile ^(a)	Contaminant	Waste Rock Pile ^(a)
Arsenic	0.07	Molybdenum	0.10
Barium	0.80	Nickel	0.02
Cadmium	ND ^(b)	Lead	0.07
Cobalt	0.01	Ruthenium	ND
Copper	0.05	Selenium	0.10
Chromium	0.02	Strontium	0.11
Iron	14	Vanadium	1.2
Mercury	ND	Zinc	0.03
Potassium	22	Uranium-238 and each	
Magnesium	3.0	daughter	238
Manganese	0.83	Thorium-232 and each	
		daughter	1.7

(a) Mass emissions = 8.65×10^5 g/yr.

(b) ND - Not detected.

3.7.2.2 Radon-222 from the Mine Area

We estimated the total radon released from the model inactive underground mine from the following parameters:

1. Radon exhalation from the waste rock pile--
 - . the exposed surface area of the waste pile = $4.1 \times 10^3 \text{ m}^2$;
 - . the average Ra-226 content of the waste pile is 110 pCi/g; and
 - . the radon flux rate for the waste pile is 13 pCi of Rn/ m^2 -sec.

Therefore, the radon released from the waste pile is

$$4.1 \times 10^3 \text{ m}^2 \times 13 \text{ pCi of Rn/m}^2\text{-sec} \times 86400 \text{ sec/day} = 4.6 \text{ mCi of Rn/day.}$$

2. Typical background release rate--
 - . the ground surface area equivalent to the area covered by the waste pile = $4.1 \times 10^3 \text{ m}^2$, and
 - . the radon flux rate for background soils in uranium mining areas = 0.83 pCi of Rn/ m^2 -sec (Table 3.72).

Therefore, the radon exhalation rate from an undisturbed area equivalent to the waste pile of a model underground mine is

$$4.1 \times 10^3 \text{ m}^2 \times 0.83 \text{ pCi of Rn/m}^2\text{-sec} \times 86400 \text{ sec/day} = 0.29 \text{ mCi of Rn/day.}$$

The net radon release rate due to the waste pile at the inactive underground mine is 4.6 minus 0.29 or about 4.3 mCi of Rn/day above normal background.

Natural ventilation will occur in most mines and usually is considered by mine ventilation engineers when planning the forced ventilation systems. The natural force that can maintain a natural air flow due to temperature differences is thermal energy. The thermal energy added to a system is converted into a pressure difference. If the pressure difference is sufficient to overcome head losses, a flow of air will occur.

Natural ventilation depends upon the difference between the temperature inside and outside of a mine and the difference between the elevation of the mine workings and the surface. Air flow by natural ventilation is generally small ($140 - 566 \text{ m}^3/\text{min}$) in shallow mines (Pe52). In deep mines, natural ventilation flows may range from 1,420 to $4,250 \text{ m}^3/\text{min}$ (Pe52). The flow in either the shallow or deep mines depends upon the depth, size, and number of

openings. The intensity of thermal energy-induced natural pressure usually ranges from a few hundredths to a few tenths cm of water in shallow (less than 460 m deep) mines (Pe52). The maximum pressure drop per 305 m of depth in deep mines is about 2.54 cm of water in winter and about 0.84 cm during the summer (Pe52).

In general, natural ventilation is subject to considerable fluctuation. It usually increases to a maximum in winter and a minimum in summer for deep mines. The typical inactive underground uranium mine would be shallow; therefore, the natural ventilation would be expected to reach its maximum in the winter and summer and its minimum in the spring and fall (air temperature in the mine closely approaches the outside temperature during the spring and fall).

A first approximation of the annual release of Rn-222 from an inactive underground mine simply would be that all Rn-222 released into the mine air will be exhausted by natural ventilation before a significant radioactive decay occurs. That is, the quantity of radon released into the mine is equal to the quantity of radon released from the mine. The quantity of Rn-222 released from the sub-ore surfaces remaining in the mined-out ore body is

$$\frac{Q \text{ (pCi Rn-222)}}{\text{sec}} = A \times \phi_{so}, \quad (3.35)$$

where A is the surface area of the mined out ore body (m^2)

ϕ_{so} = exhalation rate of the Rn-222 from sub-ore per unit area per unit time, 12 $\frac{\text{pCi}}{\text{m}^2\text{-sec}}$ (Section 3.7.1.2)

$$Q = (1.71 \times 10^4 \text{ m}^2) \frac{(12 \text{ pCi})}{\text{m}^2\text{-sec}} = 2.1 \times 10^5 \text{ pCi/sec.}$$

It should be noted that ϕ_{so} is the average radon flux physically measured from sub-ore bodies in inactive surface mines (Section 3.7.1.2). Because of safety considerations, no measurements were made from sub-ore bodies in inactive underground mines during the April 1979 field surveys. The annual Rn-222 source term from the mined-out ore body in an inactive underground uranium mine, using the preceding assumptions, is

$$Q \left(\frac{\text{Ci}}{\text{yr}} \right) = 2.1 \times 10^5 \frac{\text{pCi}}{\text{sec}} \times 3.6 \times 10^3 \frac{\text{sec}}{\text{hr}} \times 24 \frac{\text{hr}}{\text{d}} \times 365 \frac{\text{d}}{\text{yr}} \times \frac{1}{10^{12} \frac{\text{pCi}}{\text{Ci}}} = 6.6 \frac{\text{Ci}}{\text{yr}}.$$
(3.36)

The annual Rn-222 source term (Q) from the passageways, assuming an exhalation rate of $8.7 \frac{\text{pCi}}{\text{m}^2\text{-sec}}$ for overburden, is (Section 3.7.1.2)

$$(4.8 \times 10^3 \text{ m}^2) (2.7 \times 10^{-4} \frac{\text{Ci}}{\text{yr-m}^2}) = 1.3 \frac{\text{Ci}}{\text{yr}}.$$
(3.37)

The air flow rate from the mine, assuming $140 \text{ m}^3/\text{min}$ for an average shallow mine, will exchange the mine air every three hours. The average annual radon-222 concentration will be

$$7.9 \text{ Ci/yr} \times \frac{1}{7.4 \times 10^7 \text{ m}^3/\text{yr}} \times \frac{1}{1000 \text{ l/m}^3} = 107 \text{ pCi/l}.$$

The radon daughter concentration will be about 87 percent of equilibrium with the radon, assuming a mean residence time of the radon in the mine to be 1.5 hours.

Several inactive mines in the Grants, New Mexico area were monitored for radon discharges by natural ventilation. One of the mines monitored was relatively small and had a vertical shaft access. Five cased 30 cm diameter vents were found and were assumed to be connected with the mine. The shaft was covered with steel plate, but access holes were cut in the plate and one corner had been pried up. Four vents were capped with buckets. Just one cover was gas tight. One vent was partially covered with a piece of wood. Only very small flow rates due to natural ventilation were measured at the shaft and vents. The maximum radon emission from the mine per day was estimated to be $2.8 \times 10^3 \text{ } \mu\text{Ci}$. This low radon discharge rate is probably due to partial blockage of the vents and water in the mine. The mine was partially flooded, and flowing water could be seen at the bottom of the shaft. The

effect of the water would be to partially or completely close off the mine workings and substantially reduce natural ventilation. The water would also dissolve and substantially suppress the radon exhaling from the surface areas of the mine. Thus, we believe that the radon discharges from wet inactive mines via natural ventilation will be minimal.

Investigation at another inactive mine revealed that it was connected to three other inactive mines that were subsequently connected to two active mines. Ventilation fans at the connecting active mines were usually shut down after the end of the day shift and on weekends. Mine air was exhausted by natural ventilation through the shaft (highest opening) and vents of the mine investigated. A flow rate up to $88 \text{ m}^3/\text{min}$ was observed coming from the shaft, and radon-222 concentrations reached $11,000 \text{ pCi}/\ell$. The average flow rate observed over a weekend was $75 \text{ m}^3/\text{min}$, with an average radon-222 concentration of $9,800 \text{ pCi}/\ell$. The average radon emission was 1.1 Ci/day .

Figure 3.27 is a plot of the changes in the Rn-222 concentration in the air from the shaft of the inactive mine investigated. The average of the measurements of the air flow rate from the shaft was about $76 \text{ m}^3/\text{min}$. The Rn-222 concentration increased almost linearly with time for about 20 hours after the fans were shut down at the end of the day shift on April 27, 1979. The Rn-222 concentrations also leveled off at about $10,000 \text{ pCi}/\ell$. A dip, presumed to have been caused by high winds, occurred in the Rn-222 concentration curve from about 1000 to 1600 hours on April 28, 1979.

Since the curve is relatively flat at $10,000 \text{ pCi}/\ell$, it is assumed that the rate of production of the Rn-222 is equal to the rate of removal of the Rn-222 from the six mines. The average residence time of the radon in the mine air is assumed to be approximately 10 hours, and the radon daughters would be in near-equilibrium (assumed to be ≈ 90 percent). Assuming that all six interconnecting mines contributed equally to the source term measured, the release rate of Rn-222 for a single mine will be

$$10,000 \text{ pCi}/\ell \times 76,000 \ell/\text{min} \times 1440 \text{ min/day} \times 10^{-12} \text{ Ci/pCi} \div 6 \\ = 0.18 \text{ Ci/day}.$$

Based on the preceding estimation of Rn-222 and progeny released from a typical mine on the Colorado plateau and physical measurements at six connected mines, the annual radon release rate may range from 7.9 to 66 Ci/yr. These source term estimates, of course, are based on a single mine. Many mine workings are, in fact, interconnected. If these interconnected workings are assumed to constitute a single mine, then the upper limit of Rn-222 and

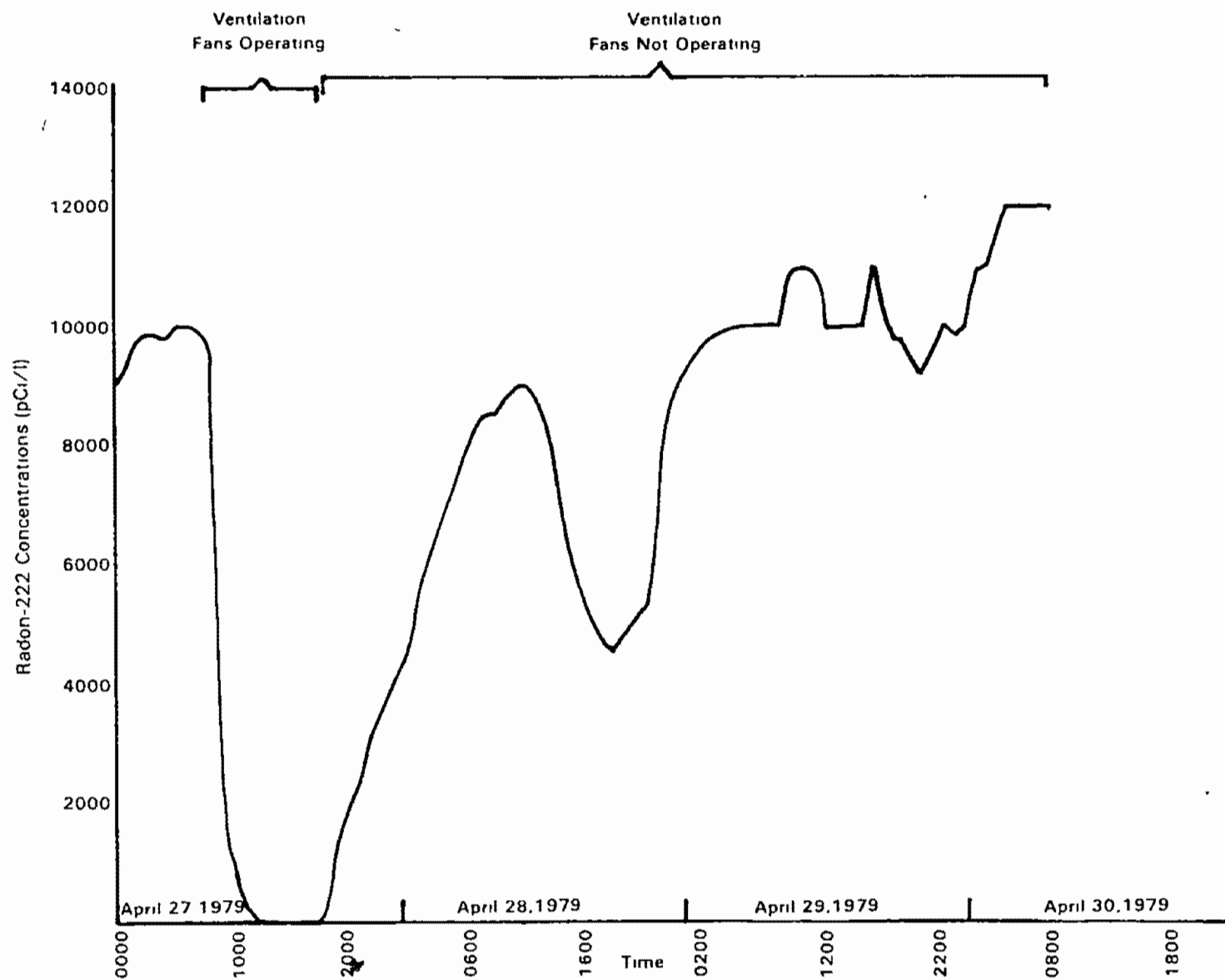


Figure 3 27 Radon-222 concentrations in mine air discharged by natural ventilation.

progeny discharge known at this time will be about 10,000 pCi/l with an annual Rn-222 source term of about 400 Ci/yr. For example, 67 percent of all inactive underground uranium mines are in or near the Uravan mineral belt and are probably dry. Their aggregate Rn-222 discharge by natural ventilation is estimated to be

$$1360 \text{ mines} \times 66 \frac{\text{Ci Rn-222}}{\text{yr-mine}} = 9.0 \times 10^4 \text{ Ci/yr.}$$

In summary, there is little information available on the discharge of Rn-222 and its progeny from the vents and entries of inactive uranium mines by natural ventilation. Some physical measurements indicate that the discharges may be substantial. It is known, through surveys conducted to support this study, that a large majority of the inactive uranium mines are not isolated from the atmosphere and are capable of discharging their Rn-222 and progeny into the local environment. It is also known that some self-sealing will probably occur at some of the mines, due to flooding, cave-ins, and subsidence. Table 3.77 summarizes estimates of the annual radon-222 releases from inactive underground uranium mines. This potential source of exposure could be practically eliminated by proper sealing of the inactive mines.

3.7.2.3 Land Surface Gamma Radiation

Gamma radiation surveys were conducted around underground mining areas in Colorado and New Mexico. Table 3.78 lists the ranges of gamma radiation exposure rates measured at some of the mines. The elevated gamma ray exposure rates on the waste piles are due primarily to plating those piles with sub-ore removed during the mining process.

Some radioactive materials originating from ore and sub-ore handling can be lost into the local environment around a mine site. Erosion of the mine wastes can also disperse contaminants into the local environment. Figure 3.28 illustrates gross gamma radiation exposure rate measurements around an inactive underground uranium mine in New Mexico. Background gamma-ray exposure rate measurements made around the mine area ranged from 12 to 15 $\mu\text{R/hr}$. According to the measurements made, exposure rate levels exceeded background from 50 to more than 100 meters from the waste piles. The area that has been contaminated far exceeds the area physically disturbed at the mine site. Gross gamma exposure rates measured on the waste piles averaged about 95

Table 3.77 Summary of radon-222 releases from inactive underground mines

Source	Estimation Methods	Annual Release, Ci
Model Mine		
Waste Rock Piles	Calculated volume & surface area; limited field measurements of radon flux	1.7
Underground workings	Radon release based on natural ventilation rate for shallow mines	
Sub-ore Surfaces	Calculated surface area; limited radon flux measurements of sub-ore	6.6
Passageways	Calculated passageway surface area; limited measurements of radon flux from overburden	1.3
Background	Field measurements of radon flux; and projected area of waste rock pile	0.11
Model Mine	Total radon source minus background	9.5
Actual Mine		
Underground workings (dry)	Field measurements	66
Underground workings (wet)	Field measurements	1.1
Waste rock piles	Calculated volume & surface area; Limited field measurements of radon flux	1.7

Table 3.78 Summary of land surface gamma radiation surveys in
Colorado and New Mexico

Location	Area	Gamma Radiation Exposure Rate (μ R/hr)
Boulder, Colorado	Waste piles	40 to 100
Uravan, Colorado	Waste piles	50 to 220
San Mateo, New Mexico	Waste pile	35 to 275
	Ore	100 to 350
	Overburden	20 to 120
	Background	10 to 13
Mesa Top Mines, New Mexico	Waste piles	25 to 290
Barbara J #1 Mine, New Mexico	Waste piles	21 to 170
	Background	12 to 15

Source: Wo79.

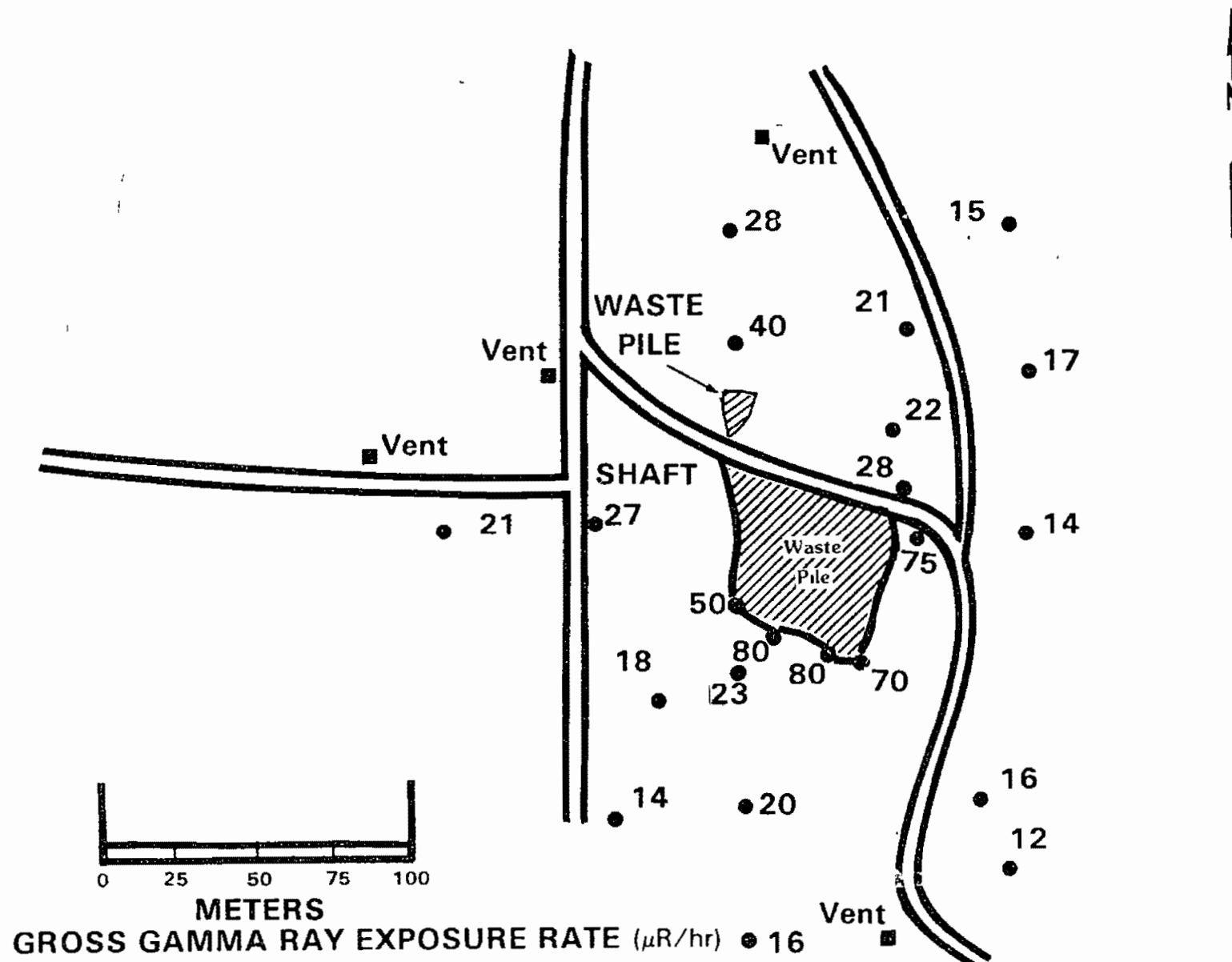


Figure 3.28 Gamma radiation survey around an inactive underground uranium mine in New Mexico.

$\mu\text{R/hr}$, which would make them unsuitable for unrestricted use.

In summary, wastes from underground uranium mining technologically enhance natural radioactivity and may be considered low-level radioactive wastes. Improperly controlled wastes will be dispersed into the surrounding environment by the mining activities and erosion.

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SECTION 4

DESCRIPTION OF MODEL MINES

4.0 Description of Model Mines

Section 1.3 describes uranium mines and their operations, and Section 3 describes the potential sources of contamination at the principal types of active and inactive mines. These discussions include an analysis of the potential sources of contamination, quantities of contaminants associated with the different sources, variations in the sources, and estimates of the values needed to define the impact that these sources may impose upon the environment and nearby populations. We attempted to define these terms and mining parameters in a way that would reflect a general view of the uranium mining industry and permit a generic assessment. The parametric values that we have chosen for this assessment are listed below. The sections of this report from which they were derived are given in parentheses.

4.1 Surface Mine

The model open pit (surface) mine will be located in Wyoming. It is the mine defined in Section 3.3 as the "average large mine." However, to define the total impact of all 63 open pit mines operating in the United States in 1978 we used the parameters developed in Section 3.3 for the "average mine."

Production Parameters (1.3.1, 3.3.1)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Ore, MT/yr	5.1×10^5	1.2×10^5
Sub-ore, MT/yr	5.1×10^5	1.2×10^5
Overburden, MT/yr	4.0×10^7	6.0×10^6

Mining Parameters (3.3.1)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Mining days per year	330	330
Mine life, yr	17	17
Ore stockpile residence time, days	41	41
Overburden management	Case 2*	Case 2*

*Case 2--Backfilling concurrent with mining - assumes 7 pits opened in 17-yr. mine life and the equivalent of one-pit overburden (2.4 yr. production) remains on the surface.

Ore Parameters (3.3.1.2)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Average grade, percent U_3O_8	0.1	0.1
Th-232 concentration, pCi/g	10	10
Activity ratio (dust/ore)	2.5	2.5
Mineralogy	Sandstone	Sandstone
Density, MT/m ³	2.0	2.0
Surface area of stockpile, m ²	6,200	3,590
Area of pad, m ²	5,300	3,340
Stockpile height, m	9.2	3.1
Thickness of ore zone, m	12	12

Sub-Ore Parameters (3.3.1.3)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Average grade, percent U_3O_8	0.015	0.015
Th-232 concentration, pCi/g	2	2
Activity ratio (dust/sub-ore)	2.5	2.5
Mineralogy	Sandstone	Sandstone
Density, MT/m ³	2.0	2.0
Surface area of stockpile, m ²	120,000	36,000
Stockpile height, m	30	30
Area of pad, hectares	11	3

Overburden Parameters (3.3.1.1)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Average grade, percent U_3O_8	0.0020	0.0020
Th-232 concentration, pCi/g	1	1
Mineralogy	Sedimentary	Sedimentary
Density, MT/m ³	2.0	2.0
Surface area of dump, m ²	1.1×10^6	3.5×10^5
Dump height, m	65	30
Area of terrain, hectares	104	33

Wastewater Discharge Parameters (3.3.2.2)

<u>Parameters (mg/l except as noted)</u>	<u>Average Mine</u>
Discharge volume, m ³ /min	2.94 (Assumed value of 3.0)
Total uranium	0.07
Radium-226, pCi/l (a)	0.41
Total suspended solids	20.88
Sulfate ^(b)	175
Zinc	0.071
Cadmium	0.004
Arsenic	0.005

(a) Concentration of Ra-226 and its daughters are reduced to 10% of the amount actually released due to irreversible sorption and precipitation.

(b) Concentration of sulfate is reduced to 20% of the amount actually released due to irreversible sorption and precipitation.

Airborne Source Terms (3.3.4)

Section 3.3.4 identifies and describes potential sources of airborne contamination at surface mines. The principal sources are dusts produced by mining operations and wind erosion and Rn-222 released by exposed uranium in the pit and overburden, sub-ore, and ore piles. The tables of Section 3.3.4 present the average annual emissions of contaminants from these sources during active mining.

<u>Source</u>	<u>Table</u>
Combustion Products	3.30
Vehicular Dusts	3.32
Dust from Mining Activities	3.33
Wind Suspended Dust	3.34
Rn-222 Emissions	3.35

4.2 Underground Mine

The model underground mine, defined in Section 3.4 as the "average large mine," will be located in New Mexico. However, to determine the total impact of all 305 underground uranium mines in the United States we used the parameters developed in Section 3.4 for the "average mine."

Production Parameters (1.3.1, 3.4.1)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Ore, MT/yr	2×10^5	1.8×10^4
Sub-ore, MT/yr	2×10^5	1.8×10^4
Waste rock, MT/yr	2.2×10^4	2.0×10^3

Mining Parameters (3.4.1)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Mining days per year	330	330
Mine life, yr	17	17
Ore stockpile residence time, days	41	41
Waste rock management	No backfill	No backfill

Ore Parameters (3.4.1.2)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Average grade, percent U_3O_8	0.10	0.10
Th-232 concentration, pCi/g	10	10
Activity ratio (dust/ore)	2.5	2.5
Mineralogy	Sandstone	Sandstone
Density, MT/m ³	2.0	2.0
Surface area of stockpile, m ²	5,800	680
Stockpile height, m	3.1	3.1
Area of pad, m ²	5,480	620

Sub-Ore Parameters (3.4.1.3)

<u>Parameter</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Average grade, percent U_3O_8	0.035	0.035
Th-232 concentration, pCi/g	2	2
Activity ratio (dust/sub-ore)	2.5	2.5
Mineralogy	Sandstone	Sandstone
Density, MT/m ³	2.0	2.0
Surface area of dump, m ²	104,900	18,800
Dump height, m	12	6
Area of pad, m ²	99,400	17,700

Waste Rock Parameters (3.4.1.1)

<u>Parameters</u>	<u>Average Large Mine</u>	<u>Average Mine</u>
Average grade, percent U_3O_8	0.0020	0.0020
Th-232 concentration, pCi/g	1	1
Mineralogy	Sedimentary	Sedimentary
Density, MT/m ³	2.0	2.0
Surface area of dump, m ²	14,100	2,700
Dump height, m	12	6
Area of terrain, m ²	12,800	2,450

Wastewater Discharge Parameters (3.4.2.2)

<u>Parameter (mg/l except as noted)</u>	<u>Average Mine</u>
Discharge volume, m ³ /min	2.78 (assume value of 2.0)
Total Uranium	1.41
Radium-226, pCi/l (a)	1.37
Lead-210, pCi/l (a)	1.46
Total suspended solids	27.8
Sulfate ^(b)	116
Zinc	0.043
Barium	0.81
Cadmium	0.007
Arsenic	0.012
Molybdenum	0.29
Selenium	0.076

(a) Concentrations of Ra-226 and its daughters are reduced to 10 percent of the amount actually released due to irreversible sorption and precipitation.

(b) Concentrations of sulfate are reduced to 20 percent of the amount actually released due to irreversible sorption and precipitation.

Airborne Source Terms (3.4.4)

Section 3.4.4 identifies and describes potential sources of airborne contamination at underground mines. The principal sources are contaminated dusts due to mining operations and wind erosion and Rn-222 that is released from the mine exhaust vents during mining and from waste rock, sub-ore, and ore pile surfaces. Average annual emissions of contaminants from these sources during active mining operations are presented in the following tables of Section 3.4.4.

<u>Source</u>	<u>Table</u>
Combustion Products	3.52
Vehicular Dusts	3.56
Dust from Mining Activities	3.54
Wind Suspended Dust	3.55
Rn-222 Emissions	3.51

4.3 In Situ Leach Mine

The following parameters are for a model (hypothetical) in situ solution mine as defined in Section 3.5:

1. Size of deposit = 52.6 hectares
2. Average thickness of ore body = 8 m
3. Average ore grade = 0.06 percent U_3O_8
4. Mineralogy = Sandstone
5. Ore density = 2 MT/m³
6. Ore body depth = 153 m
7. Mine life = 10 years (2-yr leach period in each of 5 sectors)
8. Well pattern = 5 spot
 - Injection wells = 260
 - Production wells = 200
 - Monitoring wells = 80
9. Annual U_3O_8 production = 227 MT
10. Uranium leaching efficiency = 80 percent
11. Lixiviant = Alkaline
12. Lixiviant flow capacity = 2,000 ℓ /min
13. Lixiviant bleed = 50 ℓ /min (2.5 percent)
14. Uranium in Lixiviant = 183 mg/ ℓ
15. Calcite.(CaCO₃) removal required = 2 kg calcite per kg U_3O_8

Data were insufficient to estimate aqueous releases of contaminants from these type mines. However, since these facilities are planned to operate with no aqueous discharges, releases of contaminants via this pathway, except for possible excursions, should be small. Annual releases of contaminants to the atmosphere were computed in Section 3.5.3 for the model mine and listed in Table 3.59. These estimated annual airborne releases will be used to compute dose and indicate adverse health effects that might be associated with in situ leach mining.

4.4 Inactive Surface Mine

The model inactive surface mine will be located in Wyoming. It is defined in Section 3.7.1. The model mine parameters are listed below.

Mine Parameters

1. Period of active mining = 17 years
2. Total waste rock production = 8.88×10^5 MT
3. Total ore production = 3.59×10^4 MT
4. Density of ore and waste rock = 2.0 MT/m^3
5. Size of abandoned pit:
 Volume = $4.62 \times 10^5 \text{ m}^3$
 Ground surface area = $2.03 \times 10^4 \text{ m}^2$
 Pit bottom area = $6.00 \times 10^3 \text{ m}^2$
 Depth = 36.7 m
6. Surface area and composition of waste rock pile =
 $6.33 \times 10^4 \text{ m}^2$ uniformly covered to a depth of
 0.36 m with sub-ore
7. Reclamation = none

Airborne Source Terms

Sections 3.7.1.1 and 3.7.1.2 identify and describe potential sources of airborne contamination at inactive surface uranium mines. The principal sources are contaminated, wind-suspended dust from the waste rock pile and Rn-222 released from exposed ore and sub-ore bearing surfaces in the pit and the waste rock pile. Tables 3.70 and 3.74 show average annual emissions of contaminants from these sources.

4.5 Inactive Underground Mine

The model inactive underground mine will be located in New Mexico. It is defined in Section 3.7.2, and its parameters are listed below.

Mine Parameters

1. Period of active mining = 15 yrs
2. Total waste rock production = 1.00×10^4 MT
3. Total ore production = 3.14×10^4 MT
4. Density of ore and waste rock = 2.0 MT/m^3
5. Surface area and composition of waste rock pile =
 $4.08 \times 10^3 \text{ m}^2$ uniformly covered to a depth of
0.78 m with sub-ore
6. Mine entrance and exhaust vents not sealed

Airborne Source Terms

Sections 3.7.2.1 and 3.7.2.2 identify and define potential sources of airborne contamination at inactive underground uranium mines. The principal sources are contaminated, wind-suspended dust from the waste rock pile and Rn-222 released from the unsealed mine entrance and exhaust vents and the waste rock pile. Tables 3.76 and 3.77 list average annual emission of contaminants from these sources.

SECTION 5

POTENTIAL PATHWAYS

5.0 Potential Pathways

5.1 General

5.1.1 Vegetation

Airborne particulate radioactivity may be deposited directly on the edible foliar surfaces of crops or on the soil and then migrate through the soil into the plant's root system and into an edible crop. Such crops may be consumed directly by man or by animals which are ultimately consumed by man. The use of contaminated water (either groundwater or surface) to irrigate crops may also lead to the ingestion of radionuclides from either the direct consumption of the crop or the crop-to-animal-to-man pathway.

The reconnaissance surveys of some inactive uranium mine sites indicated that no crops for human consumption were being farmed at or near any of the sites. Although the potential for man's ingestion of radionuclides in edible crops due to the direct deposition or the root uptake of either airborne particulates or contaminated mine water is a greater possibility near the active mines, farming in such areas is not extensive.

Almost every inactive and active mine site visited had range cattle and/or sheep grazing on the natural vegetation growing at the site; hence, the possible consumption of such animals could be a potential pathway for man's ingestion of radionuclides released into the environment surrounding the mine sites.

5.1.2 Wildlife

There are numerous species of mammals, birds, reptiles, and amphibians at both active and inactive uranium mine sites. Though mining may destroy their natural habitat, there are no significant radiological impacts on wildlife in these areas. Dewatering and drainage from active mines sometimes create ponds or streams that may be used by migratory waterfowl and local wildlife as a source of water, but, when mining is completed, the ponds dry up, probably without leaving any permanent or significant radiological impact on wildlife. The small lakes formed in inactive surface mine pits, however, may remain for a long period of time and have a significant environmental impact. It would be expected that sedimentation and eutrophication of the lakes would progressively diminish the impact with time by reducing the contact of ore bodies with the biosphere. The potential food pathway of animal-

to-man via wildlife hunting at these sites is also minimal. Hunting is poor and hunting restrictions are usually observed at the mine sites.

5.1.3 Land Use

Most uranium mining activities have been conducted in areas away from population centers. Most mines are located on private property or are on Federal lands such as national forests. The predominant land use is as rangeland (or forest) and only minor areas are cropland. The fraction of land used for vegetable crop production for Wyoming and New Mexico is 1.59×10^{-3} and 1.38×10^{-3} , respectively. This fraction is based on the assumption that the statewide fractions apply to uranium mining areas within each state. Average population densities are typically rural, i.e., less than one person per 2.6 km^2 .

5.1.4 Population Near Mining Areas

Uranium mines occur in clusters throughout many western states and are somewhat scattered throughout the eastern states. In order to estimate the number of persons residing within 50 miles (80km) of a mine, we used county populations where there either is or has been mining. Table 5.1 lists the states and their respective mining counties plus the numbers of inactive and active surface and underground uranium mines in each county. We derived the county population statistics from U.S. Department of Commerce census data (DOC78), which are January 1, 1975 estimates. The county areas were obtained from the same reference.

The area, $20,106 \text{ km}^2$, within a circle with a radius of 80 km usually exceeds the area of most counties. Because of this, the number of persons residing within 80 km of a mine will be underestimated using county population statistics. In other words, we consider the estimates of populations within the mining regions to be somewhat low.

Persons residing in a mining area are likely to be exposed from more than one mine because of the aforementioned clustering. To account for this, Table 5.1 lists the product (person-mines) for both active and inactive uranium mines. The total number of person-mines for inactive mines is approximately 82,000,000 persons. The total number of person-mines for active mines is approximately 14,000,000 persons. The combined equivalent population exposed to inactive and active uranium mining is approximately 96,000,000 persons.

Table 5.1 Number of uranium mines and population statistics for counties containing uranium mines

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County Population (persons)	Person-Mines	
		Inactive	Active				Inactive	Active
Alaska	Southeast ^(a)	1	0	0.03	44,501	1,282	1,282	0
Arizona	Apache	140	0	1.1	28,930	32,304	4,522,560	0
	Cochise	2	0	3.8	16,203	61,918	123,836	0
	Coconino	113	0	1.0	48,019	48,326	5,460,838	0
	Gila	18	0	2.4	12,297	29,255	526,590	0
	Graham	1	0	1.4	11,961	16,578	16,578	0
	Maricopa	3	0	41.	23,711	971,228	2,913,684	0
	Mohave	5	0	0.76	34,232	25,857	129,285	0
	Navajo	35	1	2.3	25,666	59,649	2,088,715	59,649
	Pima	2	1	19.	23,931	443,958	887,916	443,958
	Santa Cruz	3	0	4.3	3,227	13,966	41,898	0
	Yavapai	3	0	1.8	20,956	37,005	111,015	0
California	Imperial	2	0	6.8	10,984	74,492	148,984	0
	Inyo	1	0	0.77	26,237	17,259	17,259	0
	Kern	6	0	17	21,113	349,874	2,099,244	0
	Lassen	2	0	1.4	11,816	16,796	33,592	0

Table 5.1 (Continued)

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County Population (persons)	Person-Mines	
		Inactive	Active				Inactive	Active
California	Madera	1	0	7.5	5,556	41,519	41,519	0
	Mono	1	0	0.51	7,840	4,016	4,016	0
	Riverside	5	0	25.	18,586	456,916	2,284,580	0
	San Bernardino	3	0	14	52,103	696,871	2,090,613	0
	Sierra	1	0	1.2	2,481	2,842	2,842	0
	Tuolumne	1	0	4.6	5,832	25,996	25,996	0
Colorado	Boulder	7	0	68	1,937	131,889	923,223	0
	Clear Creek	4	0	4.8	995	4,819	19,276	0
	Custer	3	0	0.59	1,909	1,120	3,360	0
	Dolores	6	0	0.62	2,657	1,641	9,846	0
	Eagle	2	0	1.7	4,353	7,498	14,996	0
	El Paso	1	0	42	5,587	235,972	235,972	0
	Fremont	25	0	6.6	4,022	26,545	663,625	0
	Garfield	10	0	2.3	7,759	17,845	178,450	0
	Gilpin	4	0	5.0	383	1,915	7,660	0
	Grand	4	0	0.86	4,802	4,107	16,428	0

Table 5.1 (Continued)

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County Population (persons)	Person-Mines	
		Inactive	Active				Inactive	Active
Colorado	Gunnison	1	0	1.2	8,339	10,006	10,006	0
	Hinsdale	1	0	0.19	2,729	519	519	0
	Huerfano	2	0	1.6	4,077	6,590	13,180	0
	Jefferson	13	1	120	2,028	235,368	3,059,784	235,368
	La Plata	3	0	5.4	4,358	23,533	70,599	0
	Larimer	5	0	17	6,762	114,954	574,770	0
	Mesa	185	20	7.3	8,549	62,407	11,545,295	1,248,140
	Moffat	18	3	0.77	12,284	9,459	170,262	28,377
	Montezuma	6	1	2.7	5,423	14,642	87,852	14,642
	Montrose	479	63	3.5	5,796	20,286	9,716,994	1,278,018
	Park	7	0	0.77	5,599	4,311	30,177	0
	Pitkin	1	0	3.5	2,520	8,820	8,820	0
	Pueblo	1	0	20	6,228	124,560	124,560	0
	Rio Blanco	26	0	0.77	8,451	6,507	169,182	0
	Saguache	13	1	0.39	8,142	3,175	41,275	3,175
	San Juan	2	0	0.77	1,012	779	1,558	0

Table 5.1 (Continued)

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County Population (persons)	Person-Mines	
		Inactive	Active				Inactive	Active
Colorado	San Miguel	339	25	0.77	3,322	2,557	866,823	63,925
	Teller	3	0	3.9	1,432	5,584	16,752	0
Idaho	Custer	5	0	0.23	12,766	2,967	14,835	0
	Lemhi	1	0	0.39	11,862	6,395	6,395	0
Montana	Broadwater	1	0	0.82	3,090	2,526	2,526	0
	Carbon	11	0	1.5	5,325	7,797	85,767	0
	Fallon	1	0	0.96	4,229	4,050	4,050	0
	Hill	1	0	2.3	7,581	17,358	17,358	0
	Jefferson	3	0	1.5	4,278	6,839	20,517	0
	Madison	1	0	0.55	9,138	5,014	5,014	0
Nevada	Clark	2	0	16.2	20,393	330,714	661,428	0
	Elko	3	0	0.31	44,452	13,958	41,874	0
	Humboldt	1	0	0.25	25,128	6,375	6,375	0
	Lander	2	0	0.39	14,558	2,992	5,984	0
	Lincoln	2	0	0.19	27,114	2,647	5,294	0
	Lyon	2	0	1.9	5,257	10,508	21,016	0

Table 5.1 (Continued)

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County Population (persons)	Person-Mines	
		Inactive	Active				Inactive	Active
Nevada	Mineral	2	0	0.71	9,751	7,051	14,102	0
	Nye	1	0	0.12	46,786	5,599	5,599	0
	Washoe	6	0	8.9	16,487	144,750	868,500	0
New Jersey	Sussex	1	0	73	1,364	99,299	99,299	0
New Mexico	Catron	4	0	0.12	17,863	2,198	8,792	0
	Dona Ana	1	0	7.1	9,852	69,773	69,773	0
	Grant	3	0	2.1	10,282	22,030	66,090	0
	Harding	1	0	0.25	5,527	1,348	1,348	0
	Hidalgo	1	0	0.53	8,927	4,734	4,734	0
	McKinley	73	35	3.5	14,138	49,483	3,612,259	1,731,905
	Mora	1	0	0.93	5,025	4,673	4,673	0
	Quay	3	0	1.5	7,446	10,903	32,709	0
	Rio Arriba	8	0	1.9	15,133	28,752	230,016	0
	Sandoval	3	0	2.3	9,619	22,123	66,369	0
	San Juan	41	0	4.6	14,245	65,527	2,686,607	0
	San Miguel	3	0	1.8	12,279	21,951	65,853	0
	Santa Fe	2	0	13	4,926	64,038	128,076	0

Table 5.1 (Continued)

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County		
		Inactive	Active			Population (persons)	Person-Mines Inactive	Person-Mines Active
New Mexico	Sierra	6	0	0.67	10,790	7,189	43,134	0
	Socorro	7	0	0.57	17,102	9,763	68,341	0
	Taos	1	0	3.0	5,843	17,516	17,516	0
	Valencia	19	4	3.1	14,649	45,411	862,809	181,644
North Dakota	Billings	9	0	0.39	2,950	1,153	10,377	0
	Slope	1	0	0.39	3,172	1,360	1,360	0
	Stark	3	0	5.8	3,408	19,650	58,950	0
Oklahoma	Caddo	2	0	8.8	3,294	28,931	57,862	0
	Custer	1	0	8.3	2,538	21,040	21,040	0
Oregon	Crook	1	0	1.3	7,705	9,985	9,985	0
	Lake	2	0	0.34	21,318	7,158	14,316	0
South Dakota	Butte	3	0	1.3	5,827	7,825	23,475	0
	Custer	10	0	1.2	4,032	5,196	51,960	0
	Fall River	93	0	1.9	4,514	8,066	750,138	0
	Harding	28	0	0.39	6,946	1,879	52,612	0
	Lawrence	2	0	8.4	2,072	17,453	34,906	0
	Pennington	5	0	8.3	7,198	59,349	296,745	0

Table 5.1 (Continued)

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County Population (persons)	Person-Mines	
		Inactive	Active				Inactive	Active
Texas	Briscoe	2	0	1.2	2,264	2,794	5,588	0
	Burnet	1	0	4.4	2,577	11,420	11,420	0
	Crosby	1	0	3.9	2,359	9,085	9,085	0
	Garza	6	0	2.8	2,367	6,611	39,666	0
	Gonzales	2	0	5.8	2,735	16,342	32,684	0
	Karnes	23	10	6.6	1,963	12,955	297,965	129,550
	Live Oak	6	5	2.3	2,732	6,453	38,718	32,265
Utah	Beaver	9	1	0.77	6,692	5,152	46,368	5,152
	Box Elder	1	0	1.9	14,512	28,129	28,129	0
	Duchesne	4	0	1.5	8,430	12,645	50,580	0
	Emery	186	18	0.39	11,497	4,483	833,838	80,694
	Garfield	131	15	0.39	13,359	5,210	682,510	78,150
	Grand	164	17	0.77	9,536	7,342	1,204,088	124,814
	Iron	1	0	1.4	8,547	12,177	12,177	0
	Juab	4	0	0.52	8,837	4,574	18,296	0
	Kane	3	0	0.39	10,111	3,943	11,829	0

Table 5.1 (Continued)

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County		
		Inactive	Active			Population (persons)	Person-Mines Inactive	Person-Mines Active
Utah	Piute	10	0	0.77	1,952	1,503	15,030	0
	San Juan	241	24	0.77	19,961	15,369	3,703,929	368,856
	Sevier	2	0	2.3	4,996	11,490	22,980	0
	Uintah	14	0	1.5	11,621	17,431	244,034	0
	Washington	6	0	2.7	6,285	16,969	101,814	0
	Wayne	32	0	0.39	6,438	2,510	80,320	0
Washington	Pend Oreille	3	0	1.9	3,631	7,361	22,083	0
	Spokane	9	0	67	4,553	306,338	2,757,042	0
	Stevens	1	2	3.5	6,425	22,489	22,489	44,978
Wyoming	Albany	4	0	2.3	11,002	25,304	101,216	0
	Big Horn	9	0	1.5	8,176	12,264	110,376	0
	Campbell	55	0	1.2	12,318	14,781	812,955	0
	Carbon	16	3	0.77	20,473	15,764	252,224	47,292
	Converse	31	5	0.77	11,087	8,536	264,616	42,680
	Crook	23	0	0.77	7,464	5,747	132,181	0
	Fremont	65	13	1.2	23,817	28,580	1,857,700	371,540

Table 5.1 (Continued)

State	County	Number of Uranium Mines		Population Density (persons/km ²)	County Area (km) ²	County Population (persons)	Person-Mines	
		Inactive	Active				Inactive	Active
Wyoming	Johnson	15	0	0.39	10,813	4,217	63,255	0
	Natrona	16	2	3.9	13,835	53,956	863,296	107,912
	Niobrara	13	0	0.39	6,770	2,640	34,320	0
	Sublette	1	0	0.39	12,564	4,899	4,899	0
	Sweetwater	4	2	1.2	27,011	32,413	129,652	64,826
	Washakie	2	0	1.5	5,858	8,787	17,574	0
	Weston	1	0	1.0	6,234	6,307	6,307	0
Average Population				Total County		Total Person-Mines		Total Person-Mines
Density				Area (km) ²		Population		(Inactive)
4.4 persons/km ²				1,492,136		6,625,099		82,327,885
								14,035,161

Note.--Population statistics from (DOC78).

(a) Congressional District.

5.1.5 Population Statistics of Humans and Beef Cattle

Table 5.2 lists some population statistics for humans in New Mexico and Wyoming, humans in all uranium mining states, and beef cattle in New Mexico and Wyoming.

Table 5.2 Population statistics for humans and beef cattle

<u>Total Human and Beef Cattle Population Within 80 km Radius of Mines</u>			
	New Mexico	Wyoming	All Uranium Mining States
Human	447,412	224,195	6,625,099
Beef cattle	753,000	905,000	-----

Average Human and Beef Cattle Population Densities Within 80 km Radius of Uranium Mines (number/km²)^(a)

Human	2.4	1.3	4.4
Beef Cattle	4.1	5.1	----

^(a) Areas taken from Table 5.1: New Mexico = 183,646 km²; Wyoming = 177,422 km², and the total county area = 1,492,136 km².

5.2 Prominent Environmental Pathways and Parameters for Aqueous Releases

From a computer code prepared within EPA, we calculated annual committed dose equivalents to individuals and annual collective dose equivalents to a population for these assessments. Table 5.3 lists the aqueous pathways that were initially considered potential pathways of exposure. As indicated in Table 5.3, these pathways result in computation of dose equivalents due to inhalation, ingestion, ground surface exposure, and air submersion. For above surface crop ingestion, milk ingestion, and beef ingestion (pathways 3, 4, and 5), we considered only uptake through the plant root systems to predict

concentrations of radionuclides in crops, since essentially all irrigation is ditch irrigation. Appendix J contains a detailed explanation of the environmental transport and dosimetry models used in these analyses.

The maximum individual for the aquatic pathways is the individual at maximum risk. He is exposed to radionuclides discharged in mine effluent through pathways 2 through 10 of Table 5.3. The water contributing radionuclides to these pathways comes from a creek into which a mine discharges. The average individual is exposed to the average risk of all persons included in the population of the assessment area. He is exposed to radionuclides discharged in mine effluent through pathways 2 through 8 and 10 of Table 5.3. The water contributing radionuclides to these pathways is taken from the regional river after the creek water has been diluted in this river. The population considered in the assessment of the aquatic pathways is obtained by multiplying the regional assessment area size by the population density within this area. This assessment area contains the drainage basin for the mine effluent stream, the creek and the regional river discussed in defining the maximum and average individuals.

5.2.1 Individual Committed Dose Equivalent Assessment

Section 6 of this report contains the computed dose equivalents to the maximum individual and to the average individual. For the maximum individual, we included all pathways in Table 5.3 except drinking water (pathway 1). It is known that the releases to the aquatic environment occur through discharge of mine water to surface streams. Potentially, drinking water could be one of the most significant pathways for the maximum individual dose equivalents, if surface water containing mine wastes was drunk. However, it appears that all drinking water for both the New Mexico and the Wyoming sites comes from wells (Robert Kaufmann, 1979, U.S. Environmental Protection Agency, Las Vegas, NV, personal communication). Thus, the only way mine discharges can enter human drinking water is by percolating through the soil. Since we do not know the soil chemistry for these sites well enough to predict the ion-exchange parameters for the soil, we can not predict, realistically, the quantity of mine-related radionuclides that would reach the groundwater. We expect that these ion-exchange factors would be large for several of the radionuclides considered in these analyses and that groundwater concentrations of radionuclides discharged in mine water

would be quite small compared to concentrations in the surface water downstream from the mines. Further study is needed before dose equivalents for the maximum individual by drinking groundwater can be adequately addressed.

The following are other assumptions used to calculate maximum individual dose equivalents:

1. Ground surface concentrations of radionuclides (used for pathways 6 through 8) are for 8.5 years, the assumed midpoint of mine life. The assumed period of mine operation is 17 years. The organ annual dose equivalents for the external surface exposure pathway are based on the ground concentrations after the 8.5 years buildup time.
2. For inhaled or ingested radionuclides, the dose equivalents are the annual committed dose equivalents that will be accumulated over 70 years after intake for an adult.

We calculated dose equivalents to the average individual in the assessment area by taking the population dose equivalents (discussed in Subsection 5.2.2) and dividing by the population living in the area.

Table 5.3 Aquatic environmental transport pathways initially considered

Pathway No.	Pathway
1	Drinking water ingestion
2	Freshwater fish ingestion
3	Above surface crops ingestion - irrigated cropland
4	Milk ingestion - cows grazing on irrigated pasture
5	Beef ingestion - cows grazing on irrigated pasture
6	Inhalation - material resuspended which was deposited during irrigation
7	External dose due to ground contamination by material originally deposited during irrigation
8	External dose due to air submersion in resuspended material originally deposited during irrigation
9	Milk ingestion - cows drinking contaminated surface water
10	Beef ingestion - cows drinking contaminated surface water

5.2.2 Collective (Population) Dose Equivalent Assessment

For the population dose equivalent assessment calculations, we concluded that the pathways of concern are pathways 2, 3, 4, 5, 6, 7, 8, and 10 of Table 5.3 (detailed discussion in Appendix J, subsection J2). The size of the assessment areas for New Mexico is 19,037 km² and 13,650 km² for Wyoming. We used the following considerations to calculate population dose equivalents for the assessment area:

1. Ground surface concentrations of radionuclides are for 8.5 years, the assumed midpoint mine life. (The period of mine operation is 17 years.) The organ annual collective dose equivalent rates for the external surface exposure pathway are based on the ground concentrations after the 8.5 year buildup time.
2. For inhaled or ingested radionuclides, the dose equivalents are the annual collective dose equivalents that will be accumulated over the 70 years after intake for adults.
3. The population distributions around the sites are based on estimates by county planners (John Zaboroc, 1979, Converse Area Planning Office, Douglas, Wyoming, personal communication) and agricultural personnel (Tony Romo, 1979, Valencia County Agent, Los Lunas, New Mexico, personal communication) for 1979. The populations, assumed to remain constant in time, were estimated to be 16,230 and 64,950 persons in the Wyoming and New Mexico assessment areas, respectively.
4. Average agricultural production data for the county which contains a major portion of the assessment area are used.
5. The population in the assessment area eats food from the assessment area. We assume that any imported food is free of radionuclides.

As mentioned previously, Appendix J contains the details regarding the models and values for parameters used in these analyses.

5.3 Prominent Environmental Pathways and Parameters for Atmospheric Releases

We used the AIRDOS-EPA (Mo79) computer code to calculate radionuclide air and ground concentrations, ingestion and inhalation intakes, and working level exposures; and we used the DARTAB (Be80) computer code to calculate dose and risk from the AIRDOS-EPA intermediate output using dose and risk factors from the RADRISK (Du80) computer code. We calculated working levels associated with Rn-222 emissions assuming that Rn-222 decay products were 70 percent in equilibrium with Rn-222, a value considered representative of indoor exposure conditions (Ge78). Appendix K contains a detailed discussion of the application of the AIRDOS-EPA and RADRISK computer codes.

Figure 5.1 shows the general airborne pathways evaluated for uranium mines. We calculated doses due to air immersion, ground surface exposure, inhalation, and ingestion of radionuclides, but we did not address the resuspension pathway, since the AIRDOS-EPA code did not provide a method for calculating resuspended air concentrations or subsequent redeposition to the ground surface. We used the modification to the AIRDOS-EPA computer code made by Nelson (Ne80) to include the effect of environmental removal of radioactivity from the soil. For ingestion, transfers associated with both root uptake and foliar deposition on food and forage are considered.

5.3.1 Individual Committed Dose Equivalent Assessment

We assessed the maximum individual on the following basis:

1. The maximum individual for each source category is intended to represent an average of the individuals living close to each model uranium mine. The individual is assumed to be located about 1600 meters from the center of the model site.
2. Ground surface concentrations of radionuclides used in the assessment are those that would occur during the midpoint of the active life of the model uranium mine. Buildup times used in the assessment are 8.5 years for active surface and underground mines, 5 years for the in situ leach mine, and 26.5 years for the inactive surface and underground mines. The 26.5-year buildup time for the inactive mines is chosen to represent the midpoint of the 53-year exposure time that a resident living a lifetime in the region around the model mine is estimated to experience. The organ dose equivalent rates for the external surface exposure pathway are based on

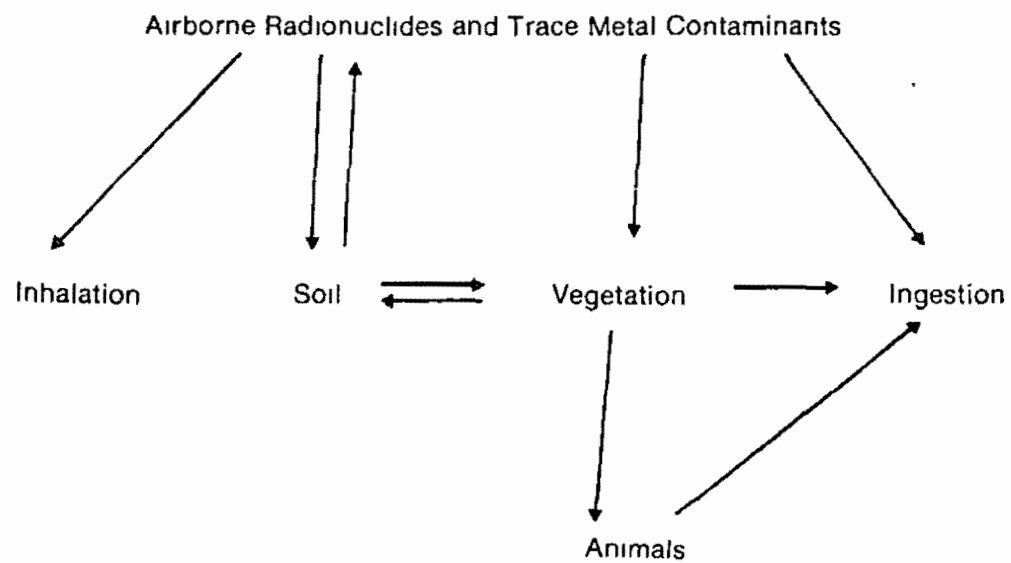


Figure 5 1 Potential airborne pathways in the vicinity of uranium mines.

the concentrations for the indicated buildup time.

3. For inhaled or ingested radionuclides, the dose equivalent rates are actually the 70-year committed dose equivalent rates for an adult receptor, i.e., the internal dose equivalent that would be delivered up to 70 years after an intake. The individual dose equivalent rates in the tables are in units of mrem/yr.
4. The individual is assumed to home grow a portion of his or her diet consistent with the rural setting for each model uranium mine site. Appendix K contains the actual fractions of home-produced food consumed by individuals for the model mine sites. The portion of the individual's diet that was not locally produced is assumed to be imported and uncontaminated by the assessment source.

5.3.2 Collective (Population) Dose Equivalent Assessment

The collective dose equivalent assessment to the population out to 80 km from the facility under consideration is performed as follows:

1. The population distribution around the model mine sites is based on the 1970 census. The population is assumed to remain constant in time.
2. Ground surface concentrations and organ dose equivalent rates for the external surface exposure pathway (as for the individual case) are those that would occur over the active life of the model mine.
3. Average agricultural production data for the state in which the model uranium mine is located are assumed.
4. The population in the assessment area eats food from the assessment area to the extent that the calculated production allows, and any balance is assumed to be imported without contamination by the assessment source.
5. Seventy-year committed dose equivalent factors for an adult receptor (as for the individual case) are used for ingestion and inhalation.

5.4 Mine Wastes Used In the Construction of Habitable Structures

Using uranium mine wastes under or around habitable structures or building habitable structures on land contaminated with uranium mine wastes can result in increased radiation exposures to individuals occupying these structures. The radium-226 present in these wastes elevates the concentrations of radon-222 and its decay products and produces increased gamma radiation inside these structures. The health risk to individuals occupying these structures is generally much greater from inhaling radon-222 decay products than the risk received from gamma radiation.

Radon-222, formed from the decay of radium-226, is an inert gas that diffuses through the soil and migrates readily through foundations, floors, and walls and accumulates in the inside air of a structure. Breathing radon-222 and its short-lived decay products (principally polonium-218, bismuth-214, and polonium-214) exposes the lungs to radiation.

The radon-222 decay product concentration (working level) inside a structure from radon-222 gas diffusing from underlying soil is extremely variable and influenced by many complex factors. These would include the radium-226 concentration of the soil, the fraction of radon-222 emanating from the soil, the diffusion coefficient of radon-222 in soil, the rate of influx of radon-222 into the structure, the ventilation rate of the structure, and the amount of plate-out (adsorption) of radon-222 decay products on inside surfaces.

The potential risks of fatal lung cancer that could occur to individuals living in homes built on land contaminated by uranium mine wastes have been estimated using measurements and calculational methodology relating radon-222 decay product concentrations inside homes to the radium-226 concentrations in outside soil (He78, Wi78). These estimates are shown in Section 6.1.5.

5.5 References

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SECTION 6

HEALTH AND ENVIRONMENTAL EFFECTS

6.0 Health and Environmental Effects

6.1 Health Effects and Radiation Dosimetry

6.1.1 Radioactive Airborne Emissions

We used data on radioactive emissions (Section 3) to estimate the public health impact of these emissions. Our assessments include estimates of the following radiation exposures and health risks:

1. Dose equivalent rates and working level exposures to the most exposed individuals (maximum individual) and to the average exposed individuals in the regional population (average individual)
2. Collective dose equivalent rates and working level exposures to the regional population
3. Lifetime fatal cancer risks to the maximum and average individuals in the regional population
4. Genetic effect risk to the descendants of the maximum and average individuals in the regional population
5. The number of fatal cancers committed in the regional population per year of model mine operation
6. The number of genetic effects committed to the descendants of the regional population per year of model mine operation

The somatic health impact risks estimated in this report are for fatal cancers only. For whole body exposure, the risk of nonfatal cancer is about the same or slightly less than for fatal cancer. Thus, for whole body doses, it is conservatively estimated that one nonfatal cancer could occur for each additional fatal cancer. The somatic health impact for the regional population (additional cancers per year) is calculated at equilibrium for continuous exposure and this is equal to the additional cancers committed over all time per year of exposure; thus we used the term committed additional cancers (see Appendix L).

The genetic effect risks estimated in this report are for effects in descendants of an irradiated parent or parents. Genetic effects per year in the regional population due to radionuclide releases from the mines are calculated for an equilibrium exposure situation. The calculated genetic effects per year at equilibrium is equal to the genetic effects committed over all time from one year exposure. Thus, the calculated additional

Table 6.1 Annual release rates (Ci) used in the dose equivalent and health effects computations for active uranium mines

Classification	Location	Average Surface Mine ^(a)			Average Large Surface Mine ^(a)		
		U	Th	Rn-222	U	Th	Rn-222
Mining activities	Pit/mine site	4.3E-3	2.2E-4	1.99E+2	2.57E-2	1.44E-3	7.97E+2
Ore	Pile site	1.01E-2	1.42E-4	4.2E+1	4.42E-2	6.20E-4	9.6E+1
Sub-ore	Pile site	4.2E-4	8.4E-6	5.0E+1	1.51E-3	3.00E-5	1.66E+2
Overburden/ waste rock	Pile site	2.25E-3	1.50E-4	4.0E+1	1.34E-2	8.94E-4	2.02E+2
Vehicular dust	Mining area	9.9E-4	3.7E-4	0	5.86E-3	2.17E-3	0
Total	All sources	1.81E-2	8.90E-4	3.31E+2	9.07E-2	5.15E-3	1.26E+3

(a) Release rates taken from Tables 3.32 to 3.35.

(b) Release rates taken from Tables 3.51 and 3.54 to 3.56.

Table 6.1 (cont.)

Average Underground Mine ^(b)			Average Large Underground Mine ^(b)			In Situ Leach Mine ^(c)		
U	Th	Rn-222	U	Th	Rn-222	U	Th	Rn-222
2.22E-4	2.8E-6	3.08E+2	2.41E-3	3.10E-5	3.42E+3	1.0E-1	0	6.50E+2
9.63E-4	1.35E-5	7.7	1.07E-2	1.50E-4	6.83E+1	N.A. ^(d)	N.A.	N.A.
1.04E-3	8.4E-6	6.1E+1	5.95E-3	4.8E-5	3.38E+2	N.A.	N.A.	N.A.
9.6E-6	6.4E-7	5.0E-1	5.10E-5	3.40E-6	2.6	N.A.	N.A.	N.A.
6.5E-5	2.4E-5	0	1.29E-4	4.80E-5	0	N.A.	N.A.	N.A.
2.30E-3	4.93E-5	3.77E+2	1.92E-2	2.80E-4	3.83E+3	1.0E-1	0	6.50E+2

^(c) Release rates taken from Table 3.59.

^(d) N.A.- Not Applicable.

Note.--Columns labeled U and Th include each daughter of the decay chain in secular equilibrium.

Table 6.2 Annual release rates (Ci) used in the dose equivalent and health effects computations for inactive uranium mines

Location	Surface Mine (a)			Underground Mine (b)		
	U	Th	Rn-222	U	Th	Rn-222
Pit/vents- portals	0	0	8.1	0	0	7.55
Waste rock/ sub-ore pile	1.48E-3	1.1E-5	1.74E+1	2.38E-4	1.7E-6	1.7

(a) Release rates taken from Tables 3.70 and 3.74.

(b) Release rates taken from Tables 3.76 and 3.77.

Note.--Column headings U and Th include each daughter of the decay chain in secular equilibrium.

genetic effects are committed effects to all future generations for one year of exposure to the regional population.

We calculated individually each major source of radionuclide airborne emissions for each model uranium mine site so that we could determine the extent that each source contributed to the total health impact. Tables 6.1 and 6.2 contain the annual release rates for each source classification (or location) that we used to calculate dose equivalent rates and health effects for active and inactive uranium mines.

The estimated annual working level exposures from Rn-222 emissions by the model uranium mines are listed in Table 6.3. The working level exposures presented for the maximum individual are the Rn-222 decay product levels to which an individual would be continuously exposed for an entire year. Working level exposure to the regional population is the sum of the exposures to all individuals in the exposed population from the annual release from the model mine.

We estimated radiological impacts of radioactive airborne emissions from the model uranium mines with the AIRDOS-EPA (Mo79), RADRISK (Du80), and DARTAB (Be80) computer codes. Appendixes K and L contain explanations of our use of these computer codes.

Where emissions for U-238 plus daughters and Th-232 plus daughters were reported (Section 3), a source term for both the parent and important daughters were input into the AIRDOS-EPA code. For example, a reported emission rate of 0.01 Ci/yr of U-238 plus daughters (U in Tables 6.1 and 6.2) would be input into the AIRDOS-EPA code as 0.01 Ci/yr of U-238, 0.01 Ci/yr of U-234, 0.01 Ci/yr of Th-230, 0.01 Ci/yr of Ra-226, 0.01 Ci/yr of Pb-214, 0.01 Ci/yr of Bi-214, 0.01 Ci/yr of Pb-210, and 0.01 Ci/yr of Po-210. A reported emission rate of 0.01 Ci/yr of Th-232 plus daughters (Th in Tables 6.1 and 6.2) would be input into the AIRDOS-EPA code as 0.01 Ci/yr of Th-232, 0.01 Ci/yr of Ra-228, 0.01 Ci/yr of Ac-228, 0.01 Ci/yr of Th-228, 0.01 Ci/yr of Ra-224, 0.01 Ci/yr of Pb-212, 0.01 Ci/yr of Bi-212, and 0.0036 Ci/yr of Tl-208. The Tl-208 source term is approximately one-third that of Bi-212 because of the branching ratio.

The maximum individual, average individual, and population dose equiv-

Table 6.3 Annual working level exposure from radon-222
emissions from model uranium mines

Source	Maximum Individual (WL) ^(a)	Average Individual (WL)	Regional Population (person-WL)
Average Surface Mine	2.3E-4	4.5E-7	6.5E-3
Average Large Surface Mine	8.4E-4	1.7E-6	2.5E-2
Average Underground Mine	4.6E-4	2.1E-6	7.5E-2
Average Large Underground Mine	4.7E-3	2.1E-5	7.6E-1
Inactive Surface Mine	1.8E-5	3.5E-8	5.0E-4
Inactive Underground Mine	1.1E-5	5.1E-8	1.8E-3
In Situ Leach Mine	4.5E-4	8.9E-7	1.3E-2

(a) Working level.

Table 6.4 Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model average surface uranium mine

Organ	Maximum Individual (mrem/yr)	Average Individual (mrem/yr)	Population (person-rem/yr)
Red marrow	2.4	5.4E-3	7.7E-2
Endosteal	3.4E+1	7.5E-2	1.1
Pulmonary	1.2E+1	6.3E-3	9.0E-2
Muscle	5.5E-1	2.0E-3	2.7E-2
Liver	1.6	6.3E-3	9.1E-2
Stomach wall	9.7E-2	8.9E-5	1.3E-3
Pancreas	5.2E-1	1.9E-3	2.7E-2
LLI ^(a) wall	4.6E-1	1.6E-3	2.3E-2
Kidney	4.2	1.8E-2	2.5E-1
Bladder wall	3.0E-1	9.7E-4	1.4E-2
ULI ^(b) wall	2.1E-1	5.2E-4	7.4E-3
SI ^(c) wall	9.4E-2	1.2E-4	1.7E-3
Ovaries	5.1E-1	1.9E-3	2.7E-2
Testes	5.4E-1	1.9E-3	2.7E-2
Spleen	6.4	2.8E-2	4.0E-1
Uterus	5.1E-1	1.9E-3	2.7E-2
Thymus	5.2E-1	1.9E-3	2.7E-2
Thyroid	5.4E-1	1.9E-3	2.7E-2
Weighted mean	4.9	5.5E-3	7.8E-2

(a) Lower large intestine wall.

(b) Upper large intestine wall.

(c) Small intestine wall.

Table 6.5 Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model average large surface uranium mine

Organ	Maximum Individual (mrem/yr)	Average Individual (mrem/yr)	Population (person-rem/yr)
Red marrow	1.35E+1	2.7E-2	3.9E-1
Endosteal	1.9E+2	3.8E-1	5.4
Pulmonary	6.6E+1	3.1E-2	4.5E-1
Muscle	3.0	9.6E-3	1.4E-1
Liver	8.9	3.2E-2	4.6E-1
Stomach wall	5.4E-1	4.5E-4	6.4E-3
Pancreas	3.0	9.6E-3	1.4E-1
LLI wall	2.5	8.2E-3	1.2E-1
Kidney	2.1E+1	9.0E-2	1.3
Bladder wall	1.7	4.9E-3	7.0E-2
ULI wall	1.1	2.6E-3	3.8E-2
SI wall	5.2E-1	6.0E-4	8.6E-3
Ovaries	2.8	9.6E-3	1.4E-1
Testes	3.0	9.6E-3	1.4E-1
Spleen	3.5E+1	1.4E-1	2.0
Uterus	2.8	9.6E-3	1.4E-1
Thymus	2.9	9.6E-3	1.4E-1
Thyroid	3.0	9.6E-3	1.4E-1
Weighted mean	2.7E+1	2.7E-2	3.8E-1

Table 6.6 Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model average underground uranium mine

Organ	Maximum Individual (mrem/yr)	Average Individual (mrem/yr)	Population (person-rem/yr)
Red marrow	5.1E-1	8.3E-4	2.9E-2
Endosteal	7.2	1.2E-2	4.1E-1
Pulmonary	2.9	5.0E-3	1.8E-1
Muscle	1.2E-1	2.3E-4	8.3E-3
Liver	3.5E-1	7.2E-4	2.7E-2
Stomach wall	2.0E-2	2.8E-5	1.0E-3
Pancreas	1.1E-1	2.2E-4	8.0E-3
LLI wall	9.4E-2	1.8E-4	6.5E-3
Kidney	9.1E-1	2.0E-3	7.4E-2
Bladder wall	6.4E-2	1.2E-4	4.4E-3
ULI wall	4.3E-2	7.3E-5	2.7E-3
SI wall	2.0E-2	2.8E-5	1.0E-3
Ovaries	1.1E-1	2.2E-4	8.0E-3
Testes	1.1E-1	2.3E-4	8.0E-3
Spleen	1.4	3.1E-3	1.1E-1
Uterus	1.1E-1	2.2E-4	7.9E-3
Thymus	1.1E-1	2.2E-4	8.0E-3
Thyroid	1.1E-1	2.3E-4	8.1E-3
Weighted mean	1.1	2.0E-3	7.1E-2

Table 6.7 Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model average large underground uranium mine

Organ	Maximum Individual (mrem/yr)	Average Individual (mrem/yr)	Population (person-rem/yr)
Red marrow	4.2	6.9E-3	2.5E-1
Endosteal	6.0E+1	9.6E-2	3.5
Pulmonary	2.5E+1	4.7E-2	1.7
Muscle	9.7E-1	1.9E-3	6.9E-2
Liver	2.9	6.0E-3	2.2E-1
Stomach wall	1.7E-1	2.3E-4	8.5E-3
Pancreas	9.4E-1	1.8E-3	6.8E-2
LLI wall	7.8E-1	1.5E-3	5.5E-2
Kidney	7.7	1.7E-2	6.2E-1
Bladder wall	5.4E-1	1.0E-3	3.6E-2
ULI wall	3.6E-1	6.0E-4	2.2E-2
SI wall	1.6E-1	2.3E-4	8.4E-3
Ovaries	9.2E-1	1.8E-3	6.6E-2
Testes	9.4E-1	1.8E-3	6.8E-2
Spleen	1.2E+1	2.6E-2	9.2E-1
Uterus	9.2E-1	1.8E-3	6.6E-2
Thymus	9.4E-1	1.8E-3	6.7E-2
Thyroid	9.4E-1	1.9E-3	6.8E-2
Weighted mean	9.8	1.8E-2	6.2E-1

Table 6.8 Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model inactive surface uranium mine

Organ	Maximum Individual (mrem/yr)	Average Individual (mrem/yr)	Population (person-rem/yr)
Red marrow	2.1E-1	4.8E-4	6.9E-3
Endosteal	2.9	6.8E-3	9.8E-2
Pulmonary	9.5E-1	5.0E-4	7.2E-3
Muscle	5.6E-2	1.8E-4	2.6E-3
Liver	1.4E-1	5.5E-4	7.8E-3
Stomach wall	1.5E-2	1.1E-5	1.6E-4
Pancreas	5.4E-2	1.8E-4	2.6E-3
LLI wall	4.4E-2	1.4E-4	2.0E-3
Kidney	3.5E-1	1.5E-3	2.1E-2
Bladder wall	3.3E-2	9.2E-5	1.3E-3
ULI wall	2.4E-2	4.7E-5	6.7E-4
SI wall	1.4E-2	1.3E-5	1.8E-4
Ovaries	5.2E-2	1.8E-4	2.5E-3
Testes	5.5E-2	1.8E-4	2.6E-3
Spleen	5.3E-1	2.3E-3	3.3E-2
Uterus	5.2E-2	1.8E-4	2.5E-3
Thymus	5.3E-2	1.8E-4	2.5E-3
Thyroid	5.5E-2	1.8E-4	2.6E-3
Weighted mean	3.9E-1	4.7E-4	6.8E-3

Table 6.9 Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a model inactive underground uranium mine

Organ	Maximum Individual (mrem/yr)	Average Individual (mrem/yr)	Population (person-rem/yr)
Red marrow	5.8E-2	9.3E-5	3.4E-3
Endosteal	8.0E-1	1.3E-3	4.6E-2
Pulmonary	2.7E-1	3.4E-4	1.3E-2
Muscle	1.6E-2	2.9E-5	1.0E-3
Liver	3.9E-2	7.9E-5	2.8E-3
Stomach wall	4.0E-3	5.2E-6	1.8E-4
Pancreas	1.5E-2	2.8E-5	1.0E-3
LLI wall	1.2E-2	2.2E-5	8.0E-4
Kidney	9.7E-2	2.1E-4	7.6E-3
Bladder wall	9.1E-3	1.6E-5	5.8E-4
ULI wall	6.6E-3	1.0E-5	3.7E-4
SI wall	3.7E-3	4.9E-6	1.8E-4
Ovaries	1.4E-2	2.7E-5	9.7E-4
Testes	1.5E-2	2.8E-5	1.0E-3
Spleen	1.5E-1	3.2E-4	1.2E-2
Uterus	1.4E-2	2.7E-5	9.8E-4
Thymus	1.5E-2	2.8E-5	1.0E-3
Thyroid	1.5E-2	2.8E-5	1.0E-3
Weighted mean	1.1E-1	1.5E-4	5.7E-3

Table 6.10 Annual radiation dose equivalents due to atmospheric radioactive particulate and Rn-222 emissions from a hypothetical in situ uranium solution mine

Organ	Maximum Individual (mrem/yr)	Average Individual (mrem/yr)	Population (person-rem/yr)
Red marrow	1.6E-1	2.7E-4	3.8E-3
Endosteal	2.8	5.0E-3	7.1E-2
Pulmonary	3.9E+1	2.0E-2	2.9E-1
Muscle	8.4E-3	2.2E-5	3.1E-4
Liver	1.9E-2	5.4E-5	7.7E-4
Stomach wall	1.6E-2	5.7E-5	8.1E-4
Pancreas	7.6E-3	2.1E-5	3.0E-4
LLI wall	6.1E-1	2.5E-3	3.5E-2
Kidney	3.3E-1	1.0E-3	1.5E-2
Bladder wall	4.8E-3	1.2E-5	1.6E-4
ULI wall	2.0E-1	8.1E-4	1.2E-2
SI wall	3.6E-2	1.4E-4	2.0E-3
Ovaries	7.3E-3	2.1E-5	3.0E-4
Testes	8.9E-3	2.2E-5	3.1E-4
Spleen	4.6E-2	1.8E-4	2.5E-3
Uterus	7.4E-3	2.1E-5	3.0E-4
Thymus	7.9E-3	2.1E-5	3.0E-4
Thyroid	8.4E-3	2.1E-5	3.1E-4
Weighted mean	1.2E+1	6.2E-3	8.8E-2

alent rates* due to atmospheric radioactive particulate and Rn-222 emissions from the model uranium mine sites are presented in Tables 6.4 through 6.10. The Rn-222 dose equivalent rate is only for the inhalation and air immersion pathways and excludes Rn-222 daughters. The impact from Rn-222 daughters is addressed separately with a working level calculation. The dose equivalent estimates are for the model sites described for use with the AIRDOS-EPA code in Appendix K. Assumptions about food production and consumption for the maximum individual were selected for a rural setting. The maximum individual dose equivalent rate occurred about 1600 meters downwind from the center of the model site. The term "population" refers to the population living within a radius of 80 kilometers of the source. Population dose equivalents are the sum of the exposures to all individuals in the exposed population for the annual release from the model uranium mine.

Dose equivalent rates in Tables 6.4 through 6.10 indicate that the red marrow, endosteal cells, lung, kidneys, and spleen are generally the highest exposed target organs. A dose equivalent rate is presented for the "weighted mean" target organ, but this calculated result was not used in the health effect calculations. We calculated "weighted mean" dose equivalents by using organ dose equivalent weighting factors (see Appendix L) and summing the results. The weighted mean dose equivalent rate was presented instead of the total body dose equivalent rate.

Individual lifetime fatal cancer risks and estimated additional fatal cancers to the regional population due to atmospheric radioactive emissions from the model uranium mine sites are presented in Tables 6.11 and 6.12. The individual lifetime risks in Table 6.11 are those that would result from one year of exposure (external and internal) and the working levels estimated for those individuals. Except for the in situ leach mine, the individual lifetime risks in Table 6.12 are those that would result from a lifetime of exposure (71 years average life expectancy). The individual lifetime risks in Table 6.12 for the in situ leach mine are based on an exposure time of 18 years, which is the expected life, including restoration, of this type of model uranium mine.

*The dose equivalent rates were not used to calculate risk and are only presented for perspective purposes. Risks of health impact were calculated directly from external and internal radionuclide exposure data.

Table 6.11 Individual lifetime fatal cancer risk for one year of exposure and estimated additional fatal cancers to the regional population due to annual radioactive airborne emissions from model uranium mines

Source	Maximum Exposed Individual	Average Exposed Individual	Regional Population
Average surface mine			
Particulates and Rn-222	6.7E-7	7.5E-10	1.1E-5
Radon-222 daughters	5.5E-6	1.1E-8	1.6E-4
Total	6.2E-6	1.2E-8	1.7E-4
Average large surface mine			
Particulates and Rn-222	3.7E-6	3.7E-9	5.4E-5
Radon-222 daughters	1.9E-5	4.1E-8	5.9E-4
Total	2.3E-5	4.5E-8	6.4E-4
Average underground mine			
Particulates and Rn-222	1.6E-7	2.8E-10	1.0E-5
Radon-222 daughters	1.1E-5	4.9E-8	1.7E-3
Total	1.1E-5	4.9E-8	1.7E-3
Average large underground mine			
Particulates and Rn-222	1.4E-6	2.5E-9	9.0E-5
Radon-222 daughters	1.1E-4	5.0E-7	1.8E-2
Total	1.1E-4	5.0E-7	1.8E-2
Inactive surface mine			
Particulates and Rn-222	5.5E-8	6.4E-11	9.1E-7
Radon-222 daughters	4.2E-7	8.3E-10	1.2E-5
Total	4.7E-7	8.9E-10	1.3E-5
Inactive underground mine			
Particulates and Rn-222	1.5E-8	2.0E-11	7.4E-7
Radon-222 daughters	2.7E-7	1.2E-9	4.4E-5
Total	2.8E-7	1.2E-9	4.5E-5
In situ leaching facility			
Particulates and Rn-222	1.6E-6	8.7E-10	1.2E-5
Radon-222 daughters	1.1E-5	2.1E-8	3.0E-4
Total	1.3E-5	2.2E-8	3.1E-4

Table 6.12 Individual lifetime fatal cancer risk due to lifetime exposure to radioactive airborne emissions from model uranium mines

Source	Maximum Exposed Individual	Average Exposed Individual (c)
Average surface mine ^(a)		
Particulates and Rn-222	1.4E-5	1.6E-8
Radon-222 daughters	1.2E-4	2.3E-7
Total	1.3E-4	2.5E-7
Average large surface mine ^(a)		
Particulates and Rn-222	6.6E-5	6.6E-8
Radon-222 daughters	3.5E-4	7.4E-7
Total	4.2E-4	8.1E-7
Average underground mine ^(a)		
Particulates and Rn-222	3.5E-6	5.8E-9
Radon-222 daughters	2.0E-4	9.0E-7
Total	2.0E-4	9.1E-7
Average large underground mine ^(a)		
Particulates and Rn-222	2.5E-5	4.4E-8
Radon-222 daughters	1.9E-3	8.6E-6
Total	1.9E-3	8.6E-6
Inactive surface mine ^(b)		
Particulates and Rn-222	3.9E-6	4.5E-9
Radon-222 daughters	3.0E-5	5.9E-8
Total	3.4E-5	6.3E-8
Inactive underground mine ^(b)		
Particulates and Rn-222	1.1E-6	1.4E-9
Radon-222 daughters	1.9E-5	8.5E-8
Total	2.0E-5	8.6E-8
In situ leaching facility ^(d)		
Particulates and Rn-222	1.6E-5	8.7E-9
Radon-222 daughters	2.0E-4	3.8E-7
Total	2.2E-4	3.9E-7

(a) Considers exposure for 17 years to active mining and 54 years to inactive mine effluents. -

(b) Considers exposure for 71 years to inactive mine effluents.

(c) Considers the average individual in the regional population within an 80-km radius of the model mine.

(d) Considers 10-year operation and 8-year restoration.

Table 6.13 Genetic effect risk to descendants for one year of parental exposure to atmospheric radioactive airborne emissions from model uranium mines

Source	Descendants of Maximum Exposed Individual (effects/ birth)	Descendants of Average Exposed Individual (effects/ birth)	Descendants of Regional Population (effects/yr)
Average surface mine	6.3E-7	2.6E-9	1.6E-5
Average large surface mine	3.7E-6	1.3E-8	7.9E-5
Average underground mine	1.4E-7	2.9E-10	4.4E-6
Average large underground mine	1.1E-6	2.4E-9	3.6E-5
Inactive surface mine	6.0E-8	2.4E-10	1.4E-6
Inactive underground mine	1.6E-8	3.4E-11	5.0E-7
In situ leach facility	8.0E-9	2.7E-11	1.6E-7

Table 6.14 Genetic effect risk to descendants for a 30-year parental exposure to atmospheric radioactive airborne emissions from model uranium mines

Source	Effects/birth	
	Descendants of Maximum Exposed Individual	Descendants of Average Exposed Individual ^(c)
Average surface mine ^(a)	1.2E-5	4.6E-8
Average large surface mine ^(a)	6.4E-5	2.2E-7
Average underground mine ^(a)	2.6E-6	5.4E-9
Average large underground mine ^(a)	2.0E-5	4.0E-8
Inactive surface mine ^(b)	1.8E-6	7.2E-9
Inactive underground mine ^(b)	5.0E-7	5.8E-10
In situ leach facility ^(d)	1.4E-7	4.8E-10

(a) Considers exposure to 17 years active mining and 13 years inactive mine effluents.

(b) Considers exposure for 30 years to inactive mine effluents.

(c) Considers the average individual in the regional population within an 80-km radius of the model mine.

(d) Considers 10-year operation and 8-year restoration.

Genetic effect risks due to atmospheric radioactive emissions from the model uranium mine sites are presented in Tables 6.13 and 6.14. The risks to descendants in Table 6.13 are those that would result from one year of exposure to the parent or parents of first generation individuals. The descendant risks in Table 6.14 are those that would result from 30 years exposure to the first generation parent or parents, except for the in situ leach mine where we used an 18-year exposure time. The 30-year time period represents the mean years of life where gonadal doses are genetically significant.

We estimated the health impact risks with the DARTAB code using exposure data from the AIRDOS-EPA code. The dose equivalent and risk conversion factors that we used with the DARTAB code are tabulated in Appendix L. The somatic risk conversion factors are based on a lifetime (71 years average lifetime) exposure time, and the genetic effect risk conversion factors are based on a 30-year exposure time. When the exposure time for calculated risks was only one year, we calculated the risk by multiplying the risk calculated by DARTAB with the ratio of the one year exposure time to the exposure times used to calculate the risk conversion factors ($1/71$ for somatic effects and $1/30$ for genetic effects to descendants of maximum and average exposed individuals).^{*} Appendix L contains a discussion of the health risk assessment methodology.

We developed several tables to present the calculated health impact risk. The percentage contributions to the fatal cancer risks for individual sources at each model uranium mine site are contained in Table 6.15 for the maximum individual and Table 6.16 for the average individual. The fatal cancer risks by source term for one year of exposure which we used to calculate percentage contributions are contained in Tables L.4 to L.6 in Appendix L. Tables L.7 to L.9 contain genetic risks by source term at each model uranium mine site. The percent of the fatal cancer risk due to radon-222 daughter concentrations at model uranium mine sites is indicated in Table 6.17. The percent of the fatal cancer risk for principal nuclides and pathways due to radioactive particulate and Rn-222 emissions at each model uranium mine site are contained in Table 6.18.

^{*}A correction factor was not needed for DARTAB calculated genetic effects committed per year to the regional population.

Table 6.15 Percent of the fatal cancer risk for the maximum individual
due to the sources of radioactive emissions at model uranium
mines

Mine type	Percent of fatal cancer risk (a,b)				Vehicular Dust
	Mining Activities	Ore	Sub-ore	Spoils	
Average surface mine	56 (95)	18 (66)	14 (98)	12 (89)	<1 (0)
Average large surface mine	59 (93)	14 (41)	12 (98)	14 (86)	1 (0)
Average underground mine	80 (\pm 100)	3 (79)	17 (97)	<1 (96)	<1 (0)
Average large underground mine	89 (\pm 100)	2 (76)	9 (96)	<1 (96)	<1 (0)
Inactive surface mine	28 ^(c) (100)	0	0	72 (84)	0
Inactive underground mine	77 ^(c) (100)	0	0	23 (77)	0
In situ leach facility	100 (87)	0	0	0	0

(a) See Table L. 4, Appendix L.

(b) Values in parentheses are percent contribution of radon-222 daughters.

(c) Emissions from abandoned pit (surface mine) or vents and portals (underground mine).

Table 6.16 Percent of the fatal cancer risk for the average individual
in the regional population due to the sources of radioactive
emissions at model uranium mines

Mine type	Percent of fatal cancer risk ^(a,b)				
	Mining Activities	Ore	Sub-ore	Spoils	Vehicular Dust
Average surface mine	58 (97)	16 (78)	14 (99)	12 (93)	<1 (0)
Average large surface mine	60 (96)	11 (64)	12 (99)	16 (92)	1 (0)
Average underground mine	81 (\cong 100)	2 (93)	16 (99)	<1 (99)	<1 (0)
Average large underground mine	89 (\cong 100)	2 (91)	9 (99)	<1 (99)	<1 (0)
Inactive surface mine	29 ^(c) (100)	0	0	71 (90)	0
Inactive underground mine	80 ^(c) (100)	0	0	20 (92)	0
In situ leach facility	100 (96)	0	0	0	0

(a) See Table L.5, Appendix L.

(b) Values in parentheses are percent contribution of radon-222 daughters.

(c) Emissions from abandoned pit (surface mine) or vents and portals (underground mines).

Table 6.17 Percent of fatal cancer risks due to radon-222
daughter concentrations at model uranium mine
sites

Source	Percent fatal cancer risk ^(a)
Average surface mine	89
Average large surface mine	84
Average underground mine	99
Average large underground mine	99
Inactive surface mine	88
Inactive underground mine	95
In situ leach facility	87

(a) Remainder due to radioactive particulate and Rn-222 emissions.

Table 6.18 Percent of the fatal cancer risk for principal nuclides and pathways due to radioactive particulate and Rn-222 emissions at model uranium mines

Mine Type	Receptor	Principal Nuclides	Percent of fatal cancer risk			
			Internal Pathways		External Pathways	
			Ingestion	Inhalation	Air Immersion	Ground Surface
Average Surface Mine	Max. Individual	U-238(20.0), U-234(22.1), Th-230(31.7), Ra-226(7.94), Po-210(7.33)	15.8	80.2	0.003	4.02
	Av. Individual or population	U-238(9.17), U-234(10.1), Th-230(22.7), Ra-226(21.3), Pb-210(6.92), Po-210(22.4)	60.1	38.1	0.005	1.81
Average Large Surface Mine	Max. Individual	U-238(20.0), U-234(22.2), Th-230(31.8), Ra-226(7.98)	15.9	80.0	0.002	4.05
	Av. Individual or Population	U-238(9.19), U-234(10.1), Th-230(22.7), Ra-226(21.4), Pb-210(6.94), Po-210(22.4)	60.5	37.7	0.004	1.83
Average Underground Mine	Max. Individual	U-238(17.9), U-234(19.8), Th-230(28.4), Ra-226(7.14), Po-210(6.59), Rn-222(13.6)	14.0	82.5	0.025	3.52
	Av. Individual or Population	U-238(12.0), U-234(13.2), Th-230(20.1), Ra-226(7.24), Po-210(7.31), Rn-222(34.6)	16.9	80.6	0.063	2.43
Average Large Underground Mine	Max. Individual	U-238(17.5), U-234(19.3), Th-230(27.7), Ra-226(6.97), Po-210(6.43), Rn-222(16.0)	13.6	82.9	0.029	3.39
	Av. Individual or Population	U-238(11.2), U-234(12.4), Th-230(18.8), Ra-226(6.76), Po-210(6.83), Rn-222(39.2)	15.7	82.0	0.071	2.25
Inactive Surface Mine ^(a)	Max. Individual	U-238(19.5), U-234(21.6), Th-230(31.0), Ra-226(9.4), Bi-214(5.31), Po-210(7.17)	16.8	75.4	0.002	7.85
	Av. Individual or Population	U-238(8.76), U-234(9.68), Th-230(21.7), Ra-226(26.1), Pb-210(6.77), Po-210(21.4)	62.2	34.3	0.003	3.48
Inactive Underground Mine ^(a)	Max. Individual	U-238(19.6), U-234(21.6), Th-230(31.1), Ra-226(9.45), Bi-214(5.33), Po-210(7.21)	16.9	75.2	0.001	7.88
	Av. Individual or Population	U-238(17.0), U-234(18.8), Th-230(28.5), Ra-226(12.7), Bi-214(4.81), Po-210(10.4)	26.3	66.6	0.004	7.09
In situ Leaching Facility	Max. Individual	U-238(45.2), U-234(50.0), U-235(2.21)	0.46	99.5	0.002	0.039
	Av. Individual or Population	U-238(43.3), U-234(47.8), U-235(2.12)	3.50	96.5	0.009	0.038

(a) Spoils source term only.

The fatal cancer health risk at each of the model uranium mine sites is dominated by the lung cancer risk from radon-222 daughter exposures (see Table 6.17). Radioactive particulates and Rn-222 contributed to a little over 10 percent of the total fatal cancer health risk at the model surface mines and at the in situ leaching facility (see Table 6.11). Essentially all the risks from the model underground mines are due to radon-222 daughter exposures. The fatal cancer health risks from the active model underground mines are greater than the risks from the active model surface mines because of the larger quantity of Rn-222 released. The risks are similar at inactive surface and underground mines.

The largest fatal cancer risk is from the average large underground mine (see Tables 6.11 and 6.12)--an estimated $1.9\text{E-}3$ lifetime fatal cancer risk to the maximum exposed individual for a lifetime exposure. The lifetime fatal cancer risk to the average individual in the regional population is estimated to be $8.6\text{E-}6$ for a lifetime exposure period. The number of estimated additional fatal cancers in the regional population per year of mine operation is estimated to be $1.8\text{E-}2$.

For the active surface mines, about 60 percent of the radon daughter impact is from the exposed pit surfaces (see Table L.4). For the active underground mines, the predominate radon daughter impact is from mine vent air. For the inactive surface mine, about 70 percent of the radon daughter impact is from waste rock pile exhalation and about 30 percent was from the pit interior surfaces. About 80 percent of the radon daughter impact for the inactive underground mine was due to radon releases from the mine vents and entrance. The release of radon from the pregnant leach surge tanks was the predominate source of the radon daughter health impact risk for the model in situ leach mine. Detailed percentages of the lifetime fatal cancer risks by source term for each model uranium mine are contained in Tables 6.15 and 6.16.

The health impact from particulate radionuclides and Rn-222 was predominately due to U-238 and daughter radionuclides (see Table 6.18). Thorium-232 and daughters were only minor contributors to the particulate and Rn-222 fatal cancer risk with Rn-222 only contributing significantly (14 to 40 percent) at active underground mines. The majority of the exposure to individuals around the model uranium mines is received from the internal pathways. Inhalation was the most important internal pathway except for the average individual and regional population impact at surface mines

where ingestion was the major pathway (see Table 6.18). For active surface mines, about 52 percent of the particulate and Rn-222 impact to the maximum individual was from the ore source term, and about 25 percent of the health impact was from the mining activities source term (see Table L.4). For active underground mines, between 28 and 46 percent of the particulate and Rn-222 impact was from the ore source term and between 26 and 41 percent of the particulate and Rn-222 impact was from the sub-ore source term. The predominant source of the particulate and Rn-222 impact from the inactive mines was particulate radionuclides in wind-suspended dust from the waste rock pile. The release of particulate radionuclides from the uranium recovery plant was the predominant source of the particulate health impact risk for the model in situ leach mine.

For perspective, the calculated fatal cancer risks can be compared to the estimated cancer risk from all causes. The American Cancer Society estimates the risk of cancer death from all causes to be 0.15 (Ba79). The maximum exposed individual around the model average large underground mine is estimated to incur an additional lifetime fatal cancer risk of 0.0019 (1.3 percent) due to radioactive airborne emissions from the model mine. There is a regional population of 36,004 persons for the model average large underground mine site located in New Mexico. The cancer death rate for the State of New Mexico for whites of both sexes was 154.5 deaths per year for 1973 to 1976 per 100,000 people (NCI78). Applying this statistic to the regional population, about 56 cancer deaths are estimated to occur each year in the regional population from all causes. Applying the approximate fatal cancer risk coefficient of 0.15 to the regional population of 36,004 persons, about 5,400 people in the regional area would normally die of cancer. About 0.018 additional cancer deaths (0.00033 percent) in the regional population are estimated per year of operation from radioactive airborne emissions at the model average large underground mine.

The risk of genetic effects from radiation exposure at model uranium mine sites is very small compared to the normal occurrence of hereditary disease. The national incidence of genetic effects is 60,000 per 10^6 births (NAS72). The normal occurrence of hereditary disease for the descendants of the regional population of 14,297 at the model average large surface mine in Wyoming is 0.06 effects per birth and 12.1 effects per year, based on 202 live births per year in the regional population. (We present statistics for the site of the average large surface mine since the largest

genetic risk for all the evaluated model uranium mines occurred at this site [see Tables 6.13 and 6.14]). We estimated the genetic effect risk to the descendants of the maximum exposed individual to be an additional $6.4\text{E-}5$ effects/birth (0.1 percent increase) for a 30-year exposure period. The genetic effect risk to the descendants of the average exposed individual in the regional population is estimated to be an additional $2.2\text{E-}7$ effects/birth (0.00036 percent increase) for a 30-year exposure period. The number of additional genetic effects committed to the descendants of the regional population per year of operation of the average large surface mine is estimated to be $7.9\text{E-}5$. The additional committed genetic effects constitute a very small increase to the 12.1 effects that will normally occur each year in the live births within the regional population.

6.1.2 Nonradioactive Airborne Emissions

To calculate atmospheric concentrations at the location of the maximum individual, we used the data on nonradioactive air pollutant emissions from Section 3. We compared these pollutant air concentrations with calculated nonoccupational threshold limit values, natural background concentrations, and average urban concentrations of selected airborne pollutants in the United States.

The "natural" background atmospheric concentration has been defined (Va71) as the concentration of pollutants in areas absent of activities by man which cause significant pollution. Variations in background levels may result from differences in mineral content of the soil, vegetation, wind conditions, and the proximity to the ocean or metropolitan areas. Based on an extensive literature survey and consideration of the abundance and distribution of the chemical elements in the ocean and earth's crust, a set of "natural" background airborne concentrations has been developed for the United States (Va71). Natural background airborne concentrations for selected pollutants are listed in the second column of Table 6.19. Also listed in the table are average concentrations of airborne pollutants in urban areas. The latter are arithmetic mean concentrations obtained from measurements taken over a period of several years (Va71).

6.1.2.1 Combustion Products

Airborne concentrations of combustion products released from diesel and gasoline-powered equipment were estimated for the site of the maximum

Table 6.19 Natural background concentrations and average urban concentrations of selected airborne pollutants in the United States

Pollutant	Natural Background Concentration, $\mu\text{g}/\text{m}^3$	Average Urban Concentration, $\mu\text{g}/\text{m}^3$
<u>Gases</u>		
CO	100	7000
NO ^x	40	141
NH ₃	10	80
SO ₂	5	62
CO ₂	594,000	NR
Hydrocarbons	NR ^(a)	500
<u>Suspended particles</u>		
Total	20 - 40	105
As	0.005	0.02 (1)
Ba	0.005	NR
Cd	0.0001	0.002
Co	0.0001	0.0005
Cr	0.001	0.015
Cu	0.01	0.09
Hg	0.0005	0.1
Fe	0.2 - 0.5	1.58
Pb	0.001	0.79
Mg	0.1	NR
Mn	0.01	0.1
Mo	0.0005	0.005
Ni	0.001	0.034
Se	0.001	NR
Sr	0.005	NR
Th	0.0005	NR
U	0.0001	NR
V	0.001	0.05
Zn	0.01	0.67
Zr	0.001	NR

(a) NR - Not Reported.

Source: Va71; except for CO₂, Ba76.

individual. The concentrations were computed using the annual release rates given in Tables 3.30 and 3.52 with dispersion parameters applicable for the model underground (New Mexico) and surface (Wyoming) mining areas (Appendix K). The estimated combustion product concentrations are low compared to the natural background and average urban concentrations (see Table 6.20). A conservative threshold limit value (TLV) was computed, as described in Section 6.1.2.3 for SO_2 , CO, and NO_2 . Of these pollutants, only the nitrogen oxide concentrations at the average large surface mine exceed the nonoccupational TLV. Considering these comparisons and the conservative nature of the analyses, combustion products released from heavy uranium mining equipment do not appear to pose a health hazard.

6.1.2.2 Nonradioactive Gases

Airborne concentrations of the three principal nonradioactive gases released from the hypothetical in situ leach mining site were computed using the source terms from Table 3.59 and the meteorological parameters and dispersion model described in Appendix K. Table 6.21 shows the estimated atmospheric concentrations at the location of a maximum individual; occupational threshold limit values (TLV's); adjusted TLV's applicable to nonoccupational exposures; and the percent the estimated concentrations are of the adjusted TLV's. The occupational TLV's have been conservatively adjusted. They were adjusted on the basis of a 168-hr week, instead of a 40-hour week and a safety factor of 100.

The results of this analysis indicate that two of the estimated concentrations fall below their respective TLV's, and the concentration of ammonium chloride is approximately equal to its TLV. Considering the conservative nature of the adjusted nonoccupational TLV on which the comparisons were made, none of the nonradioactive gases appear to be at concentrations that might pose a serious health hazard. The ammonia level is about 80 percent of the estimated "natural" background concentration and only about 10 percent of the average urban concentration (Table 6.19).

6.1.2.3 Trace Metals and Particulates in the Form of Dust

We identified seventeen trace metals and particulates in the form of dust as potential airborne emissions from uranium mines. Table 6.22 presents projected airborne concentrations of the metals and particulates at the site of the maximum individual for six mine classifications. As might

Table 6.20 Combustion product concentrations at the site of the maximum individual with comparisons, $\mu\text{g}/\text{m}^3$

Pollutant	Average underground mine	Average large underground mine	Average surface mine	Average large surface mine	Natural background concentration ^(a)	Average urban concentration ^(a)	Non-occupational TLV ^(b)
Particulates of combustion	1.4E-3	1.6E-2	9.7E-2	4.5E-1	NR ^(c)	NR	NR
SO _x	1.2E-2	1.3E-1	5.5E-1	2.2E+0	5E+0	6.2E+1	3.1E+1
CO	9.7E-2	1.1E+0	4.3E+0	1.8E+1	1.0E+2	7.0E+3	1.3E+2
NO _x	1.6E-1	1.8E+0	7.1E+0	3.0E+1	4.0E+1	1.4E+2	2.1E+1
Hydrocarbons	1.6E-2	1.8E-1	7.1E-1	3.1E+0	NR	5.0E+2	NR

^(a) See Table 6.19.

^(b) Nonoccupational TLV = TLV (mg/m³) x 40 hr/168 hr x 10⁻² x 10³ $\mu\text{g}/\text{mg}$ (ACGIH76).

^(c) NR - Not reported.

Table 6.21 A comparison of the airborne concentrations of nonradioactive gases at the hypothetical in situ leach site with threshold limit values

Contaminant	Atmospheric Concentration ^(a) ($\mu\text{g}/\text{m}^3$)	TLV ^(b) (mg/m^3)	Non- occupational ^(c) TLV ($\mu\text{g}/\text{m}^3$)	Percent of Nonoccupational TLV
NH ₃	8.1	18	43	19
NH ₄ Cl	24	10	24	100
CO ₂	60	9000	21,400	0.3

(a) Location of maximum individual.

(b) Source: ACGIH76.

(c) Nonoccupational TLV = TLV (mg/m^3) \times 40 hr/168 hr \times 10^{-2} \times 10^3 $\mu\text{g}/\text{mg}$.

be expected, large surface mine emissions usually have the greatest concentrations, and those from inactive underground mines the least. Projected metal concentrations range from a low of about $5 \times 10^{-7} \mu\text{gm}/\text{m}^3$ of cobalt from inactive underground mines to a high of about $1 \mu\text{gm}/\text{m}^3$ of potassium from large surface mines.

Table 6.23 shows where particulates (dust) or trace metal air concentrations are estimated to exceed natural background or average urban air concentrations (Table 6.19). Several trace metal air concentrations exceed "natural" background; however, only the estimated air concentration of particulates (dust) exceeds the air concentration of airborne pollutants in urban areas.

We evaluated the significance of these concentrations by comparing them with threshold limit values (TLV's) for workroom environments published by the American Conference of Governmental Industrial Hygienists (ACGIH76). These TLV's, which are for occupational workers and a 40-hour workweek, were adjusted by multiplying by 40/168 to convert them to continuous exposure values and dividing by 100 to make them applicable to the general public. Table 6.24 is a tabulation of the adjusted TLV's, the projected concentrations of metals and particulates (from Table 6.22), and the ratio of these concentrations to the adjusted TLV's. The sums of these ratios provide a measure of whether a mixture of the metals would be a significant problem, a sum greater than one indicating that the "composite" TLV has been exceeded.

Table 6.24 shows that in no case does a single metal exceed its TLV, nor do any of the mixtures exceed a "composite" TLV. Although TLV's were not available for potassium and strontium, their low toxicity and low concentrations make it unlikely that their addition to the sums would change this conclusion. For the worst case, large surface mines, the sum of ratios is only about 17 percent of the limit.

Particulates, on the other hand, present a different picture. The TLV for nonspecific particulates, nuisance dust, was chosen for comparison. It can be seen that the TLV is exceeded by a factor of six at the large model surface mine and nearly exceeded at the average model surface mine. About 50% of the exposure to dust is from vehicular traffic, and about 30% results from mining activities within the pit.

In summary, specific trace metal airborne emissions from uranium mines do not appear to present a significant hazard, either singly or as com-

Table 6.22 Stable trace metal airborne concentrations at the site of the maximum individual, $\mu\text{g}/\text{m}^3$

Trace metal	Avg. under-ground mine	Avg. large underground mine	Avg. surface mine	Avg. large surface mine	Inactive under-ground mine	Inactive surface mine
As	3.1E-5	1.9E-4	2.6E-4	1.5E-3	3.1E-6	1.5E-5
Ba	5.1E-4	1.8E-3	7.0E-3	4.2E-2	3.6E-5	1.6E-4
Co	4.0E-6	3.1E-5	1.1E-5	4.7E-5	4.5E-7	2.9E-6
Cu	3.3E-5	1.5E-4	4.4E-4	2.6E-3	2.2E-6	1.1E-5
Cr	5.0E-5	1.4E-4	1.1E-3	6.9E-3	8.9E-7	3.6E-6
Fe	9.7E-3	4.1E-2	1.4E-1	8.5E-1	6.3E-4	2.7E-3
Hg	7.2E-6	1.5E-5	1.8E-4	1.1E-3	NA ^(a)	NA
K	1.3E-2	6.3E-2	1.7E-1	1.0	9.8E-4	4.4E-3
Mg	9.4E-4	6.9E-3	2.5E-3	1.0E-2	1.3E-4	6.2E-4
Mn	7.1E-4	2.8E-3	1.1E-2	6.8E-2	3.7E-5	1.7E-4
Mo	3.3E-5	2.3E-4	1.4E-4	6.7E-4	4.5E-6	2.0E-5
Ni	4.9E-6	3.9E-5	1.4E-5	5.8E-5	8.9E-7	3.6E-6
Pb	4.1E-5	2.0E-4	5.4E-4	3.2E-3	3.1E-6	1.4E-5
Se	3.1E-5	2.2E-4	1.2E-4	5.9E-4	4.5E-6	1.9E-5
Sr	1.7E-4	5.5E-4	3.4E-3	2.1E-2	4.9E-6	2.3E-5
V	4.7E-4	3.0E-3	2.2E-3	1.8E-2	5.4E-5	2.5E-4
Zn	2.6E-5	9.6E-5	4.6E-4	2.8E-3	1.3E-6	5.2E-6
Part ^(b)	1.2	3.9	2.3E+1	1.4E+2	3.9E-2	1.7E-1

^(a)NA - Not available.

^(b)Part. - Particulates (dust).

Table 6.23 Comparison of stable trace metal airborne concentrations at the location of the maximum individual with natural background concentrations and average urban concentrations of these airborne pollutants

<u>Exceed Natural Background</u> ^(a)	<u>Exceed Average Urban Concentration</u> ^(a)
<u>Average Large Surface Mine</u>	
Ba, Cr (possible), Fe, Hg (possible), Mn, Mo, Pb, Sr, V, particulates	Particulates
<u>Average Surface Mine</u>	
Ba, Cr (possible), Mn, V	None
<u>Average Large Underground Mine</u>	
V	None
<u>Average Underground Mine</u>	
None	None
^(a) See Tables 6.19 and 6.22.	

Table 6 24 Comparison of trace metal airborne concentrations at the site of the maximum individual with threshold limit values (TLV's) in the workroom environment adjusted for continuous exposure to the general public, $\mu\text{g}/\text{m}^3$

Trace metal	Adjusted ^(a) TLV	Average Underground Mine		Average Large Underground mine		Average Surface Mine		Average Large Surface mine		Inactive Underground mine		Inactive Surface mine	
		Conc	Conc /TLV	Conc.	Conc /TLV	Conc.	Conc./TLV	Conc	Conc /TLV	Conc	Conc./TLV	Conc.	Conc./TLV
As	1.2	3.1E-5	3E-5	1.9E-4	2E-4	2.6E-4	2E-4	1.5E-3	1E-3	3.1E-6	3E-6	1.5E-5	1E-5
Ba	1.2	5.1E-4	4E-4	1.8E-3	2E-3	7.0E-3	6E-3	4.2E-2	4E-2	3.6E-5	3E-5	1.6E-4	1E-4
Co	0.24	4.0E-6	2E-5	3.1E-5	1E-4	1.1E-5	5E-5	4.7E-5	2E-4	4.5E-7	2E-6	2.9E-6	1E-5
Cu	0.48	3.3E-5	7E-5	1.5E-4	3E-4	4.4E-4	9E-4	2.6E-3	5E-3	2.2E-6	5E-6	1.1E-5	2E-5
Cr	1.2	5E-5	4E-5	1.4E-4	1E-4	1.1E-3	9E-4	6.9E-3	6E-3	8.9E-7	7E-7	3.6E-6	3E-6
Fe	12	9.7E-3	8E-4	4.1E-2	3E-3	1.4E-1	1E-2	8.5E-1	7E-2	6.3E-4	5E-5	2.7E-3	2E-4
Hg	0.12	7.2E-6	6E-5	1.5E-5	1E-4	1.8E-4	2E-3	1.1E-3	9E-3	NA	----	NA	--
Mn	NA ^(b)	1.3E-2	---	6.3E-2	---	1.7E-1	--	1.0E+0	--	9.8E-4	----	4.4E-3	--
Mg	24	9.4E-4	4E-5	6.9E-3	3E-4	2.5E-3	1E-4	1.0E-2	4E-4	1.3E-4	5E-6	6.2E-4	3E-5
Mn	12	7.1E-4	6E-5	2.8E-3	2E-4	1.1E-2	9E-4	6.8E-2	6E-3	3.7E-5	3E-6	1.7E-4	1E-5
Mo	12	3.3E-5	3E-6	2.3E-4	2E-5	1.4E-4	1E-5	6.7E-4	6E-5	4.5E-6	4E-7	2.0E-5	2E-6
Ni	0.24	4.9E-6	2E-5	3.9E-5	2E-4	1.4E-5	6E-5	5.8E-5	2E-4	8.9E-7	4E-6	3.6E-6	2E-5
Pb	0.36	4.1E-5	1E-4	2.0E-4	6E-4	5.4E-4	2E-3	3.2E-3	9E-3	3.1E-6	9E-6	1.4E-5	4E-5
Se	0.48	3.1E-5	6E-5	2.2E-4	5E-4	1.2E-4	2E-4	5.9E-4	1E-3	4.5E-6	9E-6	1.9E-5	4E-5
Sr	NA	1.7E-4	--	5.5E-4	--	3.4E-3	--	2.1E-2	--	4.9E-6	--	2.3E-5	--
V	1.2	4.7E-4	4E-4	3.0E-3	2E-3	2.2E-3	2E-3	1.8E-2	2E-2	5.4E-5	4E-5	2.5E-4	2E-4
Zn	12	2.6E-5	2E-6	9.6E-5	8E-6	4.6E-4	4E-5	2.8E-3	2E-4	1.3E-6	1E-7	5.2E-6	4E-7
Total of ratios		2E-3		1E-2		3E-2		1.7E-1		2E-4		7E-4	
Particulates:													
Dust 24 ^(c)		1.2E+0	5E-2	3.9E+0	2E-1	2.3E+1	1E+0	1.4E+2	6E+0	3.9E-2	2E-3	1.7E-1	7E-3

(a) Adjusted TLV = Occupational TLV (mg/m^3) \times 40 hr/168hr \times 10^3 $\mu\text{g}/\text{mg} \times 1/100$.

(b) NA - Not available.

(c) Limit for nuisance dust - total mass.

Source: Workroom TLV's from ACGIH76.

posite mixtures, when evaluated against adjusted threshold limit values. However, particulate emissions, at least for surface mines, require further evaluation. If model predictions can be verified by measurement, control measures are indicated.

6.1.3 Radioactive Aquatic Emissions

We used the data on radioactive releases from mine dewatering (Sections 3.3.3 and 3.4.3) to estimate the public health impact of mining operations at a typical active underground mining site (New Mexico) and a typical active surface mining site (Wyoming). The health risks estimated in this section are of fatal cancers and genetic effects to succeeding generations. Dose equivalents and health risks per year of active mine operation are estimated for the maximum and average individuals and for the population of each assessment area. These calculated dose equivalents and health risk estimates are believed to be higher than the actual dose equivalents and health risks because of the conservative assumptions required to predict movement of radionuclides in surface waters (see Section J.2 of Appendix J). Very few data are available on aquatic releases from inactive mines; hence, the significance of these releases, particularly for Colorado and Utah where inactive mines are numerous, could not be determined.

The individual and population dose equivalents presented in this section are computed using the models and parameters discussed in Appendix J. The health risk estimates are generated by the following procedures:

- a. For inhalation or ingestion of radionuclides, the quantity of radionuclides taken into the body is determined as part of the dose equivalent calculations. This quantity is multiplied by a health risk per unit intake conversion factor.
- b. For external irradiation from ground deposited radionuclides or from air submersion, the dose equivalents are calculated and multiplied by a health risk per unit dose equivalent conversion factor.

The health risk per unit intake and health risk per unit external dose equivalent conversion factors for aquatic releases are listed in Tables J.13 and J.14, Appendix J. This appendix also discusses the health risk assessment methodology used to obtain the risks presented in this section. Uranium and Ra-226 releases are given for both active mining sites. It is assumed that the stated uranium releases are entirely U-238 and that U-234 is in equilibrium with the U-238 but that Th-230 precipitates out of the

Table 6.25 Annual radiation dose equivalent rates due to aquatic releases from the New Mexico model underground mine

Organ	Maximum Individual Dose Rate (mrem/y)	Average Individual Dose Rate (mrem/y)	Population Dose Rate (person-rem/y)
Endosteal	5.6E+1	5.0	3.2E+2
Red Marrow	2.0	1.6E-1	1.1E+1
Lung	1.3	2.1E-3	1.4E-1
Liver	5.5E-1	2.9E-2	1.9
Stomach Wall	1.9E-1	3.8E-3	2.5E-1
LLI Wall ^(a)	9.4E-1	6.6E-2	4.3
Thyroid	4.5E-1	2.5E-2	1.6
Kidney	2.8E+1	2.4	1.6E+2
Muscle	4.9E-1	2.5E-2	1.6
Ovaries	4.1E-1	2.4E-2	7.8E-1
Testes	4.7E-1	2.4E-2	7.8E-1
Weighted Mean	2.2	1.5E-1	9.9

^(a) Lower large intestine wall.

Table 6.26 Annual radiation dose equivalent rates due to aquatic releases from the Wyoming model surface mine

Organ	Maximum Individual Dose Rate (mrem/y)	Average Individual Dose Rate (mrem/y)	Population Dose Rate (person-rem/y)
Endosteal	6.8E-1	2.1E-1	3.4
Red Marrow	3.8E-2	7.4E-3	1.2E-1
Lung	2.3E-2	1.0E-4	1.7E-3
Liver	3.0E-2	2.8E-3	4.5E-2
Stomach Wall	1.0E-2	2.8E-4	4.6E-3
LLI Wall ^(a)	2.9E-2	7.7E-3	1.3E-1
Thyroid	1.8E-2	1.4E-3	2.3E-2
Kidney	4.0E-1	1.1E-1	1.8
Muscle	1.9E-2	1.5E-3	2.4E-2
Ovaries	1.5E-2	1.5E-3	1.2E-2
Testes	1.8E-2	1.4E-3	1.2E-2
Weighted Mean	4.0E-2	7.1E-3	1.2E-1

^(a) Lower large intestine wall.

Table 6.27 Individual lifetime fatal cancer risk and committed fatal cancers to the population residing within the assessment areas

Source	Maximum exposed individual lifetime fatal cancer risk for operation of the mine		Average exposed individual lifetime fatal cancer risk for operation of the mine ^(a)		Committed fatal cancers for the assessment area population for operation of the mine	
	<u>1 yr.</u>	<u>17 yrs.</u>	<u>1 yr.</u>	<u>17 yrs.</u>	<u>1 yr.</u>	<u>17 yrs.</u>
Underground mine site (New Mexico)	3.3E-7	5.6E-6	2.0E-8	3.4E-7	1.3E-3	2.2E-2
Surface mine Site (Wyoming)	7.1E-9	1.2E-7	9.6E-10	1.6E-8	1.6E-5	2.7E-4

(a) The average individual risk is the cumulative population risk divided by the population residing within the assessment area.

Also, it is assumed that Rn-222, Pb-214, Bi-214, Pb-210, and Po-210 are in equilibrium with the Ra-226. For example, a reported release rate of 0.01 Ci/yr of U-238 would be reflected in the analyses as 0.01 Ci/yr of U-238 and 0.01 Ci/yr of U-234. In like manner, a release of 0.001 Ci/yr of Ra-226 would be reflected in the analyses as 0.001 Ci/yr Ra-226, 0.001 Ci/yr Rn-222, 0.001 Ci/yr Pb-214, 0.001 Ci/yr Bi-214, 0.001 Ci/yr Pb-210, and 0.001 Ci/yr Po-210.

The maximum individual, average individual, and population annual dose equivalent rates due to release of mine water containing radionuclides are given in Tables 6.25 and 6.26 for the two active uranium mine sites. The population dose equivalent rates are the sum of the dose equivalent rates to all individuals residing within the assessment areas due to the annual release from the model uranium mine. Average individual dose equivalent rates are computed by dividing the population dose equivalent rates by the number of persons in the assessment area.

The dose equivalent rates in Tables 6.25 and 6.26 indicate that the endosteal cells and kidney are the highest exposed target organs. Ingestion is the predominant exposure mode for both the endosteal cells and the kidney.

Individual lifetime fatal cancer risks and committed fatal cancers to the population within the assessment area for radionuclide releases due to mine dewatering are presented in Table 6.27. The maximum and average individual lifetime risks (columns 2 and 3, respectively) and the committed fatal cancers to the population within the assessment area (column 4) are shown for both one year of release of radionuclides due to mine dewatering and, in parenthesis, for the cumulative release over the 17 years of mine operation. To compute the 17-year risks, the one-year risks are multiplied by 17, which assumes equal annual radionuclide discharges. At both the model underground (New Mexico) and surface (Wyoming) mines, the majority of the risk is from releases of U-238, U-234, and Po-210.

A perspective on the additional fatal cancers estimated for the population (Table 6.27) can be gained by realizing that the probability of an individual dying of cancer of all types is 0.15 (Ba79). Taking the New Mexico assessment area (64,950 persons) as an example, the expected number of deaths from all forms of cancer for this population is 9,743 persons. For the 17 years of mine operations, the estimated increase in the number of deaths from cancer in the assessment area population is 0.022 deaths

(Table 6.27). This represents a 0.00023 percent increase in the expected fatal cancer occurrences in the assessment area population as a result of operation of the underground mine in New Mexico over its 17-year active life. For the Wyoming assessment area (16,230 persons), the estimated increase in the expected fatal cancer deaths due to operation of the surface mine for 17 years is 0.000011 percent.

Table 6.28 presents the genetic risks to succeeding generations, for exposure to both individuals and the population within the assessment area, caused by mine dewatering radionuclide releases. The genetic risks to succeeding generations of maximum and average exposed individuals (columns 2 and 3, respectively) and the committed genetic effects to the descendants of the present population within the assessment area (column 4) are shown for one year of releases. The mechanics and assumptions used to estimate the genetic effects are similar to those used to estimate fatal cancer risks (see Appendix J). For both the model underground (New Mexico) and surface (Wyoming) mines the majority of the risk is from releases of U-238, U-234, and Po-210.

The risks of additional genetic effects due to the discharge of contaminated mine water from model uranium mine sites are very small when compared to the normal occurrence of hereditary diseases. As given in Section 6.1.1, the natural incidence of genetic effects is 60,000 per million births (NAS72), or 0.06 effects per birth. This natural incidence rate is equivalent to 848 effects per year per million persons, considering a birth rate of 0.01413 births per person-year. Taking the New Mexico site as an example, the normal incidence of genetic effects for the assessment area population (64,950 persons) during the 17 years of operation of the mine would be 936 genetic effects. The increase in genetic effects committed to the assessment area population during the 17 years of operation is 0.015 genetic effects committed. Thus, the genetic effects committed due to aquatic wastes released during the operation of the New Mexico underground mine are only 0.0016% of the genetic effects which occur due to other causes during the mine operating life. For the Wyoming site (16,230 persons), the genetic effects committed due to aquatic wastes released during the operation of the model surface mine are only 0.0001% of the genetic effects which occur due to other causes during the mine operating life. It

Table 6.28 Genetic risks to succeeding generations of an individual and committed genetic effects to descendants of the present population residing within the assessment area

Source	Genetic effects committed to succeeding generations of an individual for operation of the mine for 1 year ^(a)		Genetic effects committed to the descendants of the present population for operation of the mine for 1 year
	Maximum Individual	Average Individual	
Underground mine site (New Mexico)	4.5E-7	3.3E-8	9.0E-4
Surface mine site (Wyoming)	1.4E-8	2.0E-9	1.4E-5

^(a) Genetic effects assume 1 birth per person.

should be noted that genetic effect risks to descendants of individuals cannot be added to somatic effect risks for these individuals.

6.1.4 Nonradioactive Aquatic Emissions

Data on nonradiological emissions from uranium mines via the water pathway are limited. Table 6.29 presents available estimates of concentrations of four trace metals plus sulfate and suspended solids in discharge streams from the model surface mine located in Wyoming and seven trace metals plus sulfate and suspended solids from the model underground mine located in New Mexico. These concentrations are calculated after dilution in the first order tributaries (Appendix J) and represent average concentrations for the assessment areas. The concentrations presented in Table 6.29 are conservative since, with the exception of sulfates, loss of contaminants due to precipitation, adsorption, and infiltration to shallow aquifers are not considered. The concentrations are calculated by diluting discharges from a mine into the first order surface streams with no losses. For sulfate, a more realistic approach is taken since only 20 percent of it is assumed to remain in solution in the surface stream, as discussed in Section 3.3.3.1.4.

Also presented in Table 6.29 are recommended agricultural water concentration limits for livestock and irrigation for several of these elements (EPA73). Drinking water limits are not presented because public water supplies are normally derived from groundwater rather than surface water, so drinking water would not be a pathway of concern for the average individual in the assessment area. Though drinking water would be a potentially significant pathway for the maximum individual, the data available for this analysis did not allow a reliable prediction of groundwater concentrations due to mine dewatering (Appendix J). For this reason, the impact of nonradioactive waterborne emission on the maximum exposed individual could not be evaluated. The ratios of the average water concentrations to these limits are also listed in Table 6.29 and show that only molybdenum from the underground mine approaches its limit (irrigation). Also, the sums of the ratios being less than one indicate that mixtures of the metals would not exceed a "composite limit" for an average individual in the assessment area.

Table 6.29 Comparison of nonradiological waterborne emissions from uranium mines with recommended agricultural water quality limits

Parameter	Recommended Limits, mg/l		Model Surface Mine			Model Underground Mine		
	Livestock	Irrigation	Avg. Water	Ratio	Ratio, Avg./	Avg. Water	Ratio	Ratio, Avg./
			Conc., mg/l	Avg./Livestock Limit	Irrigation Limit	Conc., mg/l	Avg./Livestock Limit	Irrigation Limit
Arsenic	0.2	0.1	1.4E-4	0.0007	0.0014	3.1E-4	0.0016	0.0031
Barium	NA ^(a)	NA				2.0E-2		
Cadmium	0.05	0.01	1.1E-4	0.0022	0.011	1.6E-4	0.0032	0.016
Molybdenum	NA	0.01				7.0E-3		0.70
Selenium	0.05	0.02				1.6E-3	0.032	0.08
Zinc	25	2.0	4.8E-4	0.00002	0.00024	1.1E-3	0.00004	0.00055
Uranium	NA	NA	2.0E-3			3.5E-2		
Sulfate	NA	NA	4.9			2.9		
Total suspended solids	NA	NA	5.8E-1			6.8E-1		
Totals				0.0029	0.013		0.037	0.80 (0.1) ^(b)

^(a) NA - Not available.

^(b) Excluding molybdenum.

Because of the limited number of data available, it is difficult to evaluate the significance of these discharges. Although molybdenum could be a problem, it is not possible to quantify the risk from molybdenum to the maximum individual without having estimates of drinking water concentrations. Uranium, the metal estimated to be in highest concentration (Table 6.29), has no established limits based on chemical toxicity in the United States. In Canada, the maximum acceptable concentration for uranium in drinking water based on chemical toxicity has been set at 0.02 mg/l (0.04 mg/day), considering a continuous lifetime intake rate of 2 liters of water per day (HWC78). It is reasonable to assume that limits for uranium in water used for irrigation and to water livestock would exceed the drinking water limit. Hence, based on the estimated uranium concentrations at surface (0.002 mg/l) and underground (0.035 mg/l) uranium mines, the water would probably be acceptable for irrigation and livestock watering. The other constituents, such as solids and sulfates, for which limits are not available, have minimal or no toxic properties.

It is premature to conclude the health hazard caused by non-radiological waterborne emissions from uranium mines. Before definitive conclusions can be reached, additional information is needed. Of particular interest would be data on water use patterns in the vicinity of the mines and the degree to which the mine discharges may infiltrate groundwater supplies.

6.1.5 Solid Wastes

6.1.5.1 Radium-226 Content

Solid wastes, consisting of sub-ore, waste rock, and overburden, at active and inactive uranium mines contain elevated concentrations of radium-226.* The sub-ore may contain as much as 100 pCi/g of radium-226. Even though the overburden and waste rock contain lower concentrations than the sub-ore, most of these wastes contain concentrations of radium-226 in quantities greater than 5 pCi/g (see Sections 3.3.1, 3.4.1, 3.7.1, and 3.7.2).

* The radium-226 concentration in natural soil and rock is about 1 pCi/g.

Uranium mine wastes containing radium-226 in quantities greater than 5 pCi/g have been designated as "hazardous wastes" in a recently proposed EPA regulation (43FR58946, December 18, 1978) under the Resource Conservation and Recovery Act (RCRA). This is primarily due to the fact that the use of these wastes under or around habitable structures could significantly increase the chance of lung cancer to individuals occupying these structures.

6.1.5.2 Estimates of Potential Risk

We have estimated the risk of fatal lung cancer that could occur to individuals living in houses built on land contaminated by uranium mine wastes (Table 6.30). Risks were estimated for homes built on land containing radium-226 soil concentrations ranging from 5 to 30 pCi/g. The relationship between the indoor radon-222 decay product concentration and the radium-226 concentration in soil under a structure is extremely variable and depends upon many complex factors. Therefore, the data in Table 6.30 only illustrate the levels of risks that could occur to individuals living in structures built on contaminated land. These data should not be interpreted as establishing a firm relationship between radium-226 concentrations in soil and indoor radon-222 decay product concentrations.

Table 6.30 Estimated lifetime risk of fatal lung cancer to individuals living in homes built on land contaminated by uranium mine wastes

^{226}Ra in Soil (pCi/g)	Indoor Working Levels (WL)	Lifetime Risk of Fatal Lung Cancer ^(a) (per 100 persons)
5	0.02	2.5
10	0.04	5.0
20	0.08	10
30	0.12	15

(a) Based on an individual being inside the home 75 percent of the time.

The working level concentrations in Table 6.30 were derived from calculations made by Healey (He78), who estimated that 1 pCi/g of radium-226 in underlying loam-type soil would result in about 0.004 WL inside a house with an air change rate of 0.5 per hour. These calculated working levels are in reasonable agreement with measurements made by EPA (Fig. 6.1) at 21 house sites in Florida (S.T. Windham, U.S. Environmental Protection Agency, Written Communication, 1980). The Florida data were derived from the average radium-226 concentration in soil (core samples were taken to a maximum depth of three feet at each site) and the average radon-222 decay product concentration inside each structure.

6.1.5.3 Using Radium Bearing Wastes In The Construction of Habitable Structures

Wastes containing elevated levels of radium-226 have been used at a number of locations in the construction of habitable structures. In Grand Junction, Colorado, uranium mill tailings were widely used as landfill under and around the foundations of homes and other structures causing high radon-222 decay product concentrations inside many structures. To remedy this situation, Public Law 92-314 was passed in 1972 to establish a federal-state remedial action program to correct the affected structures. In Mesa County, Colorado, which includes Grand Junction, uranium mill tailings were identified at about 6,000 locations. About 800 of these locations are expected to receive corrective action because the radon decay product concentrations inside buildings constructed at these locations exceeded the remedial action criteria (DOE79). According to the criteria, dwellings and schoolhouses would be recommended for remedial action if the indoor radon decay product concentration exceeded 0.01 WL above background; other structures would be recommended for remedial action if the indoor radon decay product concentration exceeded 0.03 WL above background.

In central Florida, structures have been built on reclaimed phosphate land. The reclaimed land is composed of phosphate mining wastes that contain elevated radium-226 concentrations. EPA estimates that about 1,500 to 4,000 residential or commercial structures are located on 7,500 acres of the total 50,000 acres of reclaimed phosphate-mined lands (EPA79). A survey of 93 structures built on reclaimed phosphate land showed that about 40 percent of the structures had indoor radon-222 decay product concentrations in excess of 0.01 WL and about 20 percent had concentrations in

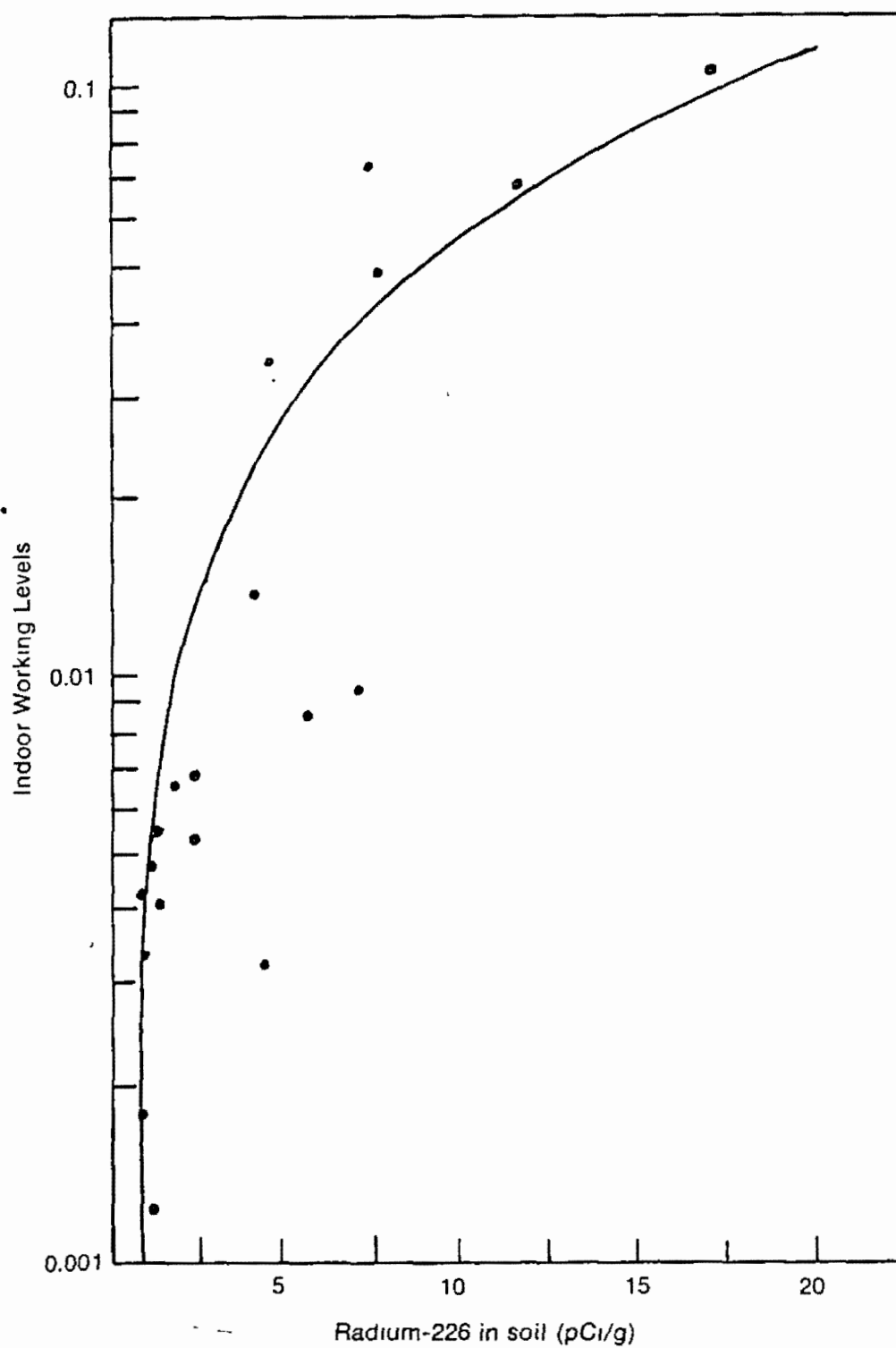


Figure 6.1 Average indoor radon-222 decay product measurements (in working levels) as a function of average radium-226 concentration in soil

excess of 0.03 WL (EPA79). Lifetime residency in a structure with a radon-222 decay product concentration of 0.03 WL could result in twice the normal 3 to 4 percent risk of fatal lung cancer.

6.1.5.3.1 Use of Uranium Mine Wastes

We do not know to what extent the wastes from uranium mines have been removed from mining sites and used in local and nearby communities. However, while surveying in 1972 for locations with higher-than-normal gamma radiation in the Western States to locate uranium mill tailings material used in local communities, EPA and AEC identified more than 500 locations where "uranium ore" was believed to be the source of the elevated gamma radiation (ORP73). The specific type of ore (mill-grade, sub-ore, low-grade waste rock) was not determined as this was beyond the scope of the survey. At some locations, however, surveyors attempted to characterize the ore by using such terms as "ore spillage," "ore specimens," "low-grade crushed ore," or "mine waste dump material." Some locations were identified as sites of former ore-buying stations (ORP73).

Since it is unlikely that valuable mill-grade ore would have been widely available for off-site use, we suspect that uranium mine waste (perhaps sub-ore) may be the source of the elevated gamma radiation levels at many of the locations where large quantities of ore material are present. Table 6.31 shows the locations where higher-than-normal gamma radiation levels were detected during these surveys and the suspected sources of the elevated levels.

6.2 Environmental Effects

6.2.1 General Considerations

Minerals are necessary to augment man's existence and welfare; in order to obtain them, some form of mining is necessary. The very nature of mining requires disturbing the land surface, but may be considered transitory. To discuss the environmental effects of uranium mining in particular, it is convenient to divide the mining operations into three phases. The first phase includes the exploration for, and the delineation of, the ore body. This involves, in most cases, substantial exploratory and development drilling. The second phase involves the preparation of the mine site and the mining process itself. This phase includes the construction of service areas, dewatering impoundments, and access roads, digging or drilling of mine entries, etc. During the actual mining process, waste

Table 6.31 Gamma radiation anomalies and causes

Location	Number of Anomalies Detected	Tailings	Cause of Anomaly			
			Uranium Ore	Radioactive Source	Natural Radioactivity	Unknown
<u>Arizona</u>						
Cane Valley (a)	19	15	4			
Cameron	3		1			2
Cutter	5		4	1		
Tuba City	17	7			3	7
State Total	44	22	9	1	3	9
<u>Colorado (b)</u>						
Cameo	3	1				2
Canon City	187	36	24		99	28
Clifton	1083	159	31	3	14	876
Collbran	145	4	2			139
Craig	86	8	7		46	25
Debeque	109	2			1	106
Delta	43	1	3		29	10
Dove Creek	83	59	17	2	2	3
Durango	354	118	18	49	67	102
Fruita	1276	58	47	1	26	1144
Gateway	17	12	1	1		3
Glade Park	1	1				
Grand Valley	110	10	2			98
Gunnison	47	3	8	1	28	7
Leadville	91	18	2		65	6
Loma	199	10	3	1	4	181
Mack	90	6	2	1		82
Mesa	123	1	1			120
Mesa Lakes	3					3
Molina	43					43
Naturita	33	10	15	5	1	2
Nucila	13	3	6		2	2
Palisade	939	107	36	3	14	779
Plateau City	28	1				27
Rifle	810	168	20	7	1	614
Salida	64	6	2		52	4
Slick Rock	9	3	5	1		
Uravan	209	208				1
Whitewater	55		4		2	49
State Total	6253	1013	256	75	453	4456
<u>Idaho</u>						
Idaho City	3				2	1
Lowman	12	9			3	
Salmon	77	1	2		65	9
State Total	92	10	2		70	10

Table 6.31 (continued)

Table 6.31 (Continued)						
Location	Number of Anomalies Detected	Tailings	Cause of Anomaly			
			Uranium Ore	Radioactive Source	Natural Radioactivity	Unknown
<u>New Mexico</u>						
Bluewater	2	1	1			
Gamerco	5				5	
Grants	101	7	49	1	25	19
Milan	41	5	23	4	1	8
Shiprock	9	8	1	0		
State Total	158	21	74	5	31	27
<u>Oregon</u>						
Lakeview	18		2		10	6
New Pine Creek	4		1			3
State Total	22		3		10	9
<u>South Dakota</u>						
Edgemont	55	43	2	1	1	8
Hot Springs	45		3		17	25
Provo	4	3		1		
State Total	104	46	5	2	18	33
<u>Texas</u>						
Campbellton	7		1		6	
Coughran	1				1	
Falls City	5	2			3	
Fashing	1		1			
Floresville	16				14	2
George West	10				10	
Karnes City	10	2			6	2
Kenedy	22	1	1		13	7
Panna Maria	3				3	
Pawnee	1		1			
Pleasanton	21		1	2	17	1
Poth	15				14	1
Three Rivers	5	1			2	2
Tilden	11				11	
Whitsett	1				1	
State Total	129	6	5	2	101	15

Table 6.31 (continued)

Table 8.31 (Continued)		Cause of Anomaly				
Location	Number of Anomalies Detected	Cause of Anomaly				
		Tailings	Uranium Ore	Radioactive Source	Natural Radioactivity	Unknown
<u>Utah</u>						
Blanding	38	10	21		3	4
Bluff	2		1			1
Cisco	2		2			
Crescent Junction	2		1			1
Green River	23	1	14		1	7
Magna	27	1	1	1	21	3
Mexican Hat	5		4	1		
Mexican Hat (Old Mill)	14	10	1	2	1	
Moab	125	15	76	7	6	21
Monticello	59	31	16	3		9
Salt Lake City ^(c)	225	70	10	5	76	64
Thompson	30	26	3	0		1
State Total	552	164	150	19	108	111
<u>Washington</u>						
Creston	3				3	
Ford	1				1	
Reardan	10				10	
Springdale	2				2	
State Total	16				16	
<u>Wyoming</u>						
Hudson	8		2		5	1
Jeffery City	28	13	9	1	3	2
Lander	86	4	8	1	53	20
Riverton	86	15	14	1	33	23
Shirley Basin	9	9				
State Total	217	41	33	3	94	46
Totals	7587	1323	537	107	904	4716

(a) From EPA report ORP/LV-75-2, August 1975. Cane Valley was not included in initial gamma survey program.

(b) Excluding Grand Junction where non-tailings anomalies were not sub-categorized according to source.

(c) Salt Lake City was not completely surveyed.

Source: ORP73.

piles are produced, mine vents drilled or reamed, and pits opened and sometimes closed. In the third or retirement phase, the site is subject to deterioration from weathering *ad infinitum*. The extent of the deterioration depends somewhat on the amount and quality of reclamation conducted during this phase.

6.2.2 Effects of Mine Dewatering

Both surface and underground mines are dewatered in order to excavate or sink shafts and to penetrate and remove the ore body. Dewatering is by ditches, sumps, and drill holes within the mine or by high capacity wells peripheral to the mine and associated shafts. Dewatering rates up to $4 \times 10^5 \text{ m}^3/\text{day}$ have been reported in the literature. Average discharge for the surface and underground mines modeled herein are 3.0 and $2.0 \text{ m}^3/\text{min}$ -ute/mine, respectively. Between 33 and 72 new mines are projected in the San Juan Basin of New Mexico alone. Total annual discharge is expected to exceed $1.48 \times 10^9 \text{ m}^3$. Calculated effects include decreased flow in the San Juan ($0.05 \text{ m}^3/\text{min}$) and the Rio Grande ($0.85 \text{ m}^3/\text{min}$) rivers. Future mining will be primarily underground and the average mine depth will increase 275 percent, i.e., from 248 m to 681 m. Average mine discharge is expected to increase from $2.42 \text{ m}^3/\text{min}$ to $13.8 \text{ m}^3/\text{min}$.

Aside from the hydraulic and water quality effects of discharging copious quantities of mine water to typically ephemeral streams, dewatering impacts are receiving increasing scrutiny because of the observed and calculated impacts on regional water availability and quality. Declines of water levels in regionally-significant aquifers of New Mexico and reduced base flow to surface streams are expected. Water quality effects relating to inter-aquifer connection and water transfer as a result of both dewatering and exploratory drilling have not been evaluated in any uranium mining area. In several Texas uranium districts, the effects of massive dewatering associated with surface mining are beginning to receive attention, but definitive studies have not yet begun and regulatory action is not expected in the near future. With respect to *in situ* leach mining, dewatering is not necessary and hence is not a concern. There is, however, some question concerning the practice of pumping large volumes of groundwater to restore aquifers. It is likely that both dewatering and aquifer restoration practices will come under increasing State regulation in water-short areas, particularly in areas of designated groundwater basins or where aquifers connect with fully-appropriated surface streams. The uncertainties surrounding environmental impacts of mining in this area can be

expected to increase, and additional, comprehensive investigations of the effects of mine dewatering and wastewater discharge are needed. Expansion in Wyoming and Texas surface and in situ leaching operations is similar, and these areas should be included in future investigations.

Uranium in water removed from mines through deliberate pumping or gravity flow is extracted for sale when the concentration is 2 to 3 mg/l or more. If there is subsequent discharge to surface water, radium-226 is also removed down to concentrations of 2 to 4 pCi/l to comply with NPDES permit conditions. Use of settling ponds at the mines also reduces total suspended solids and may reduce other dissolved constituents as a result of aeration and coprecipitation. Seepage from such settling ponds is believed to be low and, therefore, environmentally insignificant relative to groundwater. Management of waterborne solid wastes is inconsistent from one mine to another. In some cases, the solids are collected and put in with mill tailings, but in most cases they remain at the mine portal and are covered over.

For surface versus underground mines, we recognize certain inconsistencies in the parameters chosen to calculate contaminant loading of streams. Contaminant loadings from a model surface uranium mine were calculated for uranium, radium, TSS, sulfate, zinc, cadmium, and arsenic. As noted in Section 3.3.1, molybdenum, selenium, manganese, vanadium, copper, zinc, and lead are commonly associated with uranium deposits; however, there were too few data for the latter elements to develop an "average" condition. In addition, barium, iron, and magnesium can be abundant in New Mexico uranium deposits. There were insufficient data for these elements in the case of surface uranium mines in Wyoming, hence contaminant loadings were not calculated. Regional differences dictate which parameters are monitored for baseline definition and NPDES purposes. Not all potential contaminants are important in every region. For this reason and others, State and industry monitoring programs are inconsistent with respect to parameters. Since the scope of this study did not permit extensive field surveys, maximum reliance was placed on published, readily-available data.

In terms of parameters and concentrations, NPDES permit limits are inconsistent from one EPA Region to another and from one facility to another in a given Region. In part, this reflects previous screening of the effluent discharge data and natural variations in the chemistry of ore bodies.

However, the inconsistencies in parameters included and concentration limits are sufficiently large as to suggest reevaluating NPDES permits and specifying more consistent limits that more closely reflect contaminant concentrations and volumes of mine discharge.

Infiltration of most of the mine discharge in Wyoming and New Mexico is confirmed by field observations from these States. The modeling results agree with these field data. Furthermore, the modeling results, i.e., maximum infiltration, are consistent with those in the generic assessment of uranium milling (NRC79). Potable aquifers are defined under the Safe Drinking Water Act as those which contain less than 10,000 mg/l TDS. Shallow groundwater throughout the uranium regions of the U.S. meets this criterion.

Considering that essentially all of the mine effluent infiltrates and is a source of recharge to shallow potable aquifers, NPDES limits should be influenced by the drinking water regulations and ambient groundwater quality. The latter is essentially never considered with respect to mine discharges. Extensive use of soils in both the saturated and unsaturated zones as sinks for significant masses of both water and toxic chemical constituents originating in the mine discharge necessitates further evaluation of the fate of these elements. Present understanding of fractionation and resuspension processes affecting stable and radioactive trace elements greatly limits accurate prediction of health and environmental effects of mine discharge.

6.2.3 Erosion of Mined Lands and Associated Wastes

Increased erosion and sediment yield result from mining activities ranging from initial exploration through the postoperative phase. Access roads and drilling pads and bare piles of overburden/waste rock and sub-ore constitute the most significant waste sources. Dispersal is by overland flow originating as precipitation and snowmelt. To a lesser extent, wind also transports wastes and sub-ore to the offsite environment. Underground mining is much less disruptive to the surface terrain than is surface mining. Documentation of the processes and removal rates is scarce and consists of isolated studies in Texas, Wyoming, and New Mexico. Conservatively assuming that sediment yields characteristic of the areas containing the mines also apply to the mine wastes, yields of overburden, waste rock, ore, and sub-ore amount to 90,000 m³ per year. Total sediment

yield from all mining sources, including exploration and development activities, is estimated at $6.3 \times 10^6 \text{ m}^3$.

Actual erosion rates from specific sources could be considerably above or below this value owing to such variables as pile shape and slope, degree of induration and grain size, vegetative cover, and local climatic patterns and cycles. Slope instability does present serious uranium mine waste problems throughout the mountainous uranium mining areas of Colorado (S.M. Kelsey, State of Colorado, written communication, 1979). Field observations in four western states confirm that some erosion characterizes essentially every pile but that proper reclamation, particularly grading and plant cover, provides marked improvement and may actually reduce sediment loss to below pre-mining levels. Unstabilized overburden, waste rock, and sub-ore piles revegetate rather slowly, even in areas of ample rainfall such as south Texas.

Stable trace metals such as molybdenum, selenium, arsenic, manganese, vanadium, copper, zinc, and lead are commonly associated with uranium ore and may cause deleterious environmental and health effects. Mercury and cadmium are rarely present. There is no apparent relationship between the concentration of trace metals and ore grade. In New Mexico ores, selenium, barium, iron, potassium, magnesium, manganese, and vanadium are most abundant. Presently, very few data are available to characterize the trace metal concentrations in overburden rock. Results of trace metal analyses of a few grab samples from several uranium mines in New Mexico and one in Wyoming show that except for selenium, vanadium, and arsenic, no significant trend attributable to uranium mining was present (N.A. Wogman, Battelle Pacific Northwest Laboratory, Written Communication, 1979). Considering the background concentration for these elements and the limited number of analyses, the inference of offsite contamination based on these elements is indefinite.

Ore storage piles, used to hold ore at the mine for periods averaging one month, are potential sources of contamination to the environment via dusts suspended and transported by the wind, precipitation runoff, and Rn-222 exhalation--all of which can be significantly reduced by proper management. Similarly, spoil piles remaining as a result of overburden, waste rock, or sub-ore accumulations left on the land surface after mining constitute a source of contaminants for transport by wind and water. Waste particles enriched in stable and radioactive solids and Rn-222 can be

transported by wind and precipitation runoff. Such transport can be reduced through proper grading and installation of soil covers protected by vegetation or rip-rap.

Soil samples collected from ephemeral drainage courses downgrade from inactive uranium mines in New Mexico and Wyoming generally revealed no significant offsite movement of contaminants (See Appendix G). For the New Mexico mines studied, Ra-226 was elevated to about twice local background at distances of 100 to 500 meters from the mine. Water and soil samples from a surface mining site in Wyoming showed no significant offsite movement of mine-related pollution although some local transport of stockpiled ore was evident in drainage areas on and immediately adjacent to one mine pit. The strongest evidence that mine wastes are a source of local soil and water contamination is the radiochemical data and uranium in particular. Substantial disequilibrium between radium and uranium may indicate leaching and remobilization of uranium, although disequilibrium in the ore body is also suspect.

6.2.4 Land Disturbance from Exploratory and Development Drilling

About 1.3×10^6 exploratory and development drill holes have been drilled through 1977 by the uranium mining industry (see Section 3.6.1). Using the estimated land area of 0.51 hectares disturbed per drill hole (Pe79), about 6.5×10^5 hectares of land have been disturbed by drilling through 1977. To further refine the estimates of land areas disturbed, we reviewed some recent drilling areas at three mine sites. From observing 187 recent drill sites, it was concluded that 0.015 ± 0.006 hectares per drill pad were physically disturbed. The error term for the estimates is at the 95 percent confidence level. The land area disturbed by roads to gain access to the drill sites was also estimated from aerial photography and amounts to 0.17 ± 0.11 hectares. The error term for this estimate is also at the 95 percent confidence level. The total area disturbed per drill site (drill pad and access roads) is 0.19 ± 0.11 hectares. Using the latter estimates from aerial photography, the total land area disturbed from all drilling through 1977 ranges from about 1000 to 4000 km² with a mean of about 2500 km². Drilling wastes removed from the boreholes can disturb additional land areas through wind and water erosion. Ore and sub-ore remaining in the drilling wastes can, in a radiological sense, disturb land areas around the drill site from erosion. The extent of the



Figure 6 2 Example of natural reclamation of drill sites

radiological contamination at drill sites is not known and cannot presently be estimated.

Some reversal of the initial environmental damage at older drill sites was also observed from aerial photographs. Figure 6.2 contains a typical medium-to-large surface uranium mine and some adjacent drilling areas that show the effects of weathering. New drill sites are in the upper left-hand corner of the photograph. The access roads and drill pads are plainly visible. It also appears that exposed drilling wastes remain at the drill site. The area left of center in the photograph shows drill sites that are probably intermediate in age. The drilling wastes remaining have very little voluntary vegetation growing on them, and appear to have been subject to wind erosion. Weathering of the drill pads and access roads is obvious, as they are hardly discernible. It appears, in these cases, that weathering may be considered a natural reclamation phenomenon. Old drill holes are located in the lower left corner of the photograph. The drilling wastes appear to be isolated dots; the drill pads and roads are almost indistinguishable from the surrounding terrain. It appears that weathering and volunteer plant growth tend to obscure scarring caused by roads located in relatively level areas. In Figure 6.3, an underground mine site, the access roads to the adjacent drill sites required extensive excavation because of the topography. These more severe excavation "scars" will probably remain for a long period of time.

In summary, the average number of drill holes per mine can be estimated by dividing the total number of holes drilled through 1977 by the number of active and inactive mines in existence in 1977:

$$\frac{1.3 \times 10^6 \text{ drill holes}}{3300 \text{ mines}} \approx 400 \frac{\text{drill holes}}{\text{mine}} \quad (6.1)$$

The total land area physically disturbed from drilling per mine is

$$400 \frac{\text{drill holes}}{\text{mine}} \times \frac{0.19 \text{ hectares}}{\text{drill hole}} \times \frac{\text{km}^2}{100 \text{ hectares}} = \frac{0.76 \text{ km}^2}{\text{mine}} \quad (6.2)$$

In some instances, weathering and volunteer plant growth (natural reclamation) tend to restore the land areas disturbed by drilling. In others, especially on rugged topography where extensive excavation has occurred, weathering may promote extensive erosion rather than natural reclamation. Any ore or sub-ore remaining at the drill sites is subject to erosion.

6.2.5 Land Disturbance from Mining

6.2.5.1 Underground Mines

At underground mines, some land area must be disturbed to accommodate equipment, buildings, wastes, vehicle parking, and so on. The disturbed area may range widely between mines in the same area or in different geographical areas. The land area disturbed by 10 mines was estimated from aerial photographs. Nine of the mines were in New Mexico and one was in Wyoming. The disturbed land area averaged 9.3 hectares per mine site and ranged from 0.89 to 17 hectares. Access roads for each mine site consumed about 1.1 hectares on the average and ranged from 0.20 to 2.59 hectares. Subsidence or the collapse of the underground workings also causes some land disturbances. An estimated 2.8 km^2 of land has subsided as a result of uranium mining in New Mexico from 1930-71 (Pa74). A crude estimate of the land disturbed from subsidence per mine can be made by dividing the subsided area by the number of inactive underground mines in New Mexico. This amounts to about 1.5 hectares per mine. The total area (mine site, access roads, and subsidence) disturbed by an underground mine is estimated to be 12 hectares.

6.2.5.2 Surface Mines

An estimate of land disturbed from surface mining was also made from aerial photographs of eight mining sites in New Mexico and two in Wyoming. The area estimates are for a single pit or a group of interconnected pits, including the area covered by mine wastes. The average disturbed area was estimated to be about 40.5 hectares and ranged from 1.1 to 154 hectares. Access roads for the pits averaged 2.95 hectares (0.03 km^2) and ranged from 0.18 to 18 hectares. The total area disturbed per mine site is about 44 hectares.

6.2.6 Retirement Phase

The actual exploration and mining of the uranium ore constitutes a very small portion of the total existence time of a mine when considered over a large time frame. The natural forces of erosion and weathering, as well as plant growth, will eventually change any work or alterations that man has made on the landscape. For example, underground mines may eventually collapse and fill with water if they are in a water table; waste piles erode and disperse in the environment; the sharp edges of pits become

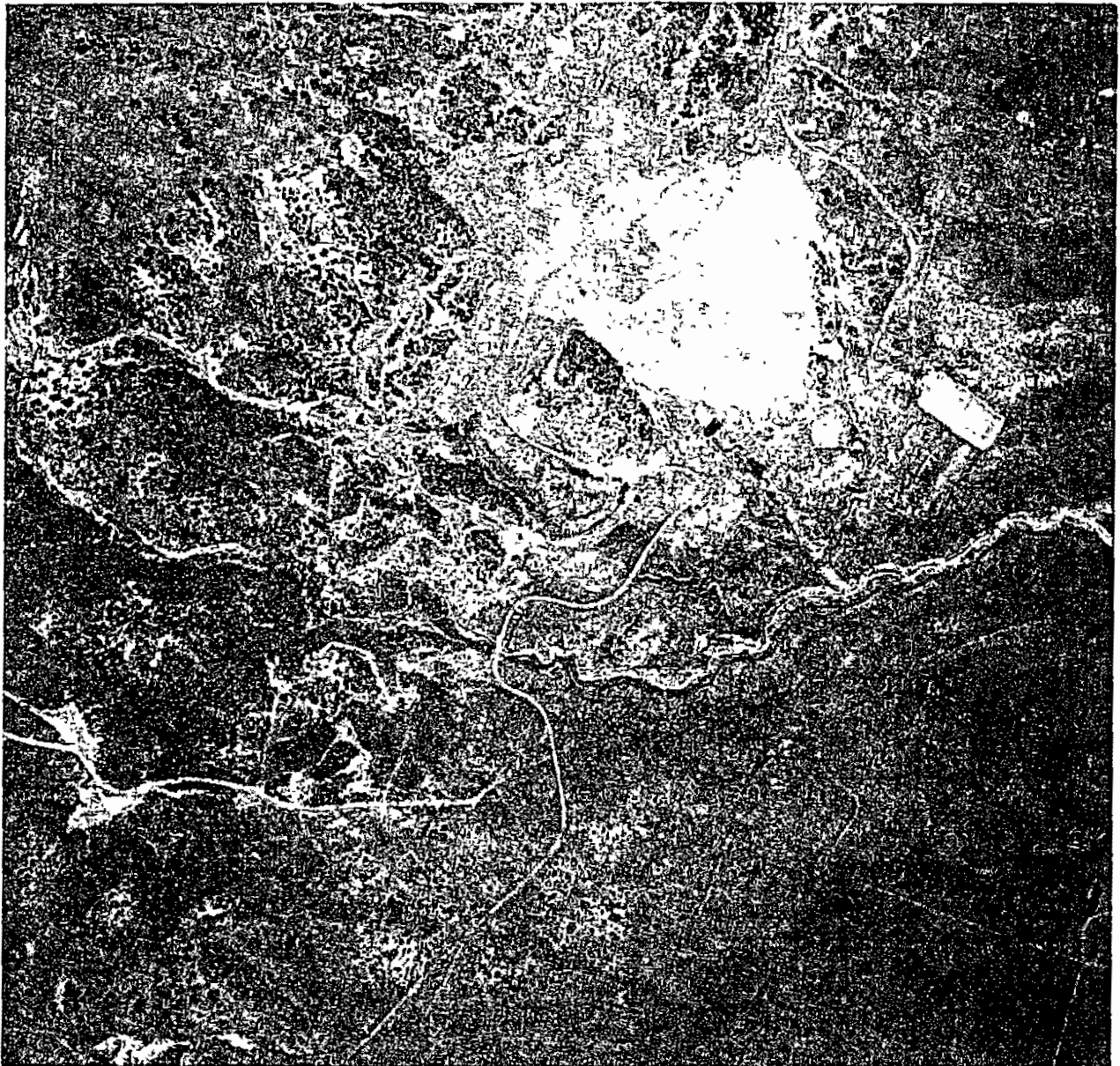


Figure 6.3 Inactive underground mine site.

smooth from wind and water erosion; lakes that are produced in pits fill up with sediment; vents and mine entries collapse, etc.

Perhaps one of the more important considerations associated with allowing a mine site to be naturally reclaimed is the dispersal of the mine wastes. Their removal from underground and subsequent storage on the surface constitute a technological enhancement of both radioactive materials and trace metals, creating a low-level radioactive materials disposal site. It appears that containment of the wastes would be preferred over their dispersal. Wastes from underground mines deposited near the entries are subject to substantial erosion. Figure 6.3 is an aerial photograph of an inactive underground uranium mine. The large light area is the waste pile and the small pile nearby is a heap-leach area. Erosion is occurring on both. A possible solution to this problem is to minimize the amount of wastes brought to the surface by backfilling mined-out areas. Another technique to minimize the dispersal of wastes into the environment by containment is to stabilize them. Unfortunately, a substantial quantity of wastes from past mining activities have been dumped in depressions and washes, which, in essence, enhances their dispersion into the environment. In retrospect, the wastes should have been stored in areas where minimal erosion would occur and then covered with sufficient topsoil to promote plant growth.

In surface mining, radiological containment can be accommodated by keeping the topsoil, waste rock, and sub-ores segregated during their removal. When backfilling, the materials can be returned to the pit in the order they were removed or in an order that would enhance the radiological quality of the ground surface. In this manner, the wastes would be contained and essentially removed from the biosphere. Figure 6.4 shows some examples of inactive and active surface mines. Some weathering and natural revegetation are noticeable around the inactive pits. Revegetation, on the other hand, appears to be relatively sparse at other inactive pits.

Erosion in inactive mining areas in New Mexico and Texas can result in deep gulying of mine waste and overburden piles. The mine wastes blanketing the foreground of Figure 6.5 are incised by an ephemeral stream that has been subsequently crossed by a roadbed in the immediate foreground. This particular mine, located in the Mesa Montanosa area immediately south of Ambrosia Lake, New Mexico, was active from 1957 to 1964. Thus, erosion occurred in about 15 years. In the background is a large mine waste pile,

the toe of which is being undercut by the same ephemeral stream (Fig. 6.6). No deliberate revegetation of the mine wastes dumped in either discrete piles or spread over the landscape (Fig. 6.7) is occurring, due in large part to the unfavorable physical and chemical characteristics of the wastes. The wastes are devoid of organic matter and are enriched in stable and radioactive trace elements, some of which are toxic to plant life. Low rainfall and poor moisture retention characteristics further suppress vegetative growth. As shown in Fig. 6.7, there is a sharp contrast between the vegetative cover on mine wastes versus that on the undisturbed rangeland in the background. Waste rock from many if not most of the mines in New Mexico, Utah, and Colorado is weakly cemented sandstone with numerous shale partings. Physical breakdown to loose, easily-eroded soil unsuitable for plant life is common (Fig. 6.8), and transport by overland flow and ephemeral streams occurs both during and long after the period of active mining (Fig. 6.9).

Depending on the degree of reclamation, if any, inactive surface mines in Texas vary considerably in the degree of erosion and revegetation. For example, the deep gullying shown in Fig. 6.10 developed in a period of one year. The mine wastes in this case were not contoured or covered to minimize gamma radiation, excessive erosion, or revegetation. In fact, the wastes were disturbed and shifted very recently in the course of constructing the holding pond (for mine water pumped from an active mine to the right of the picture) in the background. Drainage in this instance is internal, i.e., to a holding pond. In the background are more recent mine waste piles also showing deep gullying, scant vegetation, and lack of protective soil covering. Mine wastes in Texas are not completely returned to the mine primarily because of the excessive cost. As in the case of most mining operations, the bulking factor makes it physically impossible to completely dispose of the wastes in the mines.

Surface mines in Texas, particularly the older ones, also have associated overland flow to the offsite environment. Shown in Fig. 6.11 is a principal channel floored by unstabilized mine wastes and draining toward nearby grazing lands. Numerous deer and doves also were observed in the area and are actively pursued by sportsmen. The unstabilized mine in this photograph was last active several years ago, but most activity stopped in 1964. Vegetation has been very slow to reestablish and is essentially limited to a very hardy, drought-resistant willow shown in the center of the picture.



Figure 6.4 Example of active and inactive surface mining activities



Figure 6.5 Mine wastes eroded by ephemeral streams in the Mesa Montanosa area, New Mexico.

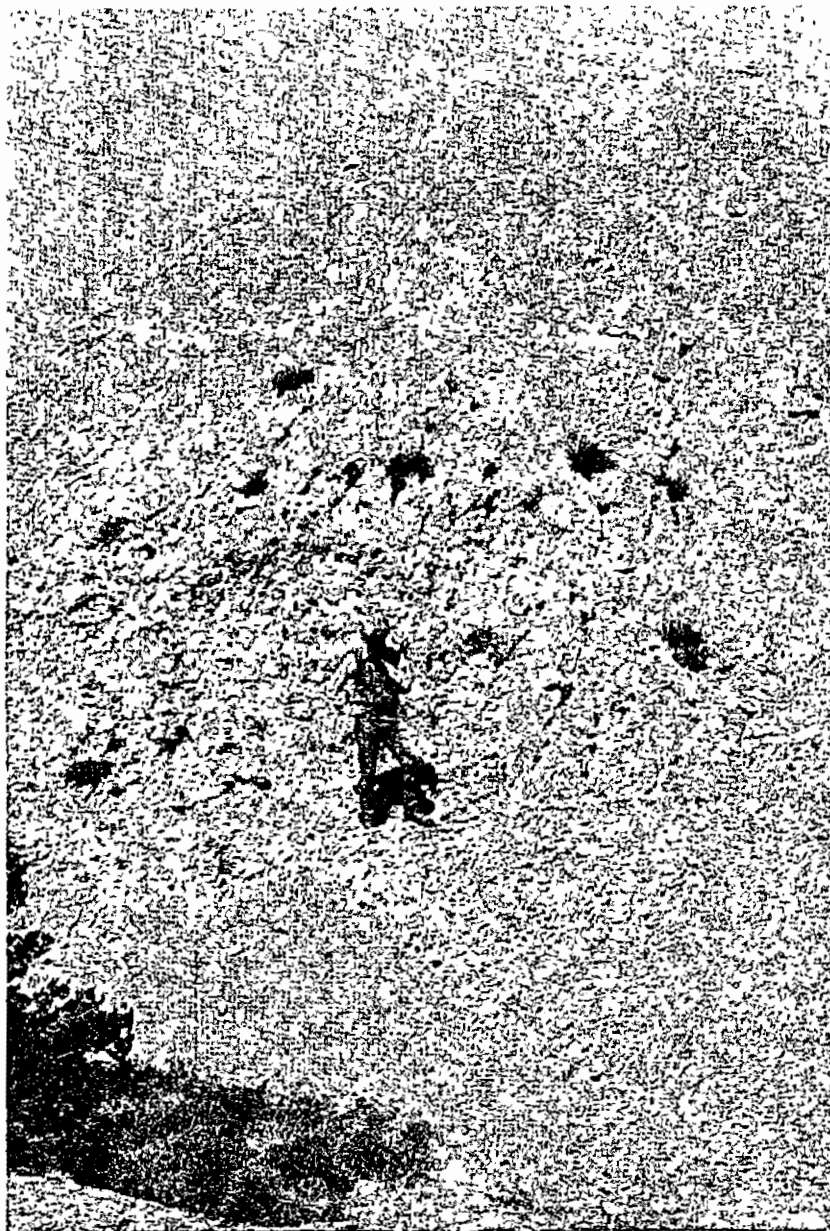


Figure 6 6 Basal erosion of a uranium mine waste pile by an ephemeral stream in the Mesa Montanosa area, New Mexico.



Figure 6.7 Scattered piles of mine waste at the Mesa Top Mine, Mesa Montanosa, New Mexico. Note the paucity of vegetation. Columnar object in background is a ventilation shaft casing

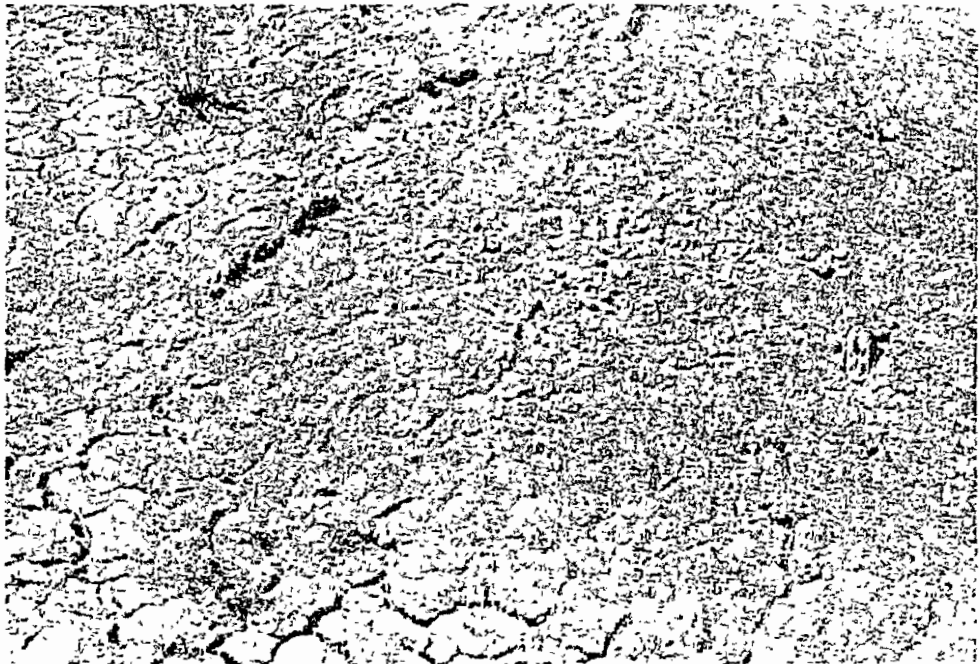


Figure 6.8 Close-up view of easily eroded sandy and silty mine waste from the Mesa Top Mine, Mesa Montanosa, New Mexico

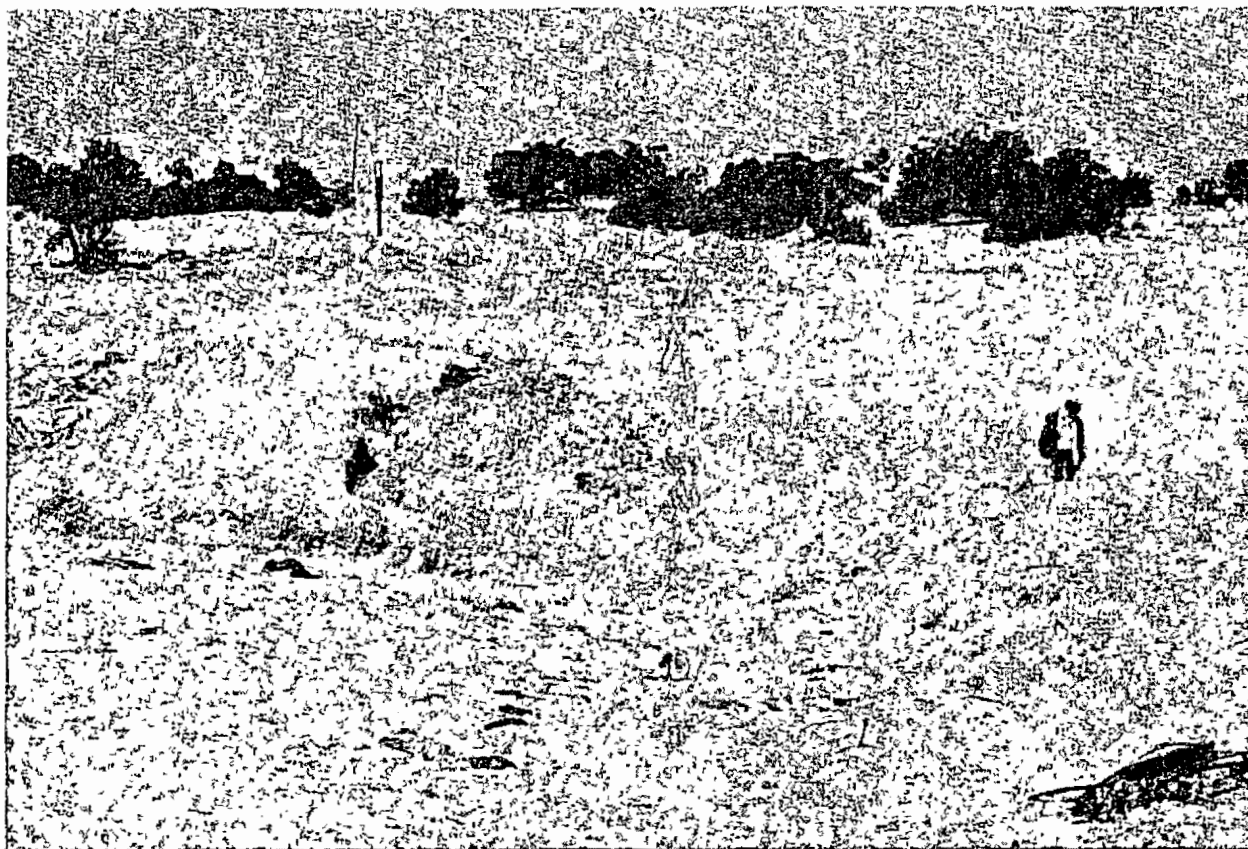


Figure 6.9 Gullying and sheet erosion of piled and spread mine wastes at the Dog Incline uranium mine, Mesa Montanosa, New Mexico.



Figure 6.10 Recent erosion of unstabilized overburden piles at the inactive Galen mine, Karnes County, Texas



Figure 6.11 Unstabilized overburden piles and surface water erosion at the Galen Mine, Karnes County, Texas.

Mines stabilized within the last few years feature improved final contouring and use of topsoil and seeding to stimulate revegetation. The reclaimed spoil piles are then available for grazing. Because backfill cannot be complete (due to economic and bulking factors), part of the mine pit remains as shown in Figs. 6.12 and 6.13, which are of the same mine. The aerial view shows extensive patches of light colored soil devoid of vegetation. Here topsoil is missing and revegetation is minimal despite the 5 years elapsed since mining. Figure 6.13 is a closeup of one portion of the mine showing deep gullying, a thin layer of dark topsoil over relatively infertile sand and silt, and the vertical mine walls. Excavations like this must be fenced. They are a hazard to livestock and people. It is likely that erosion will continue to spread away from the mine; but the rate and consequence is unknown.

Although a mine site can be reclaimed to produce an acceptable aesthetic effect, it may not be suitable in a radiological sense. At the conclusion of surface mining, the remaining pit will contain exposed sub-ore on some of the pit walls and pit floor. Because most mines at least partly fill with water and the ore zone is thereby covered, gamma radiation and radon diffusion should be markedly reduced. Although water accumulation in the pit would be expected to have elevated concentrations of trace metals and radioactive materials, this condition would probably be temporary because of the eventual covering of the pit by sedimentation from inflow of surface water and materials sloughed from the pit walls. The natural reclamation process could be enhanced by tapering the pit walls to a more gradual slope and depositing the materials on the pit floor. If sub-ores are allowed to remain near the surface, gamma exposure rates may be sufficient to prevent unlimited land use and, even if enough stabilizing materials were used to suppress the gamma radiation, radon exhalation probably could prevent unrestricted land use also. Some of the possible radiation problems could be reduced by separating the waste rock and sub-ore when hauled to the surface. The waste rock could then be used as a blanket for the sub-ore. Away from the pit proper, surface gamma readings must be below $62 \mu\text{R/hr}$ to comply with Texas State regulations. It is reasoned that, since background is about $5 \mu\text{R/hr}$, surface gamma radiation of $57 \mu\text{R/hr}$ or less would cause a total body dose of 500 mrem/yr or less.

A number of the older mines in Texas were active in the late 1950's and early 1960's--before there were requirements for stabilization. Such



Figure 6 12 Aerial view of the Manka Mine, Karnes County, Texas. Note the extent of the mine pit and associated waste piles with poor vegetative growth on bare wastes or those with insufficient topsoil cover.



Figure 6 13 Overburden pile showing the weak vegetative cover and gulying associated with improper stabilization at the Manka Mine, Karnes County, Texas. Mine stabilized in 1974

mines, one of which is shown in Fig. 6.14, are relatively shallow, contain shallow pools of water, and have high associated gamma radiation on the order of 80 to 100 μ R/hr and as much as 140 to 250 μ R/hr in some areas. The particular mine in Fig. 6.14 has maximum readings of 400 μ R/hr on the mine waste piles. In addition, the mine was used for illegal disposal of toxic wastes, primarily styrene, tars, and unidentified ceramic or refraction nodules. Some of the drums containing the wastes are shown in the rear center and right of the photograph.

Mine wastes may be used for construction and other purposes if they are not controlled or restricted (see Sections 5.4 and 6.1.5.3.1). These wastes have been used for fill in a yard and park (Appendix G). Possibly they have also been used in a school area and fairgrounds (Th79). Their use in dwelling construction has also been reported (Ha74). It is also common practice to use mine wastes for road ballast and fill in areas around mine sites. This type of usage is evident from the roads immediately adjacent to and located north and northeast of the mine shown in Fig. 6.3.

In summary, only about six percent of the land used for uranium mining has been reclaimed from 1930-71 (Pa74). For the most part, the wastes at the mine sites are spreading as a result of weathering and erosion. It appears that the wastes can be controlled or disposed of by altering some mining practices, which would require very little effort or expense on the part of the mining industry. Any reclamation of the mine sites should be keyed to long-term, natural reclamation that will continue indefinitely. Careful planning can hasten the natural reclamation process and insure long-term stability of the mine sites. Measures should be taken to prevent the removal of mine wastes.

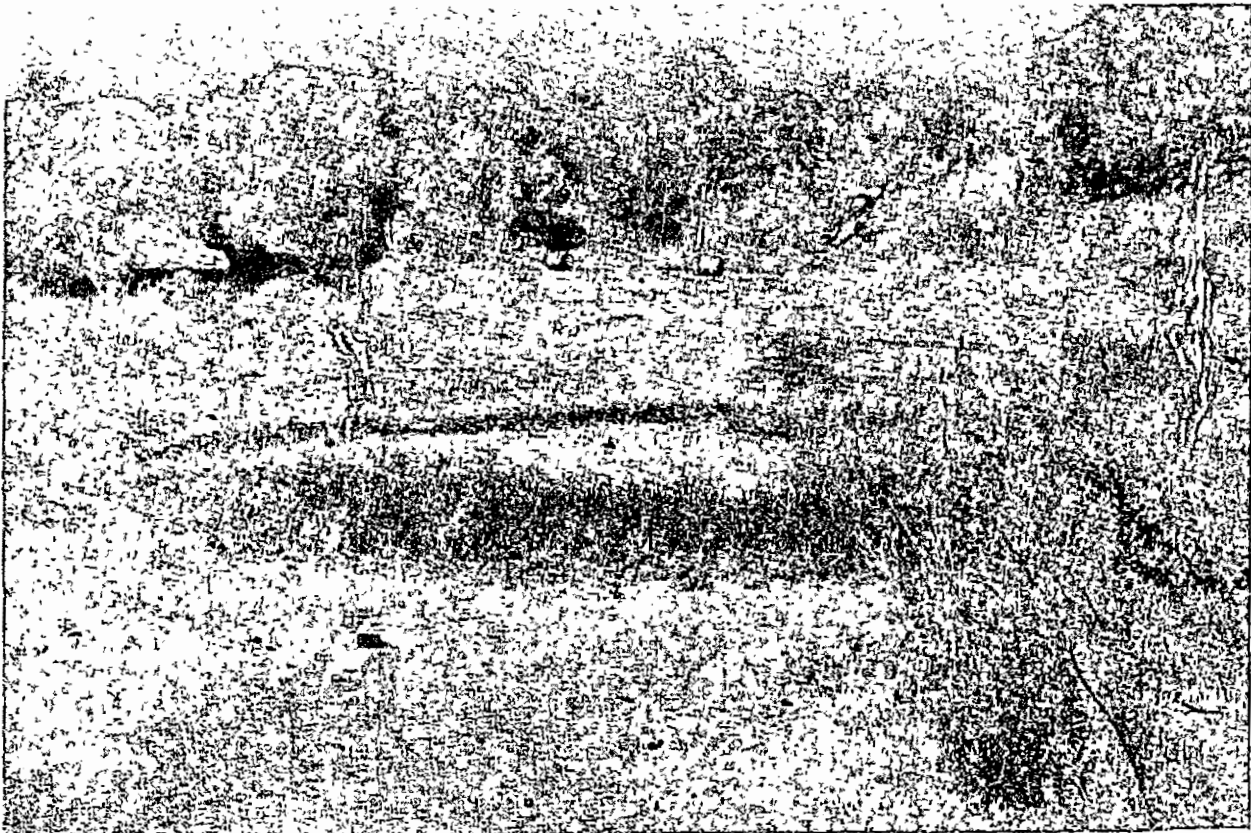


Figure 6 14 Inactive Hackney Mine, Karnes County, Texas. Drums in background contained toxic liquid wastes and styrene. Mine was active in late 1950's and early 1960.

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SECTION 7

SUMMARY AND RECOMMENDATIONS

7.0 Summary and Recommendations

7.1 Overview

This report describes the potential health and environmental effects caused by uranium mines. It considers all contaminants--solid, liquid, and airborne--and presents doses and health effects caused by wastes at both active and inactive mines. In addition to outlining the various methods of mining uranium, the report graphically depicts mine locations and lists the U.S. total of 340 active and 3,389 inactive uranium mines (Appendixes E and F) according to mine name, owner, location (state, county, section-township-range), and total ore production. Table 7.1 summarizes the mine lists.

Several facts and limitations helped shape the method and approach of this study. Little information on uranium mines is available; measurement information that is available on uranium mine wastes is frequently influenced (biased) by nearby uranium mills; there are inherent variations between uranium mines, especially between in situ mines, that complicate generic assessments of uranium mine wastes; and, finally, the law (P.L. 95-604) that mandated this study allotted only a short time in which to complete it. To accommodate these facts in our study plan, we decided to develop conceptual models of uranium mines and to make health and environmental projections from them, based upon available data from the literature; to employ conservative (maximizing) assumptions when necessary; and to supplement available information with information from discussions with persons inside and outside the agency and by doing several field studies in Texas, New Mexico, and Wyoming. Table 7.2 summarizes the sources of uranium mine contaminants that were modeled in this study.

7.2 Sources and Concentrations of Contaminants

7.2.1 Surface and Underground Mines

We calculated released radioactivity for two models of active underground and surface uranium mines. The average-large mine, the first model, reflects new and predicted future mines. The average mine, the second model, reflects the regional impact of multiple mines. The quality and

Table 7.1 Distribution of United States uranium mines by type of mine and state

State	Active				Inactive		
	Surface	Under-ground	In situ leaching	All ^(a) Others	Surface	Under-ground	All Other ^(a)
Alaska	0	0	0	0	0	1	0
Arizona	1	1	0	0	135	189	2
California	0	0	0	0	13	10	0
Colorado	5	106	0	4	263	902	52
Florida	0	0	0	0	0	0	1
Idaho	0	0	0	0	2	4	0
Minnesota	0	0	0	0	0	0	1
Montana	0	0	0	0	9	9	0
Nevada	0	0	0	0	9	12	0
New Jersey	0	0	0	0	0	1	0
New Mexico	4	35	0	3	34	142	12
N. Dakota	0	0	0	0	13	0	0
Oklahoma	0	0	0	0	3	0	0
Oregon	0	0	0	0	2	1	0
S. Dakota	0	0	0	0	111	30	0
Texas	16	0	8	1	38	0	4
Utah	13	108	0	3	378	698	17
Washington	2	0	0	0	13	0	0
Wyoming	19	6	3	2	223	32	10
Unknown	<u>0</u>	<u>0</u>	<u>0</u>	<u>0</u>	<u>6</u>	<u>5</u>	<u>2</u>
Total	60	256	11	13	1252	2036	101

(a) Includes mine water, heap leach dumps, miscellaneous, and unknown.

Table 7.2. Sources of contaminants at uranium mines

Source	Active Underground	Active Surface	Inactive Underground	Inactive Surface
<u>Waste Rock (Overburden) Pile</u>				
Wind suspended dust	M	M	M	M
Rn-222 emanation	M	M	M	M
Precipitation runoff	C	C	C	C
<u>Sub-Ore Pile</u>				
Wind suspended dust	M	M	M	M
Rn-222 emanation	M	M	M	M
Precipitation runoff	C	C	C	C
<u>Ore Stockpile</u>				
Wind suspended dust	M	M	M	M
Rn-222 emanation	M	M	M	M
Precipitation runoff	C	C	C	C
<u>Abandoned Mine Area Surfaces</u>				
Rn-222 emanation	M	M	M	M
<u>Mining Activities</u>				
Dusts	M	M	NA	NA
Combustion products	M	M	NA	NA
Rn-222	M	M	NA	NA
<u>Wastewater</u>				
Surface discharge	M	M	NA	NA
Seepage	C	C	C	C

Note.--M, Source modeled; C, considered but not modeled due to lack of information; NA, not applicable.

flow rates that were determined for water discharges from typical surface and underground mines in Wyoming and New Mexico, respectively, were used to calculate chemical loading of streams in three hydrographic units: sub-basin (containing the mines), basin, and regional basin. Infiltration of mine water to potable groundwater and suspension/solution of contaminants in flood waters are the main components of the aqueous pathway. Crude dilution and infiltration models were used to evaluate aqueous discharge from active mines. Off-site movement from inactive mines is primarily by overland flow, the contamination significance of which was evaluated with limited field and literature surveys.

Concentrations of radionuclides and stable elements in waste rock, sub-ore, and ore, selected from only a few measurements, are shown in Table 7.3. Average annual airborne emissions for the sources listed in Table 7.2 were computed for active and inactive mines using the concentrations listed in Table 7.3 and the geological and meteorological information appropriate for each region. Source terms were maximized by assuming no dust control and no spoils pile restoration. Annual emissions of airborne contaminants estimated for the various sources are given in the following tables of Section 3.

Source	<u>Tables on Active Mines</u>		<u>Tables on Inactive Mines</u>	
	Surface	Underground	Surface	Underground
Combustion Products	3.30	3.52	--	--
Vehicular Dusts	3.32	3.56	--	--
Dust from Mining				
Activities	3.33	3.54	--	--
Wind Suspended Dust	3.34	3.55	3.70	3.76
Radon-222 Emissions	3.35	3.51	3.74	3.77

Annual emissions in mine water discharged to the surface by the model average underground and surface mines are listed below.

Parameter	Surface Mine (Wyoming)	Underground Mine (New Mexico)
Flow rate, m ³ /min	3.0	2.0
Uranium-238, Ci/yr	0.037	0.49
Uranium-234, Ci/yr	0.037	0.49
Radium-226, Ci/yr ^(b)	0.00065	0.0014
Radon-222 and each short-lived daughter, Ci/yr	0.00065	0.0014
Lead-210, Ci/yr	0.00065	0.0014
Polonium-210, Ci/yr	0.00065	0.0014
Arsenic, Kg/yr	7.9	13
Barium, Kg/yr	ND ^(a)	850
Cadmium, Kg/yr	6.3	7
Molybdenum, Kg/yr	ND	300
Selenium, Kg/yr	ND	70
Sulfate, MT/yr ^(b)	276	122
Zinc, Kg/yr	112	45
Total suspended solids, MT/yr	33.0	29

(a) No data available.

(b) The values shown for radium-226 and sulfate are 10 percent and 20 percent, respectively, of those released on an annual basis. Radium is assumed to be irreversibly sorbed, and sulfate readily infiltrates.

Table 7.3. Concentration of contaminants in waste rock (overburden), ore, and sub-ore

Nonradioactive					
Stable Element	Concentration, $\mu\text{g/g}$		Stable Element	Concentration, $\mu\text{g/g}$	
	Waste Rock	Ore and Sub-ore		Waste Rock	Ore and Sub-ore
Arsenic	9	86	Manganese	485	960
Barium	290	920	Molybdenum	2.5	115
Cadmium	NA	ND	Potassium	7,000	25,000
Cobalt	NA	16	Lead	22	78
Copper	18	61	Ruthenium	NA	ND
Chromium	<51	20	Selenium	2	110
Iron	6,000	15,700	Strontium	150	130
Mercury	<8	ND	Vanadium	100	1,410
Magnesium	NA	3,500	Zinc	20	29
Radioactive					
Radioactive Contaminant	Concentration, pCi/g				
	Waste rock	Sub-ore	Ore		
U-238 and each daughter	6	(a)	285		
Th-232 and each daughter	1	2	10		

(a) The concentration of U-238 and each daughter was assumed to be 99 pCi/g at active underground mines, 40 pCi/g at active surface mines, and 110 pCi/g at inactive mines of both types.

Note.--NA, Not available; ND, Not detected.

7.2.2 In Situ Leach Mines

The sources of airborne releases that we assessed at our model in situ leach mine were the uranium recovery and packaging unit, the evaporation ponds, and the surge tank. The annual releases for these sources are listed below.

Source	Annual Airborne Release Rate
<u>Recovery Plant</u>	
Uranium-238	0.10 Ci
Uranium-234	0.10 Ci
Uranium-235	0.0048 Ci
Thorium-230	0.0017 Ci
Radium-226	0.00010 Ci
Lead-210	0.00010 Ci
Polonium-210	0.00010 Ci
Ammonia	3.2 MT
Ammonium chloride	12 MT
Carbon dioxide	680 MT
<u>Surge Tank</u>	
Radon-222	650 Ci
<u>Storage Ponds</u>	
Ammonia	100 MT
Ammonium chloride	300 MT
Carbon dioxide	80 MT

Since in situ mining is site specific and relatively new, little information is available on its wastes. Thus, only airborne releases were assessed quantitatively; liquid and solid wastes were discussed qualitatively.

Several characteristics of in situ mining, especially regarding its liquid and solid wastes, tend to minimize its release of contaminants.

First, only a small fraction of Ra-226 is leached (2.5 percent assumed); second, all liquid wastes are impounded with no planned releases; third, much of the liquid waste evaporates, except at a few sites in Texas where the wastes are injected into deep wells; and, finally, at in situ mines solid wastes accumulate at a much lower rate than they do at conventional mines. Aquifer restoration and underground excursion of the leaching solution were also discussed qualitatively. Although restoration has not yet been done at a commercial scale site, preliminary experiments indicate that proper aquifer restoration is possible. During the restoration process, Rn-222 will continue to be purged from the aquifer and should be considered a possible source of exposure.

7.2.3 Uranium Exploration

During exploration and developmental drilling, dusts are produced, Rn-222 and combustion products from drilling equipment are released, and approximately 0.2 hectares of land surface are disturbed per drill hole. The average mine site produces an estimated 6,100 kg of airborne dust, 20 kg of which is ore and subore. About 3400 Ci of Rn-222 are released annually from all development holes drilled since 1948 (4.5×10^5), which is similar to that released from one operating mine. Combustion product releases are small.

7.3 Exposure Pathways

Exposures were assessed for a hypothetical most exposed individual living about 1600 m (1-mile) from the center of the mine and for a population residing within an 80-km (50-mile) radius of the mine. The meteorological and geological parameters used were those appropriate to the respective sites.

Aqueous releases were modeled through a basin, sub-basin, and regional basin hydrographic area. Dilution by precipitation, snowmelt, and periodic flooding (typical of semiarid regions) was analyzed but not used in the model. For the model we assumed that the average annual release of contaminants is diluted by the average annual flow rate of the stream being considered. The pathways that we assessed are listed below.

Air Pathways

1. Breathing
 - a. Radioactive particulates and radon-222
 - b. Radon-222 daughters
2. External Exposure
 - a. Submersion.
 - b. Surface deposited radioactivity
3. Eating
 - a. Above-surface foods grown in the area
 - b. Milk and beef cattle grazing in the area

Water Pathways

1. Breathing
 - a. Resuspended contaminants deposited from irrigation water
2. External Exposure
 - a. Submersion in resuspended contaminants deposited from irrigation water
3. Eating
 - a. Above-surface foods grown in the area
 - b. Milk and beef cattle grazing in the area and drinking contaminated water
 - c. Fish

In addition to the risks caused by wastes at or discharged directly from the mines, we assessed the risks to occupants of habitable structures built on land containing uranium mine wastes. The radium-226 in these wastes increases the concentrations of radon-222 and its decay products and the gamma radiation inside these structures.

7.4 Potential Health Effects

7.4.1 Radioactive Airborne Emissions

The risks of fatal cancer were estimated for radioactive airborne emissions. They include the lifetime risk to the most and average exposed individuals in the regional population and the number of additional fatal cancers in the regional population caused per year of model mine operation (see Table 7.4).

The major fatal cancer risk at each of the model uranium mines is the risk of lung cancer from Rn-222 daughter exposures (Tables 6.11 and 6.12). At surface and in situ mines, radioactive particulates plus Rn-222 contribute only a little over 10 percent of the total fatal cancer risk. The principal radionuclides in the airborne particulate emissions are U-238, U-234, Th-230, Ra-226, and Po-210. The contribution from Th-232 and its daughters is minor. At underground mines, essentially all the risks are due to Rn-222 daughter exposures. Fatal cancer risks at active underground mines are greater than those at active surface mines because of the larger quantity of Rn-222 daughter products released. For inactive mines, the risks are similar at surface and underground sites.

Most of the exposure to individuals around the model uranium mines is received internally, usually by breathing. However, the average person in the region around surface mines receives most of his exposure by eating contaminated foods. The largest contributors to the radioactive particulate plus Rn-222 impact are ore and overburden at active surface mines and ore and sub-ore at the active underground mines. For the model in situ mine, the uranium processing plant was the main source of particulate radionuclides.

Of all evaluated model uranium mines, the average large underground mine (Table 7.4) causes the largest fatal cancer risk and the largest number of additional cancers in the regional population. Compared to the natural occurrence of fatal cancer from all causes (Table 7.5), we estimate an increase of 1.3 percent (0.0019) in fatal cancers over the lifetime of the maximum individual and a 0.0003 percent (0.018) increase in fatal cancers in the regional population per year. Increases in expected fatal cancers are less at all other model mine sites.

Compared to a normal occurrence of genetic effects of 0.06 effects/birth and 12.1 effects/year in the regional population (Wyoming), the computed risk of additional genetic effects from radiation exposure at the model uranium mines is very small. The average large surface mine produces the largest increase in genetic effects. We estimate the genetic risk to the descendants of the most exposed individual to be an additional $6.4\text{E-}5$ effects/birth (0.1 percent increase) for a 30-year parental exposure; $2.2\text{E-}7$ effects/birth (0.00036 percent increase) to the descendants of the average exposed individual in the regional population for the same

Table 7.4 Summary of fatal cancer risks from radioactive airborne emissions of model uranium mines

Source	Most exposed individual life-time fatal cancer risk (a)	Average exposed individual life-time fatal cancer risk (a)	Fatal cancers caused in regional population per year
Average Surface Mine	1.3E-4	2.5E-7	1.7E-4
Average Large Surface Mine	4.2E-4	8.1E-7	6.4E-4
Average Underground Mine	2.0E-4	9.1E-7	1.7E-3
Average Large Underground Mine	1.9E-3	8.6E-6	1.8E-2
Inactive Surface Mine	3.4E-5	6.3E-8	1.3E-5
Inactive Underground Mine	2.0E-5	8.6E-8	4.5E-5
In Situ Leach Mine	2.2E-4	3.9E-7	3.1E-4

(a) Lifetime exposures were calculated as follows:

Surface and underground mines: Exposure for 17 years to active mining and 54 years to inactive mine effluents.

Inactive mines: Exposure for 71 years to inactive mine effluents.

In situ leach mine: Exposure for 10-year operation and 8-year restoration.

Table 7.5 Percent additional lifetime fatal cancer risk for a lifetime exposure to the individual and the percent additional cancer deaths in the regional population per year of exposure estimated to occur as a result of uranium mining

Source	Most Exposed Individual	Average Exposed Individual	Regional Population
Average surface mine	8.7E-2	1.7E-4	7.9E-6
Average large surface mine	2.8E-1	5.4E-4	3.0E-5
Average underground mine	1.3E-1	6.1E-4	3.1E-5
Average large underground mine	1.3	5.7E-3	3.3E-4
Inactive surface mine	2.3E-2	4.2E-5	6.1E-7
Inactive underground mine	1.3E-2	5.7E-5	8.3E-7
In situ leach mine	1.5E-1	2.6E-4	1.4E-5

Note.--Comparisons are based on the risks given in Table 7.4, a national cancer risk from all causes of 0.15, and an estimate of the cancer death rate from all causes to the regional populations of New Mexico (5,400 deaths) and Wyoming (2,140 deaths).

exposure period; and $7.9\text{E-}5$ additional genetic effects committed to the descendants of the regional population per year of mine operation. The latter increase is very small compared to the 12.1 effects that will normally occur each year in the live births of the regional population.

7.4.2 Nonradioactive Airborne Emissions

Atmospheric concentrations of nonradioactive air pollutants were calculated at the location of the most exposed individual. The concentrations were compared with calculated nonoccupational threshold limit values, natural background concentrations, and average urban concentrations of selected airborne pollutants in the United States.

Of the pollutant sources investigated, three produced insignificant health hazards:

1. airborne stable trace metals
2. airborne combustion products from heavy equipment operation
3. nonradioactive gas emissions at in situ leach mines

However, at active surface mines, dust particulates (produced mainly by vehicular traffic) equal or exceed conservatively calculated nonoccupational threshold limit values and, therefore, are a potential nuisance.

7.4.3 Radioactive Aqueous Emissions

The only water from active uranium mines is that pumped from the mines and released to surface streams. The largest radiation dose* from this water to individuals in the assessment regions is to the endosteal cells (bone) (see Tables 6.25 and 6.26). It primarily comes from eating foods grown on land irrigated by streams fed by discharged mine water. Significant, but of lesser importance, are exposures due to breathing wind suspended material from irrigated land, eating fish caught in streams near the site, and external gamma radiation from land irrigated by streams fed by mine water discharges. We estimate only a small risk from eating beef and milk from cattle grazing on irrigated pasture and drinking water contaminated by mine discharges (<2 percent of the total risk from aqueous emissions). The radionuclides of major importance in the risk analyses are U-238 and U-234.

*In Section 7, "dose" is to be read as "dose equivalent"--absorbed radiation (dose) multiplied by a quality factor.

The risks of fatal cancer were estimated for radioactive aqueous discharges to surface streams from active uranium mines. The estimates included, for the 17-years of active mine operation, the cumulative risk to the most and average exposed individuals in the assessment area and the number of fatal cancers caused to persons residing within the assessment area (Table 7.6). Aqueous emissions from inactive mines and from in situ leach mines were not modeled due to a lack of data. However, we believe aqueous source terms from these mines would be low.

Drinking water may be an important source of exposure for the most exposed individual living near a uranium mine. However, we did not estimate it because we could not quantify radionuclide concentrations in potable groundwater with available information. Also, mine water probably is not consumed directly by man.

Table 7.6 Summary of the fatal cancer risks caused by radioactive aqueous emissions from model uranium mines

Source	Most exposed individual's life-time fatal cancer risk for 17 years of mine operation	Average exposed individual's life-time fatal cancer risk for 17 years of mine operation	Fatal cancers caused in the assessment area population from 17 years of mine operation
Underground mine site (New Mexico)	$5.6\text{E-}6(3.7\text{E-}3\%)^{(a)}$	$3.4\text{E-}7(2.3\text{E-}4\%)$	$2.2\text{E-}2(2.3\text{E-}4\%)$
Surface mine site (Wyoming)	$1.2\text{E-}7(8.0\text{E-}5\%)$	$1.6\text{E-}8(1.1\text{E-}5\%)$	$2.6\text{E-}4(1.1\text{E-}5\%)$

(a) All "risks" in this table are in addition to the 0.15 risk of fatal cancer from all causes.

Although aqueous discharges from the model underground mine produce greater risks than those from the model surface mine, primarily because of greater releases of U-238 and U-238 daughters, aqueous releases at either mine cause only very small cancer risks (see Table 7.6) beyond the 0.15

natural risk of fatal cancer. For example, in New Mexico (assessment population 64,950) and Wyoming (assessment population 16,230), 9,742 and 2,434 deaths from cancer from all causes are projected to occur. Aqueous mine discharges in these areas will add only 0.022 and 0.00026 estimated deaths, respectively, to these totals.

The largest increase in estimated genetic effects occurs at the underground mine site. However, compared to the natural occurrence of hereditary disease, the overall risk of additional genetic effects due to radionuclides discharged in water from the model mines is very small. Based on a natural occurrence of 0.06 effects/birth, there will be 936 genetic effects in the regional population of New Mexico during 17 years of mine operation. In contrast, there will be only 0.015 additional effects to all the descendants of the regional population because of the 17-year exposure period.

7.4.4 Nonradioactive Aqueous Emissions

Aqueous concentrations of nonradioactive pollutants were calculated for stream water we assumed was used by the average individual within the assessment area. The pathways considered are those listed in Section 7.3. Drinking water might be a significant pathway for the most exposed individual. However, we could not make a reliable prediction of increased groundwater concentrations due to mine dewatering with the available data.

A comparison of the water concentrations of several pollutants with recommended EPA limits for livestock and irrigation usage (see Table 6.29) showed that only molybdenum from the underground mine approaches its limit for irrigation. The sums of the ratios of the average water concentrations to the recommended limits are less than one, indicating that mixtures of the metals would not exceed a "composite limit" for an average individual in the assessment areas. Constituents such as solids and sulfates, for which limits are unavailable, have minimal or no toxic properties.

More information is needed before definitive conclusions can be reached about health hazards caused by nonradioactive waterborne emissions. Uranium, the metal estimated to be in highest concentration, has no established limits based on chemical toxicity in the United States. Of particular interest would be data on water use patterns near the mines and the degree to which mine discharges may infiltrate groundwater supplies.

7.4.5 Solid Wastes

We estimated the risk of fatal lung cancer to individuals living in houses built on land contaminated by uranium mine wastes as a function of the Ra-226 concentration in the wastes (see Table 7.7). How much mine waste has been used for homesite land fill as well as its level(s) of contamination are unknown. Because of the cost, it is unlikely that mill-grade ore would be available for off-site use. It is more likely that waste rock, perhaps mixed with some sub-ore, would be the material used. Considering the Ra-226 content of sub-ores and the likelihood of its being diluted with waste rock and native soil, mine wastes in residential areas would probably contain between 5 to 20 pCi/gm of Ra-226.

Table 7.7 Estimated lifetime risk of fatal lung cancer to the average person living in a home built on land contaminated by uranium mine wastes

^{226}Ra in Soil (pCi/g)	Indoor Working Levels (WL)	Lifetime Risk of Fatal Lung Cancer ^(a)
5	0.02	0.025
10	0.04	0.050
20	0.08	0.10
30	0.12	0.15

^(a) Based on the average individual being inside his home 75 percent of the time.

7.5 Environmental Impacts

We evaluated the environmental effects of uranium mining, including exploration, by reviewing completed studies, extensive communications with State and Federal agencies, field studies in Wyoming and New Mexico, reconnaissance visits to Wyoming, Colorado, New Mexico, and Texas, and imagery collection and interpretation. Underground and surface mines were examined to develop a sense of an average or typical condition with respect to mine size, land areas affected, quality and quantity of airborne and

waterborne releases, and general, qualitative appreciation for the effects of such operations on surface streams, groundwater, disturbed land areas, and natural recovery processes. In many instances, conditions can be documented, but the significance remains highly subjective and thus weakens the justification for corrective action, particularly for inactive mines.

7.5.1 Land and Water Contamination

We conclude that (1) U.S. uranium mills make little use of mine water; (2) mine drainage is to the environment, with occasional use for agriculture, sand backfilling, construction, and potable supply; (3) active surface mines in Wyoming and underground mines in New Mexico have the greatest discharge to the offsite environment; (4) inactive surface mines do not appear to adversely affect groundwater quality, although water in such mines is typically contaminated and runoff from surface accumulation of overburden and sub-ore may be a source of surface water contamination; and (5) selected inactive underground mines in Colorado and possibly adjacent portions of Utah may discharge water enriched in radionuclides and trace elements. Since the mining industry now uses terrestrial ecosystems extensively as sinks for mining-related contaminants, an appropriate government agency should monitor active mines for groundwater quality, sorption of contaminants on stream sediments, and the flushing action of flooding events.

Before and during surface and underground uranium mining, contaminated mine water is frequently discharged to arroyos and pasture lands adjacent to the mines. Less frequently, mine water is used in nearby uranium mills, in which case ultimate disposal is to the mill tailings pile where evaporation and seepage occur. However, despite this practice of mine water discharge to land and despite the existence of over 3,000 active and inactive mines and the accelerating level of exploration and mining, there are many more studies and surveys on the interaction of uranium mills and water resources than there are on uranium mines and water resources. With few exceptions, monitoring mine water quality has been related to NPDES permits.

When mines discharge water to open lands and water courses, 90 percent or more of it infiltrates the soil and the balance evaporates. Stable and radioactive contaminants subject to sorption are selectively concentrated

in nearby soils, which become a local sink. Mobile constituents such as sulfate and chloride probably percolate to the water table along with the bulk of the water, which recharges nearby shallow aquifers downgrade from the mines. Although many areas in New Mexico, Texas, Colorado, Wyoming, and Utah have received mine water discharge, studies of contaminant accretion on soils and deterioration of groundwater quality have been rather limited. Widespread contamination of groundwater has not been documented, but there are indications that local surface water and groundwater quality have been adversely effected in Colorado, Wyoming, and Texas. Studies underway in New Mexico reveal, in at least two mining districts, groundwater deteriorating because of mine drainage. Significant increases in ambient uranium and radium occurred in the Shirley Basin uranium district of Wyoming because of initial strip mining and mill processing and, to a lesser extent, in situ leaching. The long-term significance of soil loading with stable and radioactive contaminants and their cycling through the terrestrial ecosystem, including the human food chain, has not been determined for uranium mining operations.

Discharges from model active surface and underground mines average 2 to 3 m³/minute. In most cases, complete infiltration takes place in stream beds within 5 to 10 kilometers of the mines. However, when discharges from several mines are combined or if single mine discharge is several cubic meters per minute or more, infiltration and storage capacity of the alluvium in nearby channels is exceeded and perennial flows are created for distances of 20 to 30 kilometers. For example, underground uranium mines in the Grants Mineral Belt of New Mexico currently discharge 66 m³ per minute. Of this, only 12 m³ per minute are used in uranium mills; the balance is discharged to nearby washes or arroyos. Fourteen of the 20 active uranium mills make no use of mine water, which is associated with essentially every active underground mine and most active surface mines, particularly in Texas and Wyoming.

Annual contaminant loading from continuous discharge at a rate of 3 m³/minute from one surface mine in the Wyoming model area and dilution in flood flows with recurrence intervals of 2 to 25 years produce the loading and stream concentration values in Table 7.8. Chemical loading was calculated on a mass-per-time basis to estimate the effects of mine drainage.

For assessing environmental impacts, we assume that most contaminants remain on or near the land surface and are available for resuspension in periodic flash flooding in the sub-basin. Sorption, precipitation, and so on are assumed to render 90 percent of the radium-226 unavailable for further transport. Eighty percent of the sulfate is assumed to infiltrate and also becomes unavailable for further transport in flood waters.

Stream concentrations for uranium, zinc, cadmium, and arsenic are likely to be less than those shown because there will not be 100 percent resuspension of sorbed contaminants, and flood events with lesser return periods are also likely to disperse contaminants. The loading data are believed to be quite realistic; it is the temporal distribution and redistribution of the contaminants that constitute a significant unknown. These preliminary results indicate contamination of surface water with uranium, radium, sulfate, and, to a lesser extent, with cadmium and arsenic in stream waters near the mine outfall. Subsequent dilution of these initial concentrations will occur as the flow merges with that of progressively larger streams in the downgrade direction, but cadmium and sulfate may exceed the drinking water standard in flood waters as far as the regional basin. Impoundment of these initial flows can be expected considering water management practices in semiarid rangeland areas like Wyoming. Therefore, further pathway investigations, based on field data, are needed.

For the model underground mining area, we selected the Ambrosia Lake District of New Mexico. We assumed that 14 mines discharged an average of $2 \text{ m}^3/\text{minute}$ and that loading took place for two years prior to each flood. We then calculated concentrations in flood water for eight different cases—for 2, 5, 10, and 25 year floods (larger numbers indicating larger floods), with concentrations for each flood being calculated on the basis of both a 1-day and 7-day flood duration (see Table 7.9). Based upon these assumptions and calculations, it appears that concentrations in flood waters, particularly in the basin, may exceed established or suggested standards for uranium, radium, cadmium, arsenic, selenium, barium, and sulfate. However, precipitation and sorption, in addition to dilution farther downstream, probably will reduce these concentrations enough so that quality standards for drinking and irrigation water can be met. But

Table 7.8 Summary of contaminant loading and stream water quality from a model surface uranium mine

Annual Loading Per Mine (a) (Kg/yr)	Drinking Water Standard (mg/l)	Concentrations in Basin and Regional Basin Flood Flows for Floods of 2, 25, and 100 Years Return Period, mg/l			
		Basin		Regional Basin	
		Min	Max	Min	Max
Uranium 110	0.015/3.5/0.21 ^(b)	0.36	0.76	0.26	0.44
Radium-226 0.00065 Ci/yr	5 pCi/l	2.1 pCi/l	4.5 pCi/l	1.6 pCi/l	2.6 pCi/l
TSS 32,955	---	107	228	79	131
Sulfate 2.76 x 10 ⁵	250	900	1909	668	1098
Zinc 112.0	5.0	0.366	0.774	0.271	0.445
Cadmium 6.31	0.01	0.02	0.044	0.015	0.025
Arsenic 7.88	0.05	0.025	0.054	0.019	0.031

(a) Loading values shown for radium and sulfate are reduced to 10 percent and 20 percent, respectively, of the amount actually released by a mine. Irreversible sorption and precipitation affect radium and sulfate infiltrates to the water table.

(b) 0.015 mg/l : Suggested maximum daily limit based on radiotoxicity for potable water consumed at a rate of 2 liters per day on a continuous basis. 3.5 mg/l : Suggested maximum daily limit based on chemical toxicity and intake of 2 liters in any one day. 0.21 mg/l : Suggested maximum daily limit based on chemical toxicity and intake of 2 liters per day for 7 days.

Table 7.9 Summary of contaminant loading and stream water quality from a model underground uranium mine

Annual Loading, Per Mine(a) (Kg/yr)	Drinking Water Standard (mg/ℓ)	Concentrations in Basin and Regional Basin for 1-day and 7-day Floods of 2 to 25 Years Return Period, mg/ℓ			
		Basin		Regional Basin	
		Min	Max	Min	Max
Uranium 1480	0.015/3.5/0.21 ^(b)	6.9	7.1	0.045	0.046
Radium-226					
0.0014 Ci/yr	5 pCi/ℓ	6.7 pCi/ℓ	6.9 pCi/ℓ	0.044 pCi/ℓ	0.044 pCi/ℓ
Lead-210					
0.0014 Ci/yr	---	71.2 pCi/ℓ	73.4 pCi/ℓ	0.470 pCi/ℓ	0.0472 pCi/ℓ
Cadmium 7	0.01	0.03	0.03	0.0002	0.0002
Arsenic 13	0.05	0.061	0.063	0.00039	0.00041
Selenium 80	0.01	0.37	0.38	0.0026	0.0026
Molybdenum 300	---	1.4	1.4	0.0089	0.0093
Barium 850	1.0	4.0	4.2	0.26	0.27
Zinc 45	5.0	0.21	0.22	0.0014	0.0014
Sulfate 1.22×10^5	250	574	584	3.7	3.8
TSS 29,000	---	130	140	0.89	0.92

(a) Loading values shown for radium and sulfate are reduced to 10 percent and 20 percent, respectively, of the amount actually released by a mine. Irreversible sorption and precipitation affect radium and sulfate infiltrates to the water table.

(b) 0.015 mg/ℓ : Suggested maximum daily limit based on radiotoxicity for potable water consumed at a rate of 2 liters per day on a continuous basis. 3.5 mg/ℓ : Suggested maximum daily limit based on chemical toxicity and intake of 2 liters in any one day. 0.21 mg/ℓ : Suggested maximum daily limit based on chemical toxicity and intake of 2 liters per day for 7 days.

more theoretical and field evaluations are needed to confirm this.

In situ leaching has contaminated local groundwater reservoirs. We expect that this will continue because leach solution excursions from the well field do occur and because injected constituents, especially ammonium, can not be fully recovered. The NRC and agreement States recognize this situation but consider the adverse impacts outweighed by the benefits of recovering additional uranium and developing a relatively new technology.

7.5.2 Effects of Mine Dewatering

Underground mines and most surface mines are dewatered to allow for excavation or shaft sinking and ore removal. The resulting low concentration and, oftentimes, large volume effluent discharges introduce substantial masses of stable and radioactive trace elements to local soil and water systems. This extensive use of soils in both the saturated and unsaturated zones as water and contaminant sinks requires further study to determine the environmental fate of those elements. In addition to local effects, the long-term impacts on regional water availability and quality are also important. The NPDES limits relating to surface discharges are, in terms of parameters and concentrations, different from one EPA region to another and should be reevaluated to more closely reflect the impact of contaminant concentration and mine discharge. In general, the uncertainties about the environmental impact of mine dewatering can be expected to increase; and additional, comprehensive investigations of its effects are necessary.

7.5.3 Erosion of Mined Lands and Associated Wastes

From initial exploration through retirement, mining, particularly surface mining, increases erosion and sediment yield. The most significant waste sources are access roads, drilling pads, and piles of overburden/waste rock and sub-ore. Sediment and associated contaminants are dispersed mostly through the overland flow of precipitation and snowmelt water. Erosion rates vary considerably with the characteristics of the source area, i.e., pile geometry, soil and rock characteristics, amount and type of vegetative cover, topography, and local climate. There is some

erosion of all mine waste sources, although studies of ephemeral drainage courses downgrade from inactive mines in New Mexico and Wyoming usually reveal only local soil and water contamination and no significant off-site dispersal of contaminants. Proper reclamation, particularly grading and revegetation, markedly reduce erosion and, consequently, contaminant transport.

7.5.4 Exploratory and Development Drilling

The uranium industry has drilled approximately 1,300,000 exploratory and development drill holes through 1977. This amounts to about 430 drill holes per mine if averaged over all active and inactive mines. During the course of drilling, some land areas are disturbed to provide access roads to the drill sites and pads for the drill-rig placements. This has disturbed about 2500 km² (960 mi²) of land for all drilling through 1977.

Drilling wastes accumulate at each drill site. Although these wastes are sometimes placed in trenches and backfilled after drilling, the general industry practice (observed from field studies and aerial photography), apparently, is to allow the wastes to remain on the surface, subject to erosion. The extent of radiological contamination from erosion of the remaining ore and sub-ore at development drill holes is not known.

The average drilling depth has increased with time and will probably continue to do so in the future. Deeper drilling will tend to increase the probability that several aquifers may be penetrated by each drill hole. Aquifers with good quality water may be degraded by being connected, via the drill holes, with aquifers of poor quality water. Current regulations require drill holes to be plugged to prevent interaquifer exchange, but often only the first one and one-half meters of the borehole will be plugged, and regulations do not effect past drill holes. Finally, it appears, from mine site surveys and aerial photography, that very few drill sites have been reclaimed.

7.5.5 Underground Mining

The land disturbed by individual underground mines varies from 0.89 to 17 hectares (2.2 to 42 acres) with an average of 9.3 hectares (23 acres). In addition, access roads to the mines consume about 1.1 hectares (2.7 acres), and mine subsidence disturbs about 1.5 hectares (3.7 acres). A

total of about 12 hectares (30 acres) of land are disturbed by an average underground mine.

All underground uranium mining through 1977 has produced about 2.9×10^7 MT or about $1.8 \times 10^7 \text{ m}^3$ of wastes. Some of these wastes, the sub-ores, contain elevated concentrations of naturally occurring radionuclides. The sub-ores usually are removed last in the mining process and dumped on top of the waste rock where they are subject to erosion. Some radiation surveys conducted around waste piles indicate that the sub-ores are eroding and contaminating land in addition to that disturbed by the mining activities.

During our field studies in Texas, New Mexico, Wyoming, and Colorado, we saw very few mine sites where reclamation had been completed or was in progress--especially at the inactive mine sites.

7.5.6 Surface Mining

The cumulative waste from surface mining uranium between 1950 and 1978 amounts to about 1.7×10^9 MT ($1.1 \times 10^9 \text{ m}^3$). Overburden is usually used to backfill mined-out pits during contemporary mining. At older inactive mines, the mine wastes were either used for pit backfill or completely disregarded. Erosion of these waste piles may cause substantial environmental problems.

The amount of land physically disturbed at a surface mine is highly variable. The area disturbed at ten surface mines was estimated to range from 1.1 to 154 hectares (2.7 to 380 acres), averaging about 41 hectares (101 acres) per mine site. Access roads disturb about 3 hectares (7.4 acres) per mine site, bringing the total average area physically disturbed to about 44 hectares (109 acres). Field surveys of inactive mine sites indicate that mine wastes (sub-ores) erode and contaminate land areas greater than those physically disturbed. The land contamination appears to have been caused by erosion of ore stockpiles, erosion of sub-ores, and dust losses from the actual mining process.

Very few if any inactive mine sites were reclaimed. Reclamation of any mine site will have to address the radiological aspects of the mine and its wastes.

7.6 Regulatory Perspective

Except for in situ leach mining, licensed by the Nuclear Regulatory Commission (NRC), uranium mining is not licensed, per se, by a Federal agency. However, three Federal statutes have particular relevance to uranium mining. First, the Federal Water Pollution Control Act as amended (1972) requires a permit for discharges to navigable waters. Second, the Clean Air Act amendments of 1977 require a permit for pollutant air emissions. Third, proposed regulations under the Resource Conservation Recovery Act of 1977 identify hazardous wastes and stipulate their disposal for uranium mining. When promulgated, these latter regulations will strengthen current Federal and State reclamation requirements.

In situ uranium mining is licensed by those states having agreement-state status with NRC. National Pollution Discharge Elimination System (NPDES) permits are issued by EPA approved states. No state issues mining licenses per se. However, most states require mining and reclamation plans, including bonding fees, for at least state-controlled lands. Most reclamation requirements provide erosion control through slope and vegetation standards. Arizona is the only uranium mining state without reclamation requirements.

7.7 Conclusions and Recommendations

The evaluation of the potential impacts of uranium mining was performed largely by means of analytical studies of model facilities. We believe that the results give an adequate representation of the industry. In order to determine the extent of possible problems, our studies were specifically designed to give conservative results. It should be recognized that actual mines may operate under conditions producing substantially smaller impacts than the results presented.

Compared to uranium milling, health and environmental effects of mining are not as well understood, despite the existence of over 3000 active and inactive mines. We have noted throughout this report instances of the absence or inadequacy of pertinent information.

7.7.1 Conclusions

7.7.1.1 Solid Wastes

Solid uranium mining wastes are potentially hazardous when used as building materials or when buildings are constructed on land containing such wastes. The hazard arises principally from increased risk of lung cancer due to radon-222. In a 1972 survey of communities in uranium mining regions, EPA and the former Atomic Energy Commission found more than 500 locations where such wastes had been used.

7.7.1.2 Airborne Effluents

a) Individuals living very near active underground mine exhaust vents would have an increased risk of lung cancer caused by exposure to radon-222 emissions. Surface mines and in situ mines are less hazardous, and inactive mines do not have significant radon-222 emissions. Other airborne radioactive emissions from all types of mines are judged to be smaller.

b) The number of additional cancers committed per year in the regional populations due to radionuclide air emission from the approximately 340 active mines and 3300 inactive mines was estimated to be about 0.6 cancers in 1978. This number of estimated additional cancers is small, about one-third of the estimated additional cancers in regional populations due to radon emissions from the 24 inactive uranium mill tailings piles addressed by Title I of the Uranium Mill Tailings Radiation Control Act (EPA 80). (These mill tailings piles represent about 13 percent of all tailings currently existing due to U.S. uranium milling and mining). These potential effects are not of sufficient magnitude to warrant corrective measures, especially considering the large number of sites involved.

c) The following were judged to cause an insignificant health risk for all types of mines:

1. airborne nonradioactive trace metals.
2. airborne combustion products from heavy-duty equipment operations.
3. nonradioactive emissions from in situ leach sites.

d) Airborne dust near large surface mines (primarily caused by vehicular traffic) may exceed the National Ambient Air Quality Standard for particulate matter.

7.7.1.3 Waterborne Effluents

a) We estimate that an insignificant health risk accrues to populations from waterborne radioactivity from an average existing mine.

b) Uranium mine dewatering and water discharges, which are increasing as more and deeper mines are created, may in the future have significant effects on water quality. Current treatment practices are controlling the release of radioactivity into surface waters.

c) Water in inactive surface and underground mines usually contains radionuclides and trace elements in concentrations comparable to groundwater in contact with ore bodies. Some abandoned underground mines in certain areas of Colorado and Utah probably discharge such waters to nearby streams and shallow aquifers. Available data is not sufficient to conclude whether or not there is a problem.

d) We could not determine, using models, that there is no health hazard to individuals who drink water drawn from such surface or underground sources. Water discharges from active mines to nearby streams and stream channels may extensively recharge shallow aquifers, many of which are either now used or could be used for drinking water. Such determinations must be made on a site-specific basis, and take account of the additive effects of multiple mines. These studies can be made easily a part of State or utility surveillance programs.

7.7.1.4 Exploratory and Development Drilling

Harm from effluents due to exploratory and developmental drilling is probably small compared to effects of operating mines. Under current regulations and practices, however, aquifers penetrated at different levels can mix, creating the potential for degrading high quality groundwater.

7.7.2 Recommendations to Congress

1) Based on this study, we do not believe at this time that Congress needs to enact a remedial action program like that for uranium mill tailings. This is principally because uranium mine wastes are lower in radioactivity and not as desirable for construction purposes as uranium mill tailings. Nonetheless, some mining waste materials appear to have been moved from the mining sites but not to the extent that mill tailings were. .

2) Some potential problems were found that might require regulatory action but none of these appear to require new Congressional action at this time.

7.8 Other Findings

1) Regulations may be needed to control wastes at active uranium mines to preclude off-site use and to minimize the health risks from these materials. These regulations would need to address the use of the materials for construction purposes as well as ultimate disposal of the materials.

EPA proposed such regulations in 1978 under the Resource Conservation and Recovery Act (RCRA). In 1980, Congress amended RCRA to require further EPA studies before promulgating general regulations for mining wastes. An EPA study by the Office of Solid Wastes on all types of mines, including uranium mines, is currently being conducted. The amendment did not restrict EPA's authority to regulate use of uranium mine wastes in construction or reclamation of lands containing such wastes.

2) Standards are needed to control human exposure from radioactive air emissions from uranium mines. This is principally because of potential exposure to individuals living near large underground uranium mines rather than concerns regarding the exposure of regional populations. We have proposed such standards under Section 112 of the Clean Air Act.

3) EPA has conducted two field studies in 1972 and 1978 which define possible sites at which mine wastes may have been used in construction or around buildings. The information developed in these studies has been sent to State health departments. The States should conduct follow-up studies, as appropriate, to determine whether there are problems at these sites.

4) The adequacy with which NPDES permits protect individuals who may obtain drinking water near the discharge points for uranium mine dewatering should be evaluated by States. Under the Public Water Systems provision of the Safe Drinking Water Act, radionuclide ~~standards~~ now exist for drinking water.

5) Some site specific studies should be considered by States to determine the extent to which inactive uranium mines are significant water pollution sources.

6) States with uranium mines should determine the feasibility of control of fugitive dust from large surface mines and incorporate the recommendations in State Implementation Plans.

7) States should require borehole plugs in drilling operations that will prevent interaquifer mixing (exchange) and also seal drilling holes at the surface.

7.9 References

EPA80 U.S. Environmental Protection Agency, 1980, "Draft Environmental Impact Statement for Remedial Action Standards for Inactive Uranium Processing Sites (40 CFR 192), "EPA 520/4-80-011.

APPENDIX A

FEDERAL LAWS, REGULATIONS, AND GUIDES FOR
URANIUM MINING

Table A.1 Federal laws, regulations, and guides for uranium mining

Federal Agency	General		Permits		Mining					Health and Safety
	Water Use	Conservation-Preservation Statutes	Exploration Rights	Mining Rights	Environmental Quality					
					Air	Water		Land		
					Surf	UG	Solids	Reclam.		
Dept. of Int.	1	2,3,4,6,7	8	8					2,8	
BIA ^(a)			9	9					9	
BLM ^(a)		5	10	10					10	
USGS ^(a)			9						9	
Dept. of Energy		2	11	11					2	
Dept. of Agr.	1	2,8d							2,8d	
USFS ^(a)			12,13	13					12	
EPA	1	2				16		19	2	20
AIR-OAQPS ^(a)					14					
Water										
Surface OWPS ^(a)						17				
Ground OSW ^(a)							19			
Land-OSW						18	18	18	18	
Radiation-ORP ^(a)					15	15	15	15	15	20
U.S. Army										
Corps. of Engrs.	1	2				16			2	
Dept. of Labor		2								
MSHA ^(a)										22
OSHA ^(a)										21
Nuclear Reg.										
Comm. ^(b)		2		2	23 ^(b)	23 ^(b)		23 ^(b)	23 ^(b)	23 ^(b)

- (a) BIA-Bureau of Indian Affairs
 BLM-Bureau of Land Management
 USGS-United States Geological Survey
 USFS-United States Forest Service
 (b) OAQPS-Office of Air Quality, Planning and Standards
 Nuclear Regulatory Commission (NRC) regulations and guides for milling do apply to in situ extraction or mining but not conventional surface or underground mining where NRC has no authority.
- OWPS-Office of Water Planning and Standards
 OSW-Office of Solid Waste
 ORP-Office of Radiation Programs
 MSHA-Mining Safety and Health Administration
 OSHA-Occupational Safety and Health Administration

Table A.1 (continued)--Key to Federal laws, regulations and guides cited

1. See Appendix B and Appendix C for U.S. Constitution Citations, Federal Laws, and Interstate Compacts
2. National Environmental Policy Act of 1969 (Public Law 92-190)
3. Endangered Species Act of 1973 (Public Law 93-205) (Supplants Endangered Species Conservation Act of 1969)
4. National Historic Preservation Act of 1966 (Public Law 89-655) (Supplants Antiquities Act of 1906)
5. Federal Land Management and Policy Act of 1976
6. Reservoir Salvage Act of 1960 (16 USCA 469-469C)
7. Historic Sites Acts of 1935 (16 USCA 21-50)
8.
 - a. U.S. Mining Law of 1872 (30 USC 21-50)
 - b. Mineral Leasing Act of 1920 (30 USC 181 et seq)
 - c. Mineral Leasing Act for Acquired Lands (Amended) (30 USC 351-359)
 - d. Materials Act of 1947 (Amended) (30 USC 601-602)
 - e. Reorganization Plan of 1946 (60 Stat. 1099)
9. Indian Land - 30 CFR 231
10. Public Land - 30 USC 22 (43 CFR 3810, 3746, 3501, 3814.1)
11. Withdrawn Public Land - 42 USC 2097
12. National Forest Land - 16 USC 478 (43 CFR 3811.1 and 36 CFR 252)
13. National Forest Management Act of 1976 (16 USC 1600) - Regulations for land and resource management planning under this Act in the National Forest System are given in Federal Register Volume 44, Number 181, September 17, 1979
14. Clean Air Act as Amended (42 USC 1857 et seq)
15. Public Health Services Act (Reorganization No. 3, 1970; Section 301 - Environmental Monitoring)

Table A.1 (continued)--Key to Federal laws, regulations and guides cited

16. Marine Protection Research and Sanctuaries Act of 1972
17. Federal Water Pollution Control Act as Amended (33 USC 466 et seq)
18. Resource Recovery and Conservation Act of 1976 (Proposed 40 CFR 250.46-4)
19. Safe Drinking Water Act Amended (Public Law 95-523 and Public Law 95-190): (Could affect mining operation where injection of wastes is utilized)
20. Atomic Energy Act Amended (Public Law 86-373; 42 USC 2021(h), Federal Radiation Guidance functions from prior Federal Radiation Council)
21. Occupational Safety and Health Act of 1970
22. MSHA formed by transferring MESA from DOI to DOL pursuant to the Federal Mine Safety and Health Act of 1977, Public Law 91-173 as amended by Public Law 95-164
23. Nuclear Regulatory Commission Guides and Regulations for Beneficiation Processes
 - a. Regulatory Guide 3.5, Standard Format and Content of License Applications for Uranium Mills (Nov. 1977)
 - b. Regulatory Guide 3.8, Preparation of Environmental Reports for Uranium Mills (Sept. 1978)
 - c. Regulatory Guide 3.11, Design, Construction, and Inspection of Embankment Retention Systems for Uranium Mills (Dec. 1977)
 - d. Regulatory Guide 4.14, Measuring, Evaluating, and Reporting Radioactivity in Releases of Radioactive Materials in Liquid and Airborne Effluents from Uranium Mills (June 1977)
 - e. Regulatory Guide 4.15, Quality Assurance for Radiological Monitoring Programs (Normal Operations) - Effluent Streams and the Environment (Feb. 1979)
 - f. Regulatory Guide 8.11, Applications of Bioassay for Uranium (June 1975)
 - g. Regulatory Guide 8.13, Instruction Concerning Prenatal Radiation Exposure
 - h. Standards for Protection Against Radiation (10 CFR 20)
 - i. Domestic Licensing of Source Material (10 CFR 40)
 - j. Licensing and Regulatory Policy and Procedures for Environmental Protection (10 CFR 51)
 - k. Proposed Regulations: Uranium Mill Tailings Licensing (10 CFR Parts 40,150) - 44 F.R. 50012, August 24, 1979

Table A.1 (continued)--Key to Federal laws, regulations, and guides cited

- l. Staff Technical Positions: Tailings Management - "Current U.S. Nuclear Regulatory Commission Licensing Review Process: Uranium Mill Tailings Management"; Environmental Monitoring - "Proposed Branch Position for Operational Radiological Environmental Monitoring Programs for Uranium Mills"
- m. Proposed Regulatory Guide 3.11.1, Operational Inspection and Surveillance of Imbankment Retention Systems for Uranium Mill Tailings (April 1979)

APPENDIX B
FEDERAL WATER PROGRAMS AND RIGHTS ACTIVITIES
AND THEIR LEAD ADMINISTRATIVE AGENCIES

DM 2000, INC
1833 Hormel
San Antonio, Texas 78219
(210)222-9124 FAX (210)222-9065

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APPENDIX C
CONGRESSIONALLY APPROVED INTERSTATE WATER COMPACTS

Interstate water compacts

Name	Year
Arkansas River Compact	1948
Arkansas River Basin Compact	1965
Bear River Compact	1955
Belle Fourche River Compact	1943
Canadian River Compact	1950
Colorado River Compact	1922
Connecticut River Flood Control Compact	1951
Costilla Creek Compact	1963
Delaware River Basin Compact	1961
Great Lakes Basin Compact	1955
Klamath River Basin Compact	1957
La Plata River Compact	1922
Merrimack River Flood Control Compact	1956
New England Interstate Water Pollution Control Compact	1947
New York Harbor (Tri-State) Interstate Sanitation Compact	1935
Ohio River Valley Water Sanitation Compact	1939
Pecos River Compact	1948
Potomac River Basin Compact	1939
Red River of the North Compact	1937
Republican River Compact	1942
Rio Grande Compact	1938
Sabine River Compact	1953
Snake River Compact	1949
South Platte River Compact	1923
Susquehanna River Basin Compact	1970
Tennessee River Basin Water Pollution Control Compact	1955
Thames River Flood Control Compact	1957
Upper Colorado River Basin Compact	1948
Wheeling Creek Watershed Protection and Flood Prevention District Compact	1967
Yellowstone River Compact	1950

Source: Environmental Study on Uranium Mills, TRW, Inc., USEPA Contract
No. 68-03-2560, February 1979.

APPENDIX D

STATE LAWS, REGULATIONS, AND GUIDES
FOR URANIUM MINING

Table D.1 State laws, regulations, and guides for uranium mining

State	General		Mining								Safety
	NRC Agreement State	NPDES Permit State	Water Use	Permits		Environmental Quality					
				Exploration Rights	Mining Rights	Air	Water		Solids	Land Reclam.	
							Surf	UG			
COLORADO	Yes	Yes	-	-	-	-	-	-	-	-	-
Department of Health											
Water Quality Control Div.	-	-	-	-	-		6,7,8,10	8,9,10	-	-	-
Air Quality Control Div.	-	-	-	-	-	4,5	-	-	-	-	-
Department of Natural Resources											
Div. of Water Reserves (State	-	-	15	-	-	-	-	-	-	-	-
Board of Land Commissioners	-	-	-	1	1	-	-	-	-	1	-
Mined Land Reclam. Bd.	-	-	-	2,3	2,3	-	2,3	2,3	-	2,3	-
Division of Mines	-	-	-	-	-	-	-	-	-	-	14
NEW MEXICO	Yes	No	-	-	-	-	-	-	-	-	-
State Land Commission	-	-	-	1	1	-	-	-	-	-	-
Dept. of Energy and Minerals	-	-	-	-	-	-	-	-	-	2	9
Dept. of Natural Resources	-	-	3	-	-	-	-	-	-	-	-
Env. Improvement Div.	-	-	-	-	-	5,6,7,8,9	10,12,13,14	11	14	-	16
TEXAS	Yes	No	-	-	-	-	-	-	-	-	-
Dept. of Water Resources	-	-	13	-	3	-	8,9	-	4,11	4	-
R.R. Commission of Texas	-	-	-	1	1	5	5	2,7	-	2	-
General Land Office	-	-	-	14	14	-	-	-	-	15	-
Dept. of Health	-	-	-	-	-	-	6,10	-	-	-	10
Air Control Board	-	-	-	-	-	12	-	-	-	-	-
UTAH	No	No	-	-	-	-	-	-	-	-	-
State Engineer	-	-	3,4	-	-	-	-	-	-	-	-
Dept. of Social Services	-	-	-	-	-	-	-	-	-	-	-
Division of Health	-	-	-	-	-	6	1	1	-	-	5
Water Pollution Control Bd.	-	-	-	-	-	-	1	1	-	-	-
Dept. of Natural Resources	-	-	-	2	2	-	-	-	-	2	-

Table D.1 (continued)

State	GENERAL			Mining							
	NRC Agreement State	NPDES Permit State	Water Use	Permits		Environmental Quality					
				Exploration Rights	Mining Rights	Air	Water		Land		Safety
							Surf	UG	Solids	Reclam.	
WASHINGTON	Yes	Yes	-	-	-	-	-	-	-	-	-
Dept. of Natural Resources	-	-	-	1,2	1,2	-	-	-	-	2	2
Dept. of Ecology	-	-	9	-	-	-	-	-	-	-	-
Office of Water Programs	-	-	-	-	-	-	8	(No)	-	-	-
Dept. of Social Services & Health	-	-	-	-	-	-	-	-	-	-	-
Health Services Division	-	-	-	-	-	-	-	-	-	-	3
Air Quality Division	-	-	-	-	-	7	-	-	-	-	-
WYOMING	No	Yes	-	-	-	-	-	-	-	-	-
State Inspector of Mines	-	-	-	8a	8a	-	-	-	-	-	8c
State Engineers Office	-	-	1	-	-	-	-	-	-	-	-
Dept. of Env. Quality	-	-	-	-	-	-	-	-	-	-	-
Air Quality Div.	-	-	-	-	-	3a,4	-	-	-	-	-
Water Quality Div.	-	-	-	-	-	-	2,5	5,6	-	-	-
Land Quality Div.	-	-	-	3c	3c	-	-	-	3c,7	3c,7	-
Solid Waste Management	-	-	-	-	-	-	-	-	3d	-	-

Table D.1 (continued)--Key to State laws, regulations, and guides cited

Colorado

1. Mining Rules and Regulations, 1973, 1976; Uranium Mining Lease and Prospecting Permit; State Board of Land Commissioners.
2. Colorado Mined Land Reclamation Act, July 1, 1976; Mined Land Reclamation Board (Act. 32, Title 34, C.R.S. 1973, as amended).
3. Rules and Regulations, Colorado Mined Land Reclamation Board; effective July 1978.
4. Colorado Air Quality Control Act of 1979, adopted June 20, 1979. Replaces Colorado Air Pollution Control Act of 1970. Radioactive materials included in list of air pollutants.
5. Colorado Air Quality Control Regulations and Ambient Air Quality Standards, Colorado Air Pollution Control Commission. Specifically, Regulation No. 1, Emission Control Regulations for Particulates, Smokes, and Sulfur Oxides for the State of Colorado; and Regulation No. 3, Regulation Governing Air Contaminant Emission Notice, Emission Permit, and Fees for Direct Sources.
6. Regulations Establishing Basic Standards and an Antidegradation Standard and Establishing a System for Classifying State Waters, for Assigning Standards, and for Granting Temporary Modifications, Colorado Water Quality Control Commission, May 22, 1979; effective July 10, 1979.
7. Regulations for Effluent Limitations, Colorado Department of Health, Water Quality Control Commission; adopted March 18, 1975 effective August 21, 1975.
8. Regulations for the State Discharge Permit System, Colorado Department of Health, Water Quality Control Commission; adopted November 19, 1974 effective January 31, 1975, amended February 7, 1978.
9. Rules for Subsurface Disposal Systems, Colorado Department of Health, Water Quality Control Commission; revised July 6, 1976, effective October 1, 1977.
10. Guidelines for Control of Water Pollution from Mine Drainage, November 10, 1979; Water Pollution Control Commission (Ch 66, Act. 28, C.R.S. 1963 as amended 1970).
11. Colorado Rules and Regulations Pertaining to Radiation Control, April 1, 1978, Uranium Mill Licensing Guide, May 1978; Radioactive Materials License; Radiation and Hazardous Wastes Control Division (Title 25, Act. II, C.R.S. 1973, Radiation Control).

Table D.1 (continued)--Key to State laws, regulations, and guides cited.

12. Guidelines for the Design, Operation, and Maintenance of Mill Tailings Ponds to Prevent Water Pollution, March 13, 1968; Water Pollution Control Commission (Colorado Water Pollution Control Act of 1966, Ch. 44, Session Laws 1966 as amended by Ch. 217).
13. Publication of a Regulation Providing Tailings Piles from Uranium and Thorium Mills be Adequately Stabilized or Removed, Colorado Department of Public Health; effective June 10, 1966.
14. Colorado Division of Mines responsible for health and safety standards for uranium mines and mills. Regulations contained in Bulletin 20: Section 108 - "Missed Holes--Misfires," Section 110 - "Mucking," Section 12.2 "Radiation Control," Section 130 - "Safeguards," Section 140 - "Shafts and Raises."
15. Office of State Engineer, Division of Water Resources (Article 16, Section 5 - Colorado Constitution and Title 37, Article 90, Section 137 - Colorado Revised Statutes, 1973).

Table D.1 (continued)--Key to State laws, regulations, and guides cited

New Mexico

1. State Land. Leased by State Land Commission, 19-8-14 NMSA 1978.
2. State and Private Land. Mine plan filed and approved by State Mining Inspector, 67-5-1 et seq. NMSA 1978.
3. Water Permit issued by State Engineer; 72-5-1 et seq. NMSA 1978 and 72-12-1 NMSA 1978 and Desert Lands Act of 1866 as amended and 43 USC 383.
4. NRC agreement State Under 42 USC 2021. License required for source material: unrefined and unprocessed ore is not included. Specific License required for Mills, 10 CFR 40.20 - 40.31. Administered by Environmental Improvement Division (EID).
5. New Mexico delegated responsibilities and powers under Clean Air Act (40 CFR 52.1620). Ambient Air Quality Standards and Air Quality Control Regulations, State of New Mexico Health Department, Environmental Improvement Division; reissued November 1976.
6. Application for Permit and Certificate of Registration General Form for Sources Located Within the State of New Mexico, New Source Review Section, Air Quality Section, Environmental Improvement Division, revised February 1976.
7. Application for Permit to Construct or Modify and Certificate of Registration for Mineral Processing Plants Located within the State of New Mexico, New Source Review Section, Air Quality Section, Environmental Improvement Division, revised February 1976.
8. Supplementary Information and Notes for Use with Application for Permit and Certificate of Registration for Mineral Processing Plants, State of New Mexico - Environmental Improvement Division, Air Quality Section, New Source Review Section.
9. Monitoring Air Quality in Mines and Mills Underground: State Mine Inspector 69-5-7 NMSA 1978 also MSHA (30 CFR 57.5-37) Restricted Areas: Mills, EID, 74-2-13 NMSA 1978 Unrestricted Areas: EID per Clean Air Act (42 USC 7410) and State Radiation Protection Act (74-2-1 et seq. NMSA 1978).
10. New Mexico Water Quality: Not NPDES approved by EPA. State does not require permit per 74-6-5 NMSA 1978 and parts 2-100 of N.M. Water Quality Regulations if EPA issues NPDES permit.
11. Underground Water. State EID regulates pollution of underground water per 74-6-1 et seq. NMSA 1978.

Table D.1 (continued)--Key to State laws, regulations, and guides cited

12. Water Quality - Radioactivity: Mines by EID according to Sec. 2-101(b) of N.M. Water Quality Regulations; Mills by EID per NRC 10 CFR 20.106 and Appendix B.
13. Water Quality Standards on Enforcement: EPA enforces under NPDES system for effluent streams entering surface water of United States; EID enforces N.M. groundwater standards under N.M. Water Quality Control Act, 74-6-1 et seq. NMSA 1978.
14. Amended Water Quality Control Commission Regulation, Parts 1,2,3, and 4, Water Quality Control Commission; January 11, 1977, as amended June 14, 1977 and November 8, 1977.
Water Quality Standards for Interstate and Intrastate Streams in New Mexico, Water Quality Control Commission under the authority of Paragraph C, Section 74-6-4 of the New Mexico Water Quality Act (Chapter 326, Laws of 1973, as amended); adopted August 22, 1973, revised September 29, 1975, January 13, 1976, February 8, 1977 and March 14, 1978.
15. New Mexico Environmental Improvement Agency Uranium Mill License Application Guidelines, Radiation Protection Section; September 1977.
16. (a) Radiation Protection Act, Chapter 185 Laws of 1959 (as amended by Chapter 284 Laws of 1971 and by Chapter 343 Laws of 1977).

(b) New Mexico Environmental Improvement Agency Regulations for Governing the Health and Environmental Aspects of Radiation, Environmental Improvement Board, June 16, 1973.

Table D.1 (continued)--Key to State laws, regulations, and guides cited

Texas

1. Texas Uranium Surface Mining and Reclamation Act (May 1978), Rules of the Surface Mining and Reclamation Division. The Railroad Commission of Texas, July 1, 1979.
2. Surface Mining Permit Rule 102 - Elements of Permit Application, Rule 250 Reclamation Plan; Rules of the Surface Mining and Reclamation Division.
3. Application for Permit to Conduct In Situ Uranium Mining, Instructions and Procedural Information for Filing an Application for a Permit to Conduct In Situ Mining of Uranium, Texas Department of Water Resources.
4. Technical Report for In Situ Uranium Mining, Texas Department of Water Resources.
5. Surface Mining Permit, Rule 108 - Permit Approval (Rules of the Surface Mining and Reclamation Division). Permit shall be granted if application complies with Permit rules and all applicable Federal and State laws. Permit may be approved conditioned upon approval of all other required State permits or licenses.
6. Texas Department of Health (TDH) issues licenses for surface mining, in situ mining, milling and processing of uranium ores and leachates in accordance with NRC Agreement.
7. TDH implements U.S. Safe Drinking Water Act regarding public water supplies. The underground injection portion of SDWA is regulated by the Railroad Commission (Oil and Gas), Department of Water Resources (in situ mining of uranium, salt, and sulfur).
8. Texas Regulations for Control of Radiation and Texas Water Quality Standards apply to surface water throughout state.
9. Texas Department of Water Resources issues "no discharge" permits to all uranium in situ extraction processes.
10. Texas Radiation Control Act, 1971. Texas Regulations for Control of Radiation (TDH).
11. Texas Solid Waste Disposal Act, 1969 (Texas Department of Water Resources). Rules pertaining to Industrial Solid Waste Management, March 3, 1978.
12. Texas Air Control Board. Air Control Board H-76 bill introduced February 1, 1979 to include radioactive material in the definition of air contaminant and allow Board to charge fees for permits and variances.

Table D.1 (continued)--Key to State laws, regulations, and guides cited

13. Texas Water Code, Chapter 2 - "Water Use" - Texas Department of Water Resources.
14. Rules and Regulations for Prospecting and Mining State-owned minerals. General Land Office Rules 12.6.18.03.001-.006 (Feb. 17, 1976).
15. Texas Uranium Surface Mining and Reclamation, General Land Office Rules 135.18.05.001-.005.

Table D.1 (continued)--Key to State laws, regulations, and guides cited

Utah

1. Utah Water Pollution Control Act, Utah State Division of Health.
 - (a) Wastewater Disposal Regulations, Part I, Definitions and General Requirements, State of Utah, Department of Social Services, Division of Health; adopted by Utah Water Pollution Control Board, May 18, 1965, Utah State Board of Health, May 19, 1965, (Revised by Utah Water Pollution Control Committee, Nov. 2, 1978) under authority of 26-15-4 to 5 and 73-14-1 to 13, Utah Code annotated, 1953, as amended.
 - (b) Wastewater Disposal Regulations, Part II, Standards of Quality for Waters of the State, State of Utah Department of Social Services, Division of Health; adopted by Utah Water Pollution Control Board May 18, 1965, Utah State Board of Health May 19, 1965, revised by action of the Boards June 2, 1967 and June 21, 1967, further revised by action of the Utah Water Pollution Committee September 13, 1973, and by action of the Utah State Board of Health October 23, 1978.
 - (c) Wastewater Disposal Regulations, Part III, Sewers and Wastewater Treatment works. Consideration of Waste stabilization Ponds (Lagoons) for Industrial Wastes is subject to requirements determined from analysis of the engineers report and other available pertinent information in addition to sections 83-91.
 - (d) Wastewater Disposal Regulations, Part IV, Individual Wastewater Disposal Systems.
 - (e) Wastewater Disposal Regulations, Part V, Small Underground Wastewater Disposal Systems.
2. Changes and Additions to the General Rules and Regulations, adopted by the Board of Oil, Gas and Mining; March 22, 1978, effective June 1, 1978.
 - (a) Rule M-3 -- Notice of Intention to Commence Mining Operations.
 - (b) Rule M-10 -- Reclamation Standards.
3. Water Laws of Utah and Interstate Compacts and Treaties (Second Edition, 1964).
4. State Engineer, H.B. No. 167 - "Temporary Applications to Appropriate Water" - introduced in the 1979 General Session, an act enacting Section 73-3-5.5, Utah Code Annotated 1953.

Table D.1 (continued)--Key to State laws, regulations, and guides cited

5. Utah Radiation Protection Act; Utah Code Annotated, 1953; Title 26, Chapter 25 - Radiation Control.
6. Utah Air Conservation Regulations, State of Utah, Department of Social Services, Division of Health; adopted by the Utah Air Conservation Committee and the Utah State Board of Health September 26, 1971; revised January 23, 1972; July 9, 1975; May 22, 1977; February 1979; under authority of 26-15-5 and 26-24-5 Utah Code annotated, 1953, as amended.

Table D.1 (continued)--Key to State laws, regulations, and guides cited

Washington

1. Mineral Leasing Laws, Revised 1965. (Laws cover surface and underground but not in situ and heap leaching).
2. Rules and Regulations Relating to Protection and Restoration of Lands disturbed through Surface Mining, October 20, 1970 (Surface - Mined Land Reclamation Act, Ch 64, 1970, Sec. 5 RCW 78.44--only applies to surface mining on private and state-owned lands).
3. Rules and Regulations for Radiation Protection, Chapter-402-22 WAC, Specific Licenses.
4. Rules and Regulations for Radiation Protection, Sec. 402-24-220 WAC, Concentrations in Air and Water for Release to Restricted and Unrestricted Areas.
5. Rules and Regulations for Radiation Protection, Chapter 402-24 WAC, Standards For Protection Against Radiation.
6. Rules and Regulations for Radiation Protection, Chapter 402-52 WAC, Uranium and/or Thorium Mill Operation and Stabilization of Mill Tailings Piles.
7. Clean Air Act, Revised Washington Administrative Code, Rev., Chapter 70.94, RCW.
8. Water Quality Standards, State of Washington, Department of Ecology; June 19, 1973. (Revised Dec. 19, 1977). Water Pollution Control Act of 1970 (as amended).
9. Department of Ecology - Water Use -
 - (a) Water Pollution Control: Chapter 90.48 RCW
 - (b) Water Code - 1917 Act: Chapter 90.03 RCW
 - (c) Regulations of Public Groundwaters: Chapter 90.44 RCW.

Table D.1 (continued)--Key to State laws, regulations, and guides cited

Wyoming

1. Regulations and Instructions, Part I, Surface Water, Wyoming State Engineer's Office, revised January 1974.
2. Condensed Detailed Instructions for Preparation of Surface Water Applications and Accompanying Maps for Facilities (pollution control and others) for Mining and Other Industrial Operations, revised 4-28-78. Effluent Limitations and Monitoring Requirements: Wyoming's BPT for Uranium Mine Waters.
3. Wyoming Environmental Quality Act, as amended, Department of Environmental Quality: 1973 Cumulative Supplement, 1974 Session Laws, 1975 Session Laws, 1976 Session Laws, 1977 Session Laws.
 - (a) Article 2 - Air quality Regulations.
 - (b) Article 3 - Water quality.
 - (c) Article 4 - Land Quality. Guidelines No. 1-6 and 8.
 - (d) Article 5 - Solid Waste Management.
4. Wyoming Air Quality Standards and Regulations, Department Environmental Quality, filed January 25, 1979.
5. Water Quality Rules and Regulations, Department of Environmental Quality: Chapter I, Quality Standards for Wyoming Surface Waters, filed July 17, 1979; Chapter II, Discharges/Permit Regulations for Wyoming 1974; Chapter IV, Regulations for Discharge of Oil and Hazardous Substances into Water of the State of Wyoming, June 13, 1978.
6. Proposed Groundwater Regulations: WQD Chapter VIII, Quality Standards for Groundwater of Wyoming (1979); WQD Chapter IX, Wyoming Groundwater Pollution Control Permit (1979).
7. Wyoming Land Quality Rules and Regulations, Department of Environmental Quality, filed October 6, 1978, amended September 13, 1979.
8. State of Wyoming Non-Coal Mining Laws, Safety Rules and Regulations, Title 30--Mines and Minerals.
 - (a) Chapter 1 - General Provisions
 - (b) Chapter 2 - Bureau of Mining Statistics
 - (c) Chapter 3 - Mining Operations Generally (Article 4 - Safety Regulations)
 - (d) Chapter 3 - Mining Operations Generally (Article 5 - Open Cut Land Reclamation)

APPENDIX E
ACTIVE URANIUM MINES IN
THE UNITED STATES

ACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 1

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** ARIZONA *****									
RUTH 1 + 4	SILVER CREEK IND	NAVAJO	2	17 N	23.0 E	14	SURFACE	1,000 - 100,000	150
BLUE ROCK	UOCO	PIMA			0		UNDERGRO	100 - 1,000	50
***** COLORADO *****									
SCHWARTZWALDER M	COTTER CORP	JEFFERSON	25	2 S	71.0 W	06	UNDERGRO	>100,000	850
BLACK MAMA	INCE MNG CO	MESA	20	51 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
BOYANZA	ATLAS MINERALS	MESA	26	51 N	20.0 W	22	UNDERGRO	>100,000	400
CEDAR PT, 3-L CHI	W S DAWSON	MESA	23	51 N	19.0 W	22	UNDERGRO	1,000 - 100,000	200
ELIZARETH 17+10	GULF STATES ENER	MESA	29	50 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
HUBBARD HMSTD PA	HUBBARD MNG.	MESA	35	51 N	20.0 W	22	UNDERGRO	1,000 - 100,000	250
JULY	V. C. MOORES	MESA	12	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
LA SAL 4	PIONEER URAY INC	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 - 100,000	450
LIBERTY BELL	MARION BIRCH	MESA	36	51 N	19.0 W	22	UNDERGRO	1,000 - 100,000	150
LUMSDEN 1	UNION CARBIDE CP	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 - 100,000	100
MINERAL CHAN 10+	ATLAS-AMAX	MESA	7	50 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
PLB-C-G-26A	RALPH FOSTER	MESA			0		UNKNOWN	100 - 1,000	100
PLB-C-G-27	FOSTER H-GOVT LSE	MESA			0		UNDERGRO	>100,000	400
NEW VERDE	UNION CARBIDE CP	MESA	15	50 N	18.0 W	22	UNDERGRO	>100,000	300
OCTOBER ADIT	ATLAS-AMAX	MESA	9	50 N	19.0 W	22	UNDERGRO	1,000 - 100,000	450
PACK RAT 1 + 2	UNION CARBIDE	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 - 100,000	200
RAJAH 30 SHAFT	UNION CARBIDE	MESA	1	50 N	20.0 W	22	UNDERGRO	>100,000	650
RAJAH 67 + 68	UNION CARBIDE CP	MESA	36	51 N	20.0 W	22	UNDERGRO	>100,000	550
ROSEBUD	GRAHAM MNG.	MESA	36	51 N	19.0 W	22	UNDERGRO	100 - 1,000	50
THORNTON	C Y WOODWARD	MESA	31	51 N	19.0 W	22	UNDERGRO	1,000 - 100,000	300
ZEE LSE, -RAJAH 4	UNION CARBIDE	MESA	6	51 N	19.0 W	22	UNDERGRO	>100,000	450
HESSIE 2 + 3	UNION CARBIDE CP	MOFFAT	18	7 N	94.0 W	06	SURFACE	1,000 - 100,000	100
MARGE GROUP	UNION CARBIDE CP	MOFFAT	13	7 N	95.0 W	06	SURFACE	>100,000	300
BAGE-BUELLA	UNION CARBIDE CP	MOFFAT	7	7 N	94.0 W	06	SURFACE	>100,000	50
BROKEN BOW	JOHN DUFUR	MONTEZUMA			0		SURFACE	<100	50
VCA NATURITA TAI	DURITA CORP.	MONTEZUMA			0		TAIL, DYP	100 - 1,000	0
ADAK	CLEGHORN+WASHRUP	MONTROSE	12	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	250
APRIL	ATLAS MINERALS	MONTROSE	10	45 N	18.0 W	22	UNDERGRO	100 - 1,000	50
BLACK POINT	UNION CARBIDE CP	MONTROSE	13	48 N	18.0 W	22	UNDERGRO	100 - 1,000	0
BLACKBURN	UNION CARBIDE	MONTROSE	21	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
BLUE CAP	GEQ-ENERGY RES	MONTROSE	28	28 S	26.0 E	24	UNDERGRO	1,000 - 100,000	200
BOON DOCK	EARL HOTZ	MONTROSE			0		SURFACE	100 - 1,000	0
BREEZY	UNION CARBIDE CP	MONTROSE	25	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
BUCKHORN-UREKA	C + H MINING	MONTROSE	10	48 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
CANON 4, 5 + 7	E M COOPER	MONTROSE	21	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
CLIFFDMELLER	UNION CARBIDE	MONTROSE	24	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	200
COLORADIUM	UNION CARBIDE CP	MONTROSE	21	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
CRIPPLE CREEK 2	UNION CARBIDE CP	MONTROSE	21	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
DONALD L	UNION CARBIDE	MONTROSE	21	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
ECHO 2 + 3	MICHAEL GREAGOR	MONTROSE	3	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
ECHO 6	GLEN GREAGOR	MONTROSE	3	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
EQUINOX	CLEGHORN+WASHRUP	MONTROSE	12	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	350

PAGE 2

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
EULA BELLE CRAIG	UNION CARBIDE	MONTROSE	32	49 N	17.0 W	22	UNDERGRO	>100,000	750
FARMER GIRL	MONOGRAM MNG.	MONTROSE	22	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
FAWN SPRINGS 9	UNION CARBIDE CP	MONTROSE	31	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
GREGOR GROUP	ATLAS-AMAX	MONTROSE	33	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
GREASY SPOON	PIONEER URAY INC	MONTROSE			0		UNDERGRO	1,000 - 100,000	300
GUADALCANAL	UNION CARBIDE CP	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
J M	UNION CARBIDE	MONTROSE			0		UNDERGRO	100 - 1,000	200
JACK KNIFE	WILLIAMS INC	MONTROSE	21	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
LONG PARK 13	UNION CARBIDE CP	MONTROSE	27	4	1.7 W	22	UNDERGRO	1,000 - 100,000	200
LONG PARK 15	UNION CARBIDE	MONTROSE			0		UNDERGRO	1,000 - 100,000	200
LONG PARK 16	UNION CARBIDE CP	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
LUCKY GROUP	ENERGY FUELS NUC	MONTROSE	30	47 N	19.0 W	22	UNDERGRO	100 - 1,000	50
MAYBE 5 & 6	UNION CARBIDE CP	MONTROSE	26	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	400
MILL 2	UNION CARBIDE CP	MONTROSE	32	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
MINERAL JOE GROU	ATLAS MINERALS	MONTROSE	26	46 N	17.0 W	22	UNDERGRO	>100,000	300
MINERAL PARK 4,5	ATLAS MINERALS	MONTROSE	35	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
MLB-C-AH-19	UCC-GOV'T LEASE	MONTROSE			0		UNDERGRO	>100,000	600
MLB-C-BL-23	INCE MNG-GOVT LSE	MONTROSE			0		UNDERGRO	1,000 - 100,000	400
MLB-C-JD-5	GATESFOX-GOVT LSE	MONTROSE			0		UNDERGRO	1,000 - 100,000	250
MLB-C-LP-22	INCE MNG-GOVT LSE	MONTROSE			0		UNDERGRO	1,000 - 100,000	250
MLB-C-SF-11	DAWSON W-GOVT LSE	MONTROSE			0		UNDERGRO	1,000 - 100,000	150
MLB-C-SR-12	SHABAKER-GOVT LSE	MONTROSE			0		UNDERGRO	1,000 - 100,000	250
MLB-C-SR-13	FLANGANP-GOVT LSE	MONTROSE			0		UNDERGRO	1,000 - 100,000	450
MLB-C-SR-16	ALEASE DAWSON	MONTROSE			0		UNDERGRO	1,000 - 100,000	100
MLB-C-SR-16A	DYNOVE LTD	MONTROSE			0		UNDERGRO	1,000 - 100,000	100
MONOGRAM CLAIM	UNION CARBIDE	MONTROSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
NIL-TRACE	UNION CARBIDE CP	MONTROSE			0		UNDERGRO	>100,000	400
PEANUT MINES	ATLAS-AMAX	MONTROSE		45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
PEGGY	NEESHAM, GLEN	MONTROSE	6	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
PICKET CORPAL	DON ANDREWS	MONTROSE	34	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
PRINCESS	D K ANDREWS	MONTROSE	27	48 N	18.0 W	22	UNDERGRO	100 - 1,000	150
RAVEN	FOOTE MINERALS	MONTROSE			0		UNDERGRO	1,000 - 100,000	100
REX MINE	CLEGHORN & WAB	MONTROSE	3	47 N	7.0 W	22	UNDERGRO	1,000 - 100,000	400
RIMROCK 5	UNION CARBIDE CP	MONTROSE			0		UNDERGRO	1,000 - 100,000	150
RIMROCK BLUES 2	UNION CARBIDE CP	MONTROSE	2	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
RIMROCK GROUP	UNION CARBIDE CP	MONTROSE	16	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
RYE 8	NATIVE RESERVE	MONTROSE			0		UNDERGRO	1,000 - 100,000	50
SEPTEMBER MORN	UNION CARBIDE	MONTROSE	10	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
SEMO	ATLAS MINERALS	MONTROSE	24	47 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
SILVER DOLLAR	C & D EXPLORATIO	MONTROSE			0		UNDERGRO	<100	100
ST. PATRICK 9	C W BUNKER	MONTROSE	11	47 N	20.0 W	22	UNDERGRO	1,000 - 100,000	200
SUNBEAM GROUP	UNION CARBIDE	MONTROSE	10	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	280
URA	UNION CARBIDE C	MONTROSE	29	46 N	17.0 W	22	UNDERGRO	>100,000	450
WANDA 3	PATTERSON, JAMES	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	100 - 1,000	50
WHITE FACE	UNION CARBIDE CP	MONTROSE	34	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
YELLOW BIRD 1	R K DIETZ	MONTROSE	33	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
YELLOW SPOT GROU	REED MINING	MONTROSE			0		UNKNOWN	<100	0
PITCH	HOMESTAKE MNG CO	SAGUACHE	48 N		6.0 E	22	UNDERGRO	>100,000	250

ACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 3

MINING METHOD	MINES NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
UNDERGRO	BURRO	UNION CARBIDE	SAN MIGUEL	31	44 N	18.0 W	22	>100,000	500
UNDERGRO	CARNATION	PIONEER URAY INC	SAN MIGUEL	13	44 N	18.0 W	22	>100,000	200
UNDERGRO	CIVET CAT GROUP	EAGLE PEAK MNG C	SAN MIGUEL			0		1,000 - 100,000	50
UNDERGRO	DEREMO-BIGLER SH	UNION CARBIDE	SAN MIGUEL	2	42 N	20.0 W	22	>100,000	650
UNDERGRO	FALCON-BOB-DYLA	ATLAS-FOOTE	SAN MIGUEL			0		1,000 - 100,000	100
UNDERGRO	GMC	UNION CARBIDE CO	SAN MIGUEL	18	44 N	17.0 W	22	1,000 - 100,000	500
UNDERGRO	HANGOVER (SLRCK)	LORELL E ROCKWEL	SAN MIGUEL	21	43 N	19.0 W	22	1,000 - 100,000	150
UNDERGRO	HAPPY JACK (SLKPC)	MARION BIPCH	SAN MIGUEL	36	44 N	20.0 W	22	1,000 - 100,000	0
UNDERGRO	LUCKY STRIKE	HANNERT MNG.	SAN MIGUEL	19	43 N	19.0 W	22	1,000 - 100,000	150
UNDERGRO	MAGPIE 2	PIONEER URAY INC	SAN MIGUEL	25	45 N	18.0 W	22	1,000 - 100,000	250
UNDERGRO	MURIELTA	SHIPROCK, LTD	SAN MIGUEL	25	45 N	18.0 W	22	1,000 - 100,000	250
UNDERGRO	NORTH BURRO	UNION CARBIDE CP	SAN MIGUEL			0		1,000 - 100,000	300
UNDERGRO	POND	UNION CARBIDE CP	SAN MIGUEL	23	44 N	17.0 W	22	1,000 - 100,000	100
UNDERGRO	RADIUM 9,10,+ 11	ATLAS-FOOTE	SAN MIGUEL	4	43 N	19.0 W	22	1,000 - 100,000	100
UNDERGRO	RADIUM CP-BLACKB	ATLAS-FOOTE	SAN MIGUEL	32	44 N	19.0 W	22	1,000 - 100,000	0
UNDERGRO	SEARS GROUP	JOHN PERES	SAN MIGUEL			0		100 - 1,000	0
UNKNOWN	SILVER BELL	UNION CARBIDE CP	SAN MIGUEL	22	32 S	26.0 E	24	1,000 - 100,000	650
UNDERGRO	SNYDER + PETERSO	UNION CARBIDE	SAN MIGUEL	18	43 N	18.0 W	22	>100,000	750
UNDERGRO	STRAWBERRY ROAD	ATLAS MINERALS	SAN MIGUEL	32	43 N	19.0 W	22	1,000 - 100,000	100
UNDERGRO	SUMMIT INCLINE 1	ATLAS-APAX	SAN MIGUEL	28	43 N	19.0 W	22	1,000 - 100,000	300
UNDERGRO	SUN CUP (PUCKETT)	DOLORES BENCH LT	SAN MIGUEL	31	4 N	1.7 W	22	1,000 - 100,000	300
UNDERGRO	SUNDAY GROUP	UNION CARBIDE CP	SAN MIGUEL	13	44 N	18.0 W	22	>100,000	300
UNDERGRO	UNTAN	W. D. TRIPP	SAN MIGUEL	28	43 N	19.0 W	22	1,000 - 100,000	250
UNDERGRO	WILKARTH	UNION CARBIDE CP	SAN MIGUEL	24	44 N	17.0 W	22	1,000 - 100,000	0
UNDERGRO	WINDSHEPT	UNION CARBIDE CP	SAN MIGUEL	15	44 N	17.0 W	22	1,000 - 100,000	100
***** NEW MEXICO *****									
UNDERGRO	ANN LEE (2A,14-9)	UNITED NUCLEAR	MCKINLEY	28	14 N	9.0 W	22	>100,000	650
UNDERGRO	BUCKY (14-14-10)	COBB NUCLEAR INC	MCKINLEY	14	14 N	10.0 W	22	>100,000	200
UNDERGRO	CLIFFSIDE 36 14N	KERR-MCGEE CORP	MCKINLEY	36	14 N	9.0 W	22	>100,000	1400
UNDERGRO	DOG-FLEA (20-13-	M + M MINING CO	MCKINLEY	20	13 N	9.0 W	22	>100,000	200
HWATPROD	IX CIRCUIT GRANT	KERR MCGEE	MCKINLEY			0		<100	0
UNDERGRO	MAC 1 12,13-14,3	UN - HOMESTAKE	MCKINLEY	12	15 N	14.0 W	22	1,000 - 100,000	400
HWATPROD	MAINE WATER	UNITED NUC HOM	MCKINLEY			0		<100	0
SURFACE	MLB-NH-B-1	WARNOCK-GOVT LSE	MCKINLEY			0		>100,000	50
UNDERGRO	N E CHURCHROCK	UNITED NUCLEAR	MCKINLEY	34	17 N	16.0 W	22	>100,000	1600
UNDERGRO	NAVAJO RES. NW	KERR-MCGEE CORP.	MCKINLEY	34	17 N	17.0 W	22	1,000 - 100,000	2250
UNDERGRO	SANDSTONE(27-14-	UNITED NUCLEAR	MCKINLEY	34	13 N	9.0 W	22	>100,000	750
UNDERGRO	SEC 1 13 9	KERR MCGEE	MCKINLEY	1	13 N	9.0 W	22	>100,000	1400
UNDERGRO	SEC 12 14 10 BWO	COBB NUCL IND	MCKINLEY	12	14 N	10.0 W	22	>100,000	650
UNDERGRO	SEC 12-13-14W2,N	GULF MINERAL RES	MCKINLEY	12	15 N	14.0 W	22	1,000 - 100,000	450
UNDERGRO	SEC 18+20-14-9	KERR MCGEE	MCKINLEY	20	14 N	9.0 W	22	>100,000	800
UNDERGRO	SEC 19 14W 9W	KERR-MCGEE CORP.	MCKINLEY	19	14 N	9.0 W	22	>100,000	450
UNDERGRO	SEC 19-13-09(HOP	RANCHERS EXPL.	MCKINLEY	19	13 N	9.0 W	22	1,000 - 100,000	500
UNDERGRO	SEC 21-13N-13W	WESTERN NUCLEAR	MCKINLEY	21	13 N	13.0 W	22	>100,000	400
UNDERGRO	SEC 25-14W-10W	UNITED NUC HOM	MCKINLEY	25	14 N	10.0 W	22	>100,000	700

ACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 4

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** NEW MEXICO (CONT'D) *****									
SEC 30, 14-9 (E +	KERR MCGEE	MCKINLEY	30	14 N	9,0 W	22	UNDERGRO	>100,000	550
SEC 32 15N 11W	COBB NUCLEAR INC	MCKINLEY	32	15 N	11,0 W	22	UNDERGRO	1,000 - 100,000	250
SEC 35, 14N-9W	KERR MCGEE	MCKINLEY	35	14 N	9,0 W	22	UNDERGRO	>100,000	1200
SEC 35, 17N-16W N	KERR MCGEE CORP.	MCKINLEY	34	17 N	16,0 W	22	UNDERGRO	1,000 - 100,000	1650
SEC 7-13N-8W	RANCHERS EXP, DEV	MCKINLEY	7	13 N	8,0 W	22	UNDERGRO	>100,000	1350
SEC 8 13N 9W	KOPPEN	MCKINLEY	8	13 N	9,0 W	22	UNDERGRO	1,000 - 100,000	300
SEC. 13-13-11 5	TODILTO EXPL, +D	MCKINLEY	13	13 N	11,0 W	22	UNDERGRO	1,000 - 100,000	50
SEC. 15-14-10	UNITED NUCLEAR	MCKINLEY	15	14 N	10,0 W	22	UNDERGRO	>100,000	500
SEC. 13-14N-10W	UNIT, NUC, MHSTKE	MCKINLEY	13	14 N	10,0 W	22	UNDERGRO	>100,000	500
SEC. 17, 14N-9W, 82	KERR MCGEE	MCKINLEY	17	14 N	9,0 W	22	UNDERGRO	>100,000	850
SEC. 19-13-10 NWQ	TODILTO EXPL + D	MCKINLEY	19	13 N	10,0 W	22	SURFACE	>100,000	50
SEC. 19-13-9 SEQ	RESERVE OIL + NMR	MCKINLEY	19	13 N	9,0 W	22	UNDERGRO	>100,000	200
SEC. 21, 13-9 DQPI	RANCHERS EXPL.	MCKINLEY	21	13 N	9,0 W	22	UNDERGRO	1,000 - 100,000	250
SEC. 23-14N-10W	UNKNOWN	MCKINLEY	23	14 N	10,0 W	22	UNDERGRO	>100,000	700
SEC. 24-26-14N-10	KERR-MCGEE CORP.	MCKINLEY	24	14 N	10,0 W	22	UNDERGRO	>100,000	650
SEC. 26, 14N-9W, 82	KERR-MCGEE CORP.	MCKINLEY	26	14 N	9,0 W	22	UNDERGRO	1,000 - 100,000	1150
SEC. 29-14-9 (MU)	KERR-MCGEE CORP.	MCKINLEY	29	14 N	9,0 W	22	UNDERGRO	>100,000	600
SEC. 32, 14N-09W	UNITED NUCLEAR	MCKINLEY	32	14 N	9,0 W	22	UNDERGRO	>100,000	600
ENDS JOHNSON DUM	RAY WILLIAMS	SAN JUAN			0		DUMPS	1,000 - 100,000	0
L-BAR	BOHIO-RESERVE	VALENCIA	24	11 N	5,0 W	22	UNDERGRO	>100,000	600
PAGUATE-JACKPIL	ANACONDA CO.	VALENCIA		11 N	5,0 W		SURFACE	>100,000	150
PAGUATE-JACKPIL	ANACONDA CO	VALENCIA			0		UNDERGRO	>100,000	450
ST. ANTHONY MINE	UNITED NUCLEAR	VALENCIA	19	11 N	4,0 W	22	SURFACE	>100,000	250
***** TEXAS *****									
HURT LEASE	CONOCO PIONEER	ATASCOSA			0		SURFACE	>100,000	100
BOONE-ANDERSON L	INTRCNTAL FNERGY	BEE			0		IN-SITU	1,000 - 100,000	250
BENAVIDES	WYO MINERAL CORP	DUVAL			0		IN-SITU	1,000 - 100,000	200
PALANGANA DOME	UNION CARBIDE CP	DUVAL			0		IN-SITU	<100	300
REICKER-ZAMZON	CONOCO PIONEER	KARNES			0		SURFACE	1,000 - 100,000	100
BUTLER LEASE	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	100
CARMODY-ROSEN17	CONOCO- PIONEER	KARNES			0		SURFACE	1,000 - 100,000	300
DICKSON (UL 764)	CONOCO PIONEER	KARNES			0		SURFACE	1,000 - 100,000	200
FALLS CITY TAIL	SOLUTION ENGINRG	KARNES			0		TAIL, DMP	>100,000	0
FRANKLIN (UL1754	CONOCO PIONEER	KARNES			0		SURFACE	1,000 - 100,000	200
KOTZUR-2260	CONOCO-PIONEER	KARNES		09	17,0		SURFACE	>100,000	100
PAWELEK D.B. 7	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	200
PAWELEK-LYSSA 7	CONOCO- PIONEER	KARNES			0		SURFACE	>100,000	100
THOMAS-KNANDEL-D	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	150
WINERICH C.D. 4	CONOCO- PIONEER	KARNES			0		SURFACE	>100,000	100
BURNS MNG, PROSPE	U. S. STEEL	LIVE OAK			0		IN-SITU	>100,000	500
CHAPMAN	CONOCO-PIONEER	LIVE OAK			0		SURFACE	>100,000	100
CLAY WEST PROPER	U. S. STEEL	LIVE OAK			0		IN-SITU	>100,000	350
FELDER TRACT	EXXON CO., USA	LIVE OAK			0		SURFACE	>100,000	100
HOUSE-1734	CONOCO-PIONEER	LIVE OAK		11	15,0		SURFACE	>100,000	200
LAMPRECHT, A.H.-2	WYOMING MINERALS	LIVE OAK			0		IN-SITU	>100,000	250

ACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 5

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** TEXAS (CONT'D) *****									
MC LEAN-BOWMAN	EXXON CO., USA	LIVE OAK			0		SURFACE	1,000 - 100,000	100
SMITH, W.A. 2-670	CONOCO-PIONEER	LIVE OAK			0		SURFACE	1,000 - 100,000	100
ZAMZOW	INTRNTNL ENERGY	LIVE OAK			0		IN-SITU	>100,000	150
O'HERN LEASE-75	MOBIL OIL CO	WEBB			0		IN-SITU	>100,000	500
***** UTAH *****									
MONARCH	HOLLINGSHEAD MNG	BEAVER	36	29 S	16,0 W	24	UNDERGRO	1,000 - 100,000	100
ALLEN	WATTERSON MINING	EMERY			0		UNDERGRO	1,000 - 100,000	100
COMETOITE	INDUSTRIAL MININ	EMERY			0		UNDERGRO	1,000 - 100,000	50
DEDE 6,7+8	ATLAS MINERALS	EMERY			0		UNDERGRO	1,000 - 100,000	500
DELTA MINE	UTAH WEST INC.	EMERY			0		UNDERGRO	>100,000	400
DESEPT MOON 3	WATTERSON MINING	EMERY			0		UNDERGRO	1,000 - 100,000	50
DEXTER GROUP	JOHN ADAMS	EMERY			0		UNDERGRO	1,000 - 100,000	100
DIRTY DEVIL 2	INDUSTRIAL NUCL	EMERY			0		UNDERGRO	1,000 - 100,000	100
DOG 3	JOHN ADAMS	EMERY			0		SURFACE	1,000 - 100,000	50
FLAT TOP LODGE	MINERALS FVAL+IN	EMERY			0		SURFACE	1,000 - 100,000	50
INCLINE 1,2,+7	ATLAS MINERALS	EMERY			0		UNDERGRO	>100,000	200
INCLINE 6	ATLAS MINERALS	EMERY			0		UNDERGRO	>100,000	200
JACK RAB-ACE-N-H	ATLAS MINERALS	EMERY			0		UNDERGRO	1,000 - 100,000	500
JOSHUA 1	TROJAN MINING CO	EMERY			0		UNDERGRO	<100	0
TEMPLE MOUNTAIN	RON CROSS	EMERY			0		UNDERGRO	1,000 - 100,000	0
THUNDERBIRD	INDUSTRIAL MININ	EMERY			0		UNDERGRO	1,000 - 100,000	50
VANADIUM KING 1	TEMPLE ROCK MNG.	EMERY			0		UNDERGRO	1,000 - 100,000	100
YELLOW DAISY	DAN L POWELL	EMERY			0		UNDERGRO	100 - 1,000	200
YELLOW QUEEN	AUGUST PRIEBE, JR	EMERY			0		UNDERGRO	1,000 - 100,000	50
CONGRESS 29+40	TED EKKER	GARFIELD			0		SURFACE	1,000 - 100,000	100
CONGRESS-DAISY J	ENERGY FUELS NUC	GARFIELD			0		UNDERGRO	100 - 1,000	100
CONGRESS-EAGLE	DONALD D HANNI	GARFIELD			0		UNDERGRO	1,000 - 100,000	50
DAISY JUNE GROUP	TED EKKER	GARFIELD			0		UNDERGRO	1,000 - 100,000	50
DELMONTE GROUP	DENNIS EKKER	GARFIELD			0		UNDERGRO	1,000 - 100,000	50
DONALD DUCK 2	DEE RAEY	GARFIELD			0		UNDERGRO	100 - 1,000	50
ELENORA 1	ENERGY FUELS NUC	GARFIELD			0		SURFACE	<100	100
LUCKY STRIKE	PLATEAU RESOU LT	GARFIELD			0		UNDERGRO	100 - 1,000	0
MIDAS-CENTIPEDE	SHUMWAY ASSOC.	GARFIELD			0		UNDERGRO	1,000 - 100,000	150
MINNIE PEARL	LEO D JACKSON	GARFIELD			0		UNDERGRO	1,000 - 100,000	50
MOCKINGBIRD CLAI	J + R MNG.	GARFIELD			0		UNDERGRO	1,000 - 100,000	100
P+P	J + R MNG.	GARFIELD			0		UNDERGRO	1,000 - 100,000	0
POISON SPRING 2	HUNT, A + SONS	GARFIELD			0		SURFACE	<100	50
SEC 36-31-11 BLM	MINERALS RECVRY	GARFIELD			0		UNDERGRO	1,000 - 100,000	50
TRACKYTE GROUP	ENERGY FUELS NUC	GARFIELD			0		UNDERGRO	1,000 - 100,000	100
BLACK JACK	LYNN ZUFELT	GRAND	34	22 S	21,0 E	24	UNDERGRO	100 - 1,000	50
BLACKSTONE 5 + 6	ALFRED FROST	GRAND	31	22 S	22,0 E	24	UNDERGRO	1,000 - 100,000	150
CACTUS RAT	ATLAS-AMAX	GRAND	33	22 S	22,0 E	24	UNDERGRO	1,000 - 100,000	50
COBALT	L + D SHUMWAY	GRAND			0		SURFACE	<100	50
CORRAL 1 + 2	COTTER CORP.	GRAND	27	24 S	20,0 E	24	UNDERGRO	>100,000	150
CORVUSITE	SHIPROCK LTD	GRAND	19	25 S	26,0 E	24	UNDERGRO	1,000 - 100,000	150

ACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 6

MINES NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
EAST DOLLAR	JOE GOMEZ	GRAND			0		UNDERGRO	100 - 1,000	0
MC GROUP	PIONEER URANUM	GRAND			0		UNDERGRO	<100	500
MINERAL POLAR 22	ATLAS-AMAX	GRAND	26	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	50
PINTO	ZIAK, PAUL	GRAND	33	27 S	20.0 E	24	SURFACE	<100	0
POLAR GROUP	UNION CARBIDE CP	GRAND	10	25 S	25.0 E	24	UNDERGRO	1,000 - 100,000	100
POLAR MESA	UNION CARBIDE CP	GRAND		25 S	25.0 E	24	UNDERGRO	1,000 - 100,000	0
RED HEAD	JERRY STOCKS	GRAND	22	24 S	23.0 F	24	UNDERGRO	100 - 1,000	150
SEC. 32-CANE CRFE	ATLAS-AMAX	GRAND	32	26 S	21.0 E	24	UNDERGRO	1,000 - 100,000	150
SHINARUMP 2	GENERAL ELECTRIC	GRAND	34	24 S	20.0 E	24	UNDERGRO	1,000 - 100,000	50
SLICK ROCK 1,2+3	JIM C BUTT	GRAND	28	23 S	20.0 E	24	UNDERGRO	100 - 1,000	0
UTAH ALLOY GROUP	SILLMAN, KENNETH	GRAND	6	23 S	22.0 E	24	UNDERGRO	1,000 - 100,000	50
BINGHAM	WYOMING MARL CP	SALT LAKE			0		SURFACE	1,000 - 100,000	0
ARE 4	RICHARD V ENSLE	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
ATOMIC KING GROU	URANIUM PROD CO	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
BEAVER (DEER CK)	UNION CARBIDE	SAN JUAN	31	28 S	25.0 E	24	UNKNOWN	1,000 - 100,000	450
BIG BUCK 11A + 1	BOB SHUMWAY	SAN JUAN	14	30 S	24.0 E	24	UNDERGRO	>100,000	400
BLUE 1	EAST STANDARD UR	SAN JUAN	14	33 S	21.0 E	24	UNDERGRO	1,000 - 100,000	150
BLUE BIRD	DALE DILLON	SAN JUAN	4	37 S	21.0 F	24	UNDERGRO	1,000 - 100,000	150
RUGS 1-4	MERWIN SHUMWAY	SAN JUAN	4	37 S	21.0 E	24	UNDERGRO	1,000 - 100,000	100
CAMEL (HIDEOUT)	JIM C. BUTT	SAN JUAN			0		UNDERGRO	100 - 1,000	150
CANARY MINE	JEPPEY STOCKS	SAN JUAN			0		UNDERGRO	100 - 1,000	50
CLIFF HOUSE	GLEN J SHUMWAY	SAN JUAN	4	37 S	21.0 E	24	UNDERGRO	1,000 - 100,000	50
COLUMBIA SHAFT G	ATLAS MINERALS	SAN JUAN	27	29 S	24.0 E	24	UNDERGRO	>100,000	500
COLUMBUS-RIM	ATLAS-AMAX	SAN JUAN			0		UNDERGRO	>100,000	400
COTTONWOOD GROUP	DAN SHUMWAY	SAN JUAN			0		UNDERGRO	100 - 1,000	100
COTTONWOOD GROUP	MERWIN SHUMWAY	SAN JUAN	4	37 S	21.0 F	24	UNDERGRO	1,000 - 100,000	50
DUNTY	CLYDE R. SANCHEZ	SAN JUAN	26	36 S	25.0 E	24	UNDERGRO	1,000 - 100,000	0
EASTER	GLEN J SHUMWAY	SAN JUAN			0		UNDERGRO	100 - 1,000	0
FAR WEST MINE	ATLAS MINERALS	SAN JUAN	28	29 S	24.0 E	24	UNDERGRO	>100,000	550
FRY 4	JIM C. BUTT	SAN JUAN		37 S	16.0 E	24	UNDERGRO	1,000 - 100,000	150
GENEVA	CLAYTON STOCKS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
GLADE GROUP	ENERGY FUELS NUC	SAN JUAN			0		SURFACE	1,000 - 100,000	100
HAPPY JACK (HCA)	ATLAS MINERALS	SAN JUAN	8	35 S	15.0 E	24	UNDERGRO	>100,000	200
HILLSIDE	ENERGY FUELS NUC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
HOLE IN THE ROCK	BLACK + SHUMWAY	SAN JUAN	16	40 S	22.0 E	24	UNDERGRO	1,000 - 100,000	50
HUMBURG	ATLAS-AMAX	SAN JUAN	30	31 S	25.0 E	24	UNDERGRO	1,000 - 100,000	100
JEAN 1	JIM C BUTT	SAN JUAN	15	31 S	21.0 F	24	UNDERGRO	1,000 - 100,000	50
JIMBO BOB	SHUMWAY ASSOC.	SAN JUAN	6	32 S	25.0 E	24	UNDERGRO	1,000 - 100,000	250
KING EDWARD	EUGENE SHUMWAY	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
LAST CHANCE	W. R. WILLIAMS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	50
LAURA GROUP	LAURA MNG. CO.	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
LEF GROUP	ENERGY FUELS NUC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	50
LISBON MINE	RIO ALGOM CORP	SAN JUAN	27	29 S	24.0 E	24	UNDERGRO	>100,000	2600
LOCUST	LIGHTON-GRANT	SAN JUAN	4	3 S	2.5 E	24	UNDERGRO	100 - 1,000	100
LONG SHOT	DAN SHUMWAY	SAN JUAN			0		SURFACE	100 - 1,000	0
MARKEY	CALVIN BLACK EN	SAN JUAN	19	37 S	15.0 E	24	UNDERGRO	>100,000	1000
MI VIDA	MINERALS WEST	SAN JUAN	11	30 S	24.0 E	24	UNDERGRO	>100,000	50
NORTH ALICE	JRS MINING CO	SAN JUAN	21	29 S	24.0 E	24	UNDERGRO	>100,000	400

ACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH *****									
NOTCH 1 + 4	JUNCTION BIT-TOO	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
OLD RATTLER	ENERGY FUELS NUC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
PANDORA	ATLAS-AMAX	SAN JUAN	5	29 S	25.0 E	24	UNDERGRO	>100,000	450
PATS PROPERTY	M + M MINING CO	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
POSEY GROUP	ENERGY FUELS NUC	SAN JUAN		37 S	15.0 E	24	UNDERGRO	1,000 - 100,000	200
PRETTY GIRL	DERYL SHUMWAY	SAN JUAN			0		UNKNOWN	100 - 1,000	0
RADIUM KING MINE	LEECO GAS + OIL	SAN JUAN			0		UNDERGRO	1,000 - 100,000	500
RAINBOW 41	DONALD V BLAKE	SAN JUAN			0		SURFACE	100 - 1,000	50
RAINBOW LEDGE	JIM C BUTT	SAN JUAN			0		UNDERGRO	100 - 1,000	0
RED + BLACK LIZA	LOWELL E ROCKWEL	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
RED APRIL	AMPLS WEST INC	SAN JUAN			0		SURFACE	100 - 1,000	50
RED DEVIL	SUNRAY MINING CO	SAN JUAN	21	27 S	24.0 E	24	UNDERGRO	100 - 1,000	0
RYANORE	CLAYTON STOKES	SAN JUAN	5	31 S	25.0 E	24	UNDERGRO	1,000 - 100,000	50
SAGE	ATLAS MINERALS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	700
BALLY JO 1 + 2	DERYL SHUMWAY	SAN JUAN	15	37 S	21.0 E	24	UNDERGRO	1,000 - 100,000	50
SAN JUAN	MORARD BALSLEY	SAN JUAN	4	29 S	23.0 E	24	UNDERGRO	1,000 - 100,000	150
SANDY GROUP	RICHARD KIPP	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
SEC SEC 2 CHINLE	UNION CARRIDE	SAN JUAN	2	30 S	24.0 E	24	UNDERGRO	>100,000	700
SEC 2 + 31 REDD	ATLAS MINERALS	SAN JUAN	02	29 S	24.0 E	24	UNDERGRO	>100,000	450
SEC 28 CALLINAN	BIRCH + STOKES	SAN JUAN			0		UNDERGRO	1,000 - 100,000	700
SEC-2-28S-22F	BLAKE MCG CO	SAN JUAN			0		UNDERGRO	1,000 - 100,000	0
SHOKEY	LYLE DAVIS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
SNOW FLAKE	UNION CARRIDE CP	SAN JUAN	28	27 S	23.0 E	24	UNDERGRO	1,000 - 100,000	0
SNOWBALL	DELA SHUMWAY	SAN JUAN	36	29 S	24.0 E	24	UNDERGRO	1,000 - 100,000	450
STRAMBERY	GLEN J SHUMWAY	SAN JUAN	35	35 S	24.0 E	24	UNDERGRO	1,000 - 100,000	100
V-8 AND GOLD BUT	ROBERT YOUNG	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
VANADIUM QUEEN]-	MURPHY MNG CO	SAN JUAN	29	28 S	24.0 E	24	UNDERGRO	1,000 - 100,000	150
WATERFALL	JIM C BUTT	SAN JUAN	6	31 S	25.0 E	24	UNDERGRO	1,000 - 100,000	0
WEE HOPF	UNION CARRIDE CO	SAN JUAN	25	36 S	16.0 E	24	UNDERGRO	1,000 - 100,000	250
WHITE CAN, 1-CAN	ATLAS MINERALS	SAN JUAN	15	32 S	24.0 E	24	UNDERGRO	1,000 - 100,000	150
WILSON LEASE	EAGLE PEAK MINIM	SAN JUAN	6	31 S	25.0 E	24	UNDERGRO	>100,000	650
WOOD LEASE	ENERGY FUELS NUC	SAN JUAN			0		UNKNOWN	100 - 1,000	700
WOODSHOE 1+2	REX POWELL	SAN JUAN			0		UNDERGRO	1,000 - 100,000	0
YANKEE GIRL	LANHART + SHUMWAY	SAN JUAN	12	36 S	24.0 E	24	UNDERGRO	1,000 - 100,000	150
YELLOW CAKE		SAN JUAN	34	27 S	23.0 E	24	SURFACE	<100	250
YELLOW CIRCLE GR		SAN JUAN					UNDERGRO	1,000 - 100,000	100
***** WASHINGTON *****									
MIDNITE-ROYD	DAWN MNG CO	STEVENSON	6	28 N	38.0 E	33	SURFACE	>100,000	100
PETERS LBF-SHRWO	WESTERN NUCLEAR	STEVENSON	35	28 N	37.0 E	33	SURFACE	>100,000	250
***** WYOMING *****									
F GRP SEC28 ARE	PATHFINDER	CARBON	2A	28 N	78.0 W	06	SURFACE	>100,000	350
JENKINS STATE LE	URAN.SUPPLY BRVC	CARBON	11	27 N	78.0 W	06	SURFACE	1,000 - 100,000	250

ACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 8

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** WYOMING (CONT'D) *****									
SEC.33+4,28N-78W	GETTY OIL	CARBON	3	27 N	78.0 W	05	SURFACE	>100,000	350
BEAR CREEK B-4,5	ROCKY MT ENERGY	CONVERSE	35	38 N	73.0 W	6	SURFACE	>100,000	150
HIGHLAND PROJ O.	EXXON CO USA	CONVERSE	29	36 N	73.0 W	06	SURFACE	>100,000	300
HIGHLAND PROJ.U.	EXXON CO., USA	CONVERSE	20	36 N	73.0 W	06	UNDERGRO	>100,000	550
SEC.33,37N-73W	KERR-MCGEE CORP.	CONVERSE	33	37 N	73.0 W	06	SURFACE	1,000 - 100,000	200
SEC.36,36N-74W	KERR-MCGEE CORP.	CONVERSE	36	36 N	74.0 W	06	UNDERGRO	1,000 - 100,000	650
BIG EAGLE	PATHFINDER MINES	FREMONT	27	27	2.9		SURFACE	>100,000	300
CAP 1,2+3	FEDERAL AMERICAN	FREMONT	27	33 N	89.0 W	6	SURFACE	100 - 1,000	200
CLYDE, BPET, LOC	FEDERAL AMERICA	FREMONT	23	32 N	91.0 W	06	SURFACE	>100,000	200
DICK GROUP	UNION CARBIDE CP	FREMONT	30	33 N	90.0 W	06	SURFACE	>100,000	250
GOLDEN GOOSE	WESTERN NUCLEAR	FREMONT	21	28 N	92.0 W	06	SURFACE	>100,000	350
HAL-BART-EGL GRI	PATHFINDER	FREMONT	6	32 N	90.0 W	06	SURFACE	>100,000	200
LUCKY MC GROUP	PATHFINDER	FREMONT	2	33 N	90.0 W	6	SURFACE	>100,000	300
MC INTOSH E+W 8N	WESTERN NUCLEAR	FREMONT			0		SURFACE	>100,000	200
OLA GROUP	UNION CARBIDE CP	FREMONT	32	33 N	90.0 W	06	SURFACE	>100,000	250
PAY DIRT	WESTERN NUCLEAR	FREMONT	20	28 N	92.0 W	06	UNDERGRO	>100,000	150
RAVINE	WESTERN NUCLEAR	FREMONT			0		IN-SITU	1,000 - 100,000	0
SEC 16 MINE WATE	WESTERN NUCLEAR	FREMONT			0		IN-SITU	<100	0
SEISMIC RES. SEC	WESTERN NUCLEAR	FREMONT	17	28 N	92.0 W	06	UNDERGRO	>100,000	350
SHEEP MOUNTAIN 1	WESTERN NUCLEAR	FREMONT	22	28	92.0	22	UNDERGRO	>100,000	1550
SUNSET GROUP	FEDERAL AMERICA	FREMONT	6	32 N	90.0 W	6	SURFACE	>100,000	200
IRIGARAY	WYOMING MINERALS	JOHNSON	18	46	77.0	6	IN-SITU	1,000 - 100,000	250
PAY ALJOB-HEAP L	UNION CARBIDE	NATRONA			0		HL-DUMPS	>100,000	0
PAY ALJOB-PROTOR	UNION CARBIDE	NATRONA			0		LOWGRADE	1,000 - 100,000	0
STAR 1-7	UNION CARBIDE CP	NATRONA	14	33 N	89.0 W	06	SURFACE	>100,000	50
SUSIE	UNION CARBIDE CP	NATRONA	14	33 N	89.0 W	06	UNDERGRO	1,000 - 100,000	150
ARE/CRE(SWEETWAT	MINERALS EXPLTN	SWEETWATER		24 N	93.0 W	06	SURFACE	<100	200
CRE	MINERALS EXPL.CO	SWEETWATER	16	24 N	93.0 W	06	SURFACE	<100	100

APPENDIX F
INACTIVE URANIUM MINES IN
THE UNITED STATES

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 1

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** ALASKA *****		
CUB (ALASKA)	STANDARD METALS	SOUTHEAST
0	UNDERGRO	1,000 - 100,000
***** ARIZONA *****		
APACHE MINE	NAVAJO TRIBE	APACHE
ARROWHEAD 1	NAVAJO TRIBE	APACHE
BARTON 3	NAVAJO TRIBE	APACHE
BETTIE 1	NAVAJO TRIBE	APACHE
BLACK 1	NAVAJO TRIBE	APACHE
BLACK 2	NAVAJO TRIBE	APACHE
BLACK MUSTACHE	NAVAJO TRIBE	APACHE
BLACK ROCK	NAVAJO TRIBE	APACHE
BLACK ROCK POINT	NAVAJO TRIBE	APACHE
BLUCK K MIKE BRO	NAVAJO TRIBE	APACHE
BLUESTONE 1	NAVAJO TRIBE	APACHE
CAPTAN BENALLY	NAVAJO TRIBE	APACHE
CARSON	NAVAJO TRIBE	APACHE
CATO 1	NAVAJO TRIBE	APACHE
CATO 2	NAVAJO TRIBE	APACHE
CHEF MEZ 1	NAVAJO TRIBE	APACHE
CHESTER GROUP	CODDLEY, CFCIG J.	APACHE
CHESTER MINE 1	NAVAJO TRIBE	APACHE
CHINNEY MINE 1	NAVAJO TRIBE	APACHE
CISCO 1 CAMP NES	NAVAJO TRIBE	APACHE
CLAIM 28	NAVAJO TRIBE	APACHE
CLAIM 31	NAVAJO TRIBE	APACHE
CLAIM 7 & 10	NAVAJO TRIBE	APACHE
CLEVELAND 1	NAVAJO TRIBE	APACHE
COVE 1	NAVAJO TRIBE	APACHE
COVE 2	NAVAJO TRIBE	APACHE
COVE 4	NAVAJO TRIBE	APACHE
COVE MESA	NAVAJO TRIBE	APACHE
COVE MESA 1	NAVAJO TRIBE	APACHE
COVE MESA 10	NAVAJO TRIBE	APACHE
COVE MESA 2	NAVAJO TRIBE	APACHE
COVE MESA VCA	NAVAJO TRIBE	APACHE
COVE MINES	NAVAJO TRIBE	APACHE
DAN TAYLOR 1	NAVAJO TRIBE	APACHE
DENNY LFF 3	NAVAJO TRIBE	APACHE
EAST MESA	NAVAJO TRIBE	APACHE
EAST MESA 1 & 2	NAVAJO TRIBE	APACHE
ETBITTY 1	NAVAJO TRIBE	APACHE
EURIDA	NAVAJO TRIBE	APACHE
FALL DOWN MESA	NAVAJO TRIBE	APACHE
FLAG MESA 1	NAVAJO TRIBE	APACHE
FRANK JR.	NAVAJO TRIBE	APACHE
FRANK NO. 1	NAVAJO TRIBE	APACHE

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 2

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** ARIZONA (CONT'D) *****									
G + G	FOSTER, GEORGE	APACHE	18	12 N	29,0 E	14	SURFACE	<100	50
HANLEY 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	250
HARVEY BEGAY 3	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	50
HARVEY BLACKWATE	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	100
HARVEY BLACKWATE	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	50
HAZEL	NAVAJO TRIBE	APACHE			0		SURFACE	<100	0
HOWARD NEZ 1	NAVAJO TRIBE	APACHE			0		SURFACE	<100	0
JAUNITA 3	PAULSELL, PAT D.	APACHE	14	18 N	25,0 E	14	SURFACE	<100	30
JIM LEE 1	NAVAJO TRIBE	APACHE			0		SURFACE	100 - 1,000	100
JIM LEE SMILEY	NAVAJO TRIBE	APACHE	3	32 N	23,0 E	14	SURFACE	<100	0
JIMMIE BILEEN 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	50
JIMMIE KING	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	0
JOHN KEE TRACT 4	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	50
JOHN LEE BENALLY	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	150
JOHN P. YAZZIE 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	1,000 - 100,000	50
JOHNNY MC COY 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	0
KASEWOOD HANE 1	NAVAJO TRIBE	APACHE	36	33 N	24,0 E	14	UNDERGRO	<100	50
KNIFE EDGE	NAVAJO TRIBE	APACHE	29	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	100
LAST CHANCE	NAVAJO TRIBE	APACHE	12	40 N	26,0 E	14	UNDERGRO	1,000 - 100,000	100
LEASE 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	100
LOOKOUT POINT	NAVAJO TRIBE	APACHE			0		UNDERGRO	1,000 - 100,000	300
MC KENZIE 3	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	0
MELVIN BEVALLY 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	50
MESA 1	NAVAJO TRIBE	APACHE	22	9 N	7,0 E	14	UNDERGRO	1,000 - 100,000	100
MESA 1 1-2	NAVAJO TRIBE	APACHE	22	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	250
MESA 1 1-4	NAVAJO TRIBE	APACHE	22	36 N	29,0 E	14	UNDERGRO	100 - 1,000	200
MESA 1 3-4 MINE	NAVAJO TRIBE	APACHE	28	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	350
MESA 1 MINE 14A	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	250
MESA 1 MINE 16	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	250
MESA 1 MINE 22	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	250
MESA 1 MINE 24	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	200
MESA 2	NAVAJO TRIBE	APACHE	21	36 N	29,0 E	14	UNDERGRO	>100,000	300
MESA 2 1-2 MINE	NAVAJO TRIBE	APACHE	21	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	150
MESA 2 MINE 4	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	0
MESA 3 MINE 1	NAVAJO TRIBE	APACHE	20	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	200
MESA 4	NAVAJO TRIBE	APACHE	16	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	50
MESA 4 1-2	NAVAJO TRIBE	APACHE	8	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	100
MESA 4 1-4	NAVAJO TRIBE	APACHE	17	36 N	29,0 E	14	UNDERGRO	100 - 1,000	250
MESA 4 MINE 19	NAVAJO TRIBE	APACHE			0		SURFACE	<100	150
MESA 5	NAVAJO TRIBE	APACHE	8	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	200
MESA 6	NAVAJO TRIBE	APACHE	8	36 N	29,0 E	14	UNDERGRO	1,000 - 100,000	200
MIKE BRODIE 1	NAVAJO TRIBE	APACHE			0		SURFACE	<100	200
MILDRED 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	0
MONUMENT 2	NAVAJO TRIBE	APACHE			0		SURFACE	>100,000	50
MONUMENT 3	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	250
MONUMENT HEAP LE	NAVAJO TRIBE	APACHE			0		HL-DUMPS	100 - 1,000	50
MONUMENT VALLEY	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	50
N S M 2	HILL, ADAIR	APACHE	34	15 N	26,0 E	14	UNDERGRO	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 3

MIKE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** ARIZONA (CONT'D) *****									
NAKAI CHEE BEGAY	NAVAJO TRIBE	APACHE	12	36 N	28.0 E	14	UNDERGRO	100 - 1,000	150
OAK SPGS GRAVEL	NAVAJO TRIBE	APACHE	32	39 N	31.0 E	14	UNDERGRO	1,000 - 100,000	100
PETTIGREW PERMIT	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	0
PLOT 1+2-AFC PLO	NAVAJO TRIBE	APACHE	12	40 N	28.0 E	14	UNDERGRO	1,000 - 100,000	300
PLOT 10	NAVAJO TRIBE	APACHE	31	39 N	31.0 E	14	UNDERGRO	1,000 - 100,000	50
PLOT 12 SYCACUSE	NAVAJO TRIBE	APACHE	30	39 N	31.0 E	14	UNDERGRO	100 - 1,000	100
PLOT 4(WILLIAM P	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	100
PLOT 6 RATTLESNA	NAVAJO TRIBE	APACHE		12 N	7.0 W	21	UNDERGRO	1,000 - 100,000	50
PLOT 12 RATTLESNA	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	150
POPE 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	50
R F + R	NAVAJO TRIBE	APACHE	19	39 N	31.0 E	14	UNDERGRO	1,000 - 100,000	100
RATTLESNAKE	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	150
RICHARD KING 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	100
ROCK HAT	NAVAJO TRIBE	APACHE			0		SURFACE	<100	350
ROCKY SPRING	NAVAJO TRIBE	APACHE			0		SURFACE	<100	300
ROUGH ROCK SLOPE	NAVAJO TRIBE	APACHE	1	34 N	23.0 E	14	UNDERGRO	<100	50
RUBEN 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	100
SAM POINT	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	0
SANDY K	NAVAJO TRIBE	APACHE			0		SURFACE	<100	0
SCHOOL ROY CLAIM	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	50
SHEEPSKIN MESA	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	100
SHIPROCK	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	200
SHORTY 1 RATTLES	NAVAJO TRIBE	APACHE			0		UNDERGRO	1,000 - 100,000	150
SILENTMAN 1	NAVAJO TRIBE	APACHE			0		SURFACE	<100	100
SIMPSON 181	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	300
STEP MESA	NAVAJO TRIBE	APACHE	30	36 N	29.0 W	22	UNDERGRO	1,000 - 100,000	350
THOMAS BEGAY 1	NAVAJO TRIBE	APACHE	36	34 N	23.0 E	14	SURFACE	<100	0
TODAKOMZIE 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	250
TODECHEENIE 1	NAVAJO TRIBE	APACHE	35	34 N	23.0 E	14	SURFACE	1,000 - 100,000	50
TODICHINEL	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	350
TOME THLANY BEGA	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	100
TOM JOE 7	NAVAJO TRIBE	APACHE	12	36 N	28.0 E	14	UNDERGRO	<100	150
TOM KLEC CLAIM	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	0
TOM MORGAN 1	NAVAJO TRIBE	APACHE			0		SURFACE	<100	0
TOM NAKAI CHEE 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	100
TOM NAKAI CHEE 6	NAVAJO TRIBE	APACHE			0		UNDERGRO	1,000 - 100,000	100
TOM WILSON	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	0
TONECHLIE	NAVAJO TRIBE	APACHE			0		SURFACE	<100	150
TOMY 1	NAVAJO TRIBE	APACHE			0		SURFACE	<100	200
TOPAHA	NAVAJO TRIBE	APACHE	27	36 N	29.0 E	14	UNDERGRO	100 - 1,000	0
TREE MESA	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	100
TSOSIE 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	100
UPPER RED WASH	NAVAJO TRIBE	APACHE			0		UNDERGRO	100 - 1,000	200
VCA RATTLESNAKE	NAVAJO TRIBE	APACHE			0		SURFACE	<100	0
WARHOOP GROUP	APACHE MINING CO	APACHE	30	33 N	29.0 E	14	SURFACE	100 - 1,000	50
WEST MESA	NAVAJO TRIBE	APACHE			0		UNDERGRO	<100	100
WEST RESERVATION	NAVAJO TRIBE	APACHE			0		UNDERGRO	1,000 - 100,000	100
WILLY WATERS	NAVAJO TRIBE	APACHE			0		UNDERGRO	1,000 - 100,000	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 4

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** ARIZONA (CONT'D) *****									
ZONA 1	NAVAJO TRIBE	APACHE			0		UNDERGRO	1,000 - 100,000	250
STAR 1	UOCO	COCHISE			0		UNDERGRO	<100	100
WINDMILL 1	SMITH + ROYSTON	COCHISE			0		SURFACE	<100	50
A + B 13	NAVAJO TRIBE	COCOA	14	31 N	9.0 E	14	SURFACE	<100	50
A + B 2	NAVAJO TRIBE	COCOA	5	28 N	9.0 E	14	SURFACE	100 - 1,000	50
A + B 3	NAVAJO TRIBE	COCOA	21	28 N	9.0 E	14	SURFACE	100 - 1,000	50
A + B 5	NAVAJO TRIBE	COCOA	3	31 N	9.0 E	14	SURFACE	100 - 1,000	50
A + B 7	NAVAJO TRIBE	COCOA	20	31 N	9.0 E	14	SURFACE	<100	50
A MALONEY 2	NAVAJO TRIBE	COCOA			0		SURFACE	<100	50
ALYCE TOLENO	NAVAJO TRIBE	COCOA			0		SURFACE	1,000 - 100,000	50
AMOS CHEE 3	NAVAJO TRIBE	COCOA	13	25 N	11.0 E	14	UNDERGRO	<100	100
AMOS CHEE 8	NAVAJO TRIBE	COCOA	34	27 N	10.0 E	14	UNDERGRO	100 - 1,000	50
B, + B, 11A	PARIA URAN. + OIL	COCOA			0		SURFACE	<100	100
BAKER	UNKNOWN CONTROL	COCOA			0		SURFACE	<100	0
BIG BLUE	ALPINE URANIUM C	COCOA			0		UNDERGRO	<100	150
BOYD TISI 1	NAVAJO TRIBE	COCOA	31	28 N	10.0 E	14	SURFACE	<100	50
BOYD TISI 2	NAVAJO TRIBE	COCOA	25	29 N	9.0 F	14	SURFACE	100 - 1,000	50
CASEY 3	NAVAJO TRIBE	COCOA	3	29 N	9.0 F	14	SURFACE	<100	50
CHARLES HUSKON 1	NAVAJO TRIBE	COCOA	12	26 N	10.0 E	14	SURFACE	100 - 1,000	50
CHARLES HUSKON 1	NAVAJO TRIBE	COCOA			0		SURFACE	100 - 1,000	50
CHARLES HUSKON 2	NAVAJO TRIBE	COCOA	8	29 N	10.0 F	14	SURFACE	1,000 - 100,000	50
CHARLES HUSKON 4	NAVAJO TRIBE	COCOA	11	26 N	10.0 E	14	SURFACE	1,000 - 100,000	50
CHARLES HUSKON 5	NAVAJO TRIBE	COCOA	1	30 N	9.0 E	14	SURFACE	100 - 1,000	50
CHARLES HUSKON 9	NAVAJO TRIBE	COCOA	2	26 N	10.0 E	14	SURFACE	100 - 1,000	50
CLIFF CANYON	BAKER, RILEY	COCOA			0		UNDERGRO	<100	0
COPPER 1 + WILLA	KEANEY + FOLSON	COCOA			0		UNDERGRO	<100	50
E HUSKON 34	NAVAJO TRIBE	COCOA	8	29 N	10.0 E	14	SURFACE	1,000 - 100,000	50
E HUSKON 35	NAVAJO TRIBE	COCOA	25	28 N	10.0 F	14	SURFACE	<100	100
EARL HUSKON 1	NAVAJO TRIBE	COCOA	27	32 N	9.0 E	14	SURFACE	100 - 1,000	50
EARL HUSKON 3	NAVAJO TRIBE	COCOA	26	32 N	9.0 F	14	SURFACE	1,000 - 100,000	50
EL PEQUITO	EL PEQUITO MNG C	COCOA	11	40 N	7.0 E	14	SURFACE	100 - 1,000	50
ELWOOD CANYON 2	NAVAJO TRIBE	COCOA	18	29 N	10.0 E	14	UNDERGRO	100 - 1,000	100
ELWOOD THOMPSON	NAVAJO TRIBE	COCOA	1	26 N	10.0 E	14	UNDERGRO	1,000 - 100,000	100
EMMETT LEE 1	NAVAJO TRIBE	COCOA	2	26 N	10.0 E	14	SURFACE	100 - 1,000	50
EMMETT LEE 3	NAVAJO TRIBE	COCOA	12	26 N	10.0 E	14	SURFACE	100 - 1,000	100
FOLEY 5(YAZZIE)1	NAVAJO TRIBE	COCOA	24	29 N	10.0 E	14	UNDERGRO	100 - 1,000	100
GRUB CLAIM 14 #2	FILLMORE, ROBERT	COCOA	16	27 N	10.0 E	14	SURFACE	<100	50
HENRY SLOAN 1	NAVAJO TRIBE	COCOA	35	32 N	9.0 E	14	SURFACE	100 - 1,000	50
HOBSTEN REZ	NAVAJO TRIBE	COCOA	5	27 N	12.0 E	14	UNDERGRO	<100	50
HOWARD 1	LAUDERDALE MNG +	COCOA	7	27 N	10.0 F	14	SURFACE	<100	50
HUSKON 1	NAVAJO TRIBE	COCOA	23	29 N	9.0 E	14	SURFACE	1,000 - 100,000	50
HUSKON 10	NAVAJO TRIBE	COCOA	28	28 N	10.0 E	14	SURFACE	1,000 - 100,000	50
HUSKON 11	NAVAJO TRIBE	COCOA	33	28 N	10.0 E	14	SURFACE	1,000 - 100,000	50
HUSKON 12	NAVAJO TRIBE	COCOA	15	29 N	9.0 E	14	SURFACE	1,000 - 100,000	50
HUSKON 14	NAVAJO TRIBE	COCOA	35	29 N	9.0 E	14	SURFACE	<100	50
HUSKON 17	NAVAJO TRIBE	COCOA	15	29 N	9.0 E	14	SURFACE	1,000 - 100,000	50
HUSKON 2	NAVAJO TRIBE	COCOA	24	29 N	9.0 F	14	SURFACE	1,000 - 100,000	50
HUSKON 3	NAVAJO TRIBE	COCOA	12	28 N	9.0 E	14	SURFACE	1,000 - 100,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 5

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** ARIZONA (CONT'D) *****									
HUSKON 6	NAVAJO TRIBE	COCONINO	27	30 N	9.0 E	14	SURFACE	100 - 1,000	50
HUSKON 7	NAVAJO TRIBE	COCONINO	19	28 N	10.0 E	14	SURFACE	1,000 - 100,000	50
HUSKON 8	NAVAJO TRIBE	COCONINO	30	28 N	9.0 E	14	SURFACE	100 - 1,000	50
J SEYALLIE	NAVAJO TRIBE	COCONINO	12	26 N	10.0 E	14	UNDERGRO	1,000 - 100,000	50
JACK DANIELS 1	NAVAJO TRIBE	COCONINO	11	29 N	9.0 E	14	SURFACE	1,000 - 100,000	50
JACK DANIELS 3	NAVAJO TRIBE	COCONINO			0 E	14	SURFACE	<100	50
JACK DANIELS 4	NAVAJO TRIBE	COCONINO	11	29 N	9.0 E	14	SURFACE	<100	50
JACK DANIELS 5	NAVAJO TRIBE	COCONINO	11	29 N	9.0 E	14	SURFACE	100 - 1,000	50
JACK HUSKON 3	NAVAJO TRIBE	COCONINO	3	28 N	10.0 E	14	SURFACE	1,000 - 100,000	150
JACKPOT 1	NAVAJO TRIBE	COCONINO	14	27 N	10.0 E	14	SURFACE	100 - 1,000	50
JACKPOT 40	NAVAJO TRIBE	COCONINO	14	27 N	10.0 E	14	SURFACE	100 - 1,000	50
JACKPOT 5	NAVAJO TRIBE	COCONINO	14	27 N	10.0 E	14	SURFACE	<100	50
JEEPSTER 1	NAVAJO TRIBE	COCONINO	26	30 N	9.0 E	14	SURFACE	1,000 - 100,000	50
JIMMY BOONE	NAVAJO TRIBE	COCONINO	1	39 N	7.0 F	14	SURFACE	<100	0
JUAN HORSE 3	NAVAJO TRIBE	COCONINO	30	24 N	10.0 E	14	SURFACE	1,000 - 100,000	50
JUAN HORSE 4	NAVAJO TRIBE	COCONINO	31	29 N	10.0 E	14	SURFACE	1,000 - 100,000	100
JULIUS CHEE 2	NAVAJO TRIBE	COCONINO	3	26 N	10.0 E	14	SURFACE	100 - 1,000	50
JULIUS CHEE 3	NAVAJO TRIBE	COCONINO	10	26 N	10.0 E	14	UNDERGRO	100 - 1,000	50
JULIUS CHEE 4	NAVAJO TRIBE	COCONINO	2	26 N	10.0 E	14	SURFACE	1,000 - 100,000	50
JUNE	NAVAJO TRIBE	COCONINO	26	39 N	7.0 E	14	SURFACE	<100	50
KACHINA 6	NAVAJO TRIBE	COCONINO	2	29 N	9.0 E	14	SURFACE	1,000 - 100,000	100
L. LITTLEMAN 1+2	NAVAJO TRIBE	COCONINO	30	9 N	6.0 F	14	SURFACE	<100	50
L. LITTLEMAN 2	NAVAJO TRIBE	COCONINO	24	29 N	9.0 E	14	SURFACE	1,000 - 100,000	50
L. LITTLEMAN 3	NAVAJO TRIBE	COCONINO	35	29 N	9.0 E	14	SURFACE	<100	50
LIBA GROUP	BLACK, C. B.	COCONINO	4	27 N	10.0 E	14	UNDERGRO	1,000 - 100,000	50
LUSTER 1	LAUDERDALE MNG +	COCONINO	17	27 N	10.0 E	14	UNDERGRO	100 - 1,000	50
M JOHNSON 4	NAVAJO TRIBE	COCONINO	11	32 N	9.0 E	14	UNDERGRO	<100	50
MANUEL DENETSONE	NAVAJO TRIBE	COCONINO	2	28 N	10.0 E	14	UNDERGRO	100 - 1,000	50
MAX HUSKON 1 7	NAVAJO TRIBE	COCONINO	23	32 N	9.0 F	14	SURFACE	<100	50
MAX JOHNSON 1	NAVAJO TRIBE	COCONINO	23	29 N	9.0 E	14	SURFACE	1,000 - 100,000	50
MAX JOHNSON 10	NAVAJO TRIBE	COCONINO	20	29 N	9.0 E	14	SURFACE	100 - 1,000	50
MAX JOHNSON 7	NAVAJO TRIBE	COCONINO	34	27 N	10.0 E	14	SURFACE	100 - 1,000	50
MAX JOHNSON 9	NAVAJO TRIBE	COCONINO	24	29 N	9.0 E	14	SURFACE	1,000 - 100,000	50
MONTZUMA 1	NAVAJO TRIBE	COCONINO	1	28 N	9.0 E	14	SURFACE	<100	50
MONTZUMA 2	NAVAJO TRIBE	COCONINO	4	29 N	9.0 F	14	SURFACE	100 - 1,000	50
MONTZUMA 7A	NAVAJO TRIBE	COCONINO	3	29 N	9.0 E	14	SURFACE	<100	0
MONTZUMA 7B	NAVAJO TRIBE	COCONINO	5	29 N	9.0 F	14	SURFACE	<100	50
MONTZUMA 7C	NAVAJO TRIBE	COCONINO	33	30 N	9.0 E	14	SURFACE	<100	50
MURPHY 1,7,10,16	GLASSCOCK, HOWELL	COCONINO	22	27 N	10.0 E	14	UNDERGRO	1,000 - 100,000	50
NAVAJO 26	UNKNOWN	COCONINO	18	26 N	10.0 E	14	UNDERGRO	<100	50
ORPHAN LODGE (GCA	COTTER CORPORATN	COCONINO	14	31 N	2.0 E	14	UNDERGRO	>100,000	1600
PAUL HUSKIE 20	NAVAJO TRIBE	COCONINO	22	28 N	9.0 E	14	SURFACE	<100	50
PAUL HUSKIE 21	NAVAJO TRIBE	COCONINO		N	0 E	14	SURFACE	<100	50
RAINBOW	ALPINE URANIUM C	COCONINO			0		SURFACE	<100	150
RAMCO 20	NAVAJO TRIBE	COCONINO	11	27 N	10.0 E	14	SURFACE	1,000 - 100,000	50
RAMCO 21	NAVAJO TRIBE	COCONINO	11	27 N	10.0 E	14	SURFACE	1,000 - 100,000	50
RAMCO 22	NAVAJO TRIBE	COCONINO	2	27 N	10.0 F	14	SURFACE	1,000 - 100,000	50
RAMCO 24	NAVAJO TRIBE	COCONINO	12	26 N	10.0 E	14	UNDERGRO	1,000 - 100,000	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 6

WINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** ARIZONA (CONT'D) *****									
REDWING	MARBLE CANYON MG	COCONINO	34	41 N	7.0 E	14	UNDERGRO	<100	50
RIDENOUR	HUALAPAI TRIBE	COCONINO	6	31 N	8.0 W	14	UNDERGRO	<100	50
RIVERVIEW	UNKNOWN CONTROLR	COCONINO	8	26 N	10.0 E	14	UNDERGRO	100 - 1,000	50
RYAN 1	NAVAJO TRIBE	COCONINO	14	28 N	10.0 E	14	SURFACE	100 - 1,000	0
RYAN 2	NAVAJO TRIBE	COCONINO	11	27 N	10.0 E	14	SURFACE	1,000 - 100,000	100
S4,9,16,21CAMERO	WESTERN NUCLEAR	COCONINO	4	27 N	10.0 E	14	SURFACE	100 - 1,000	50
SAM 7	CONSUMERS AGENCY	COCONINO			0		SURFACE	<100	0
SEC 1 EM SWO	MORDELL, A.C.	COCONINO	1	27 N	9.0 E	14	UNDERGRO	<100	50
SHARON LYNN	URANIUM MNG.CO.	COCONINO	33	18 N	23.0 E	14	SURFACE	<100	50
SUN VALLEY MINE	GRAND CANYON URA	COCONINO			0		UNDERGRO	100 - 1,000	50
THOMAS 1	NAVAJO TRIBE	COCONINO	21	38 N	7.0 E	14	SURFACE	100 - 1,000	50
TOMMY	NAVAJO TRIBE	COCONINO	23	39 N	7.0 E	14	SURFACE	<100	50
VERMILLION	UNITED DYEL.CO.	COCONINO			0		UNDERGRO	<100	50
WARD TERRACE TRA	NAVAJO TRIBE	COCONINO			0		UNDERGRO	<100	0
YAZZIE 1	NAVAJO TRIBE	COCONINO	14	27 N	10.0 E	14	SURFACE	100 - 1,000	50
YAZZIE 101	NAVAJO TRIBE	COCONINO	19	29 N	10.0 E	14	SURFACE	1,000 - 100,000	50
YAZZIE 102	NAVAJO TRIBE	COCONINO	19	28 N	10.0 E	14	UNDERGRO	1,000 - 100,000	50
YAZZIE 2	NAVAJO TRIBE	COCONINO	14	29 N	10.0 E	14	SURFACE	1,000 - 100,000	50
YAZZIE 312	NAVAJO TRIBE	COCONINO	25	29 N	9.0 E	14	SURFACE	1,000 - 100,000	50
YELLOW JEEP 7 A+	NAVAJO TRIBE	COCONINO	25	29 N	11.0 E	14	UNDERGRO	100 - 1,000	100
REE CAVE	BURFE, ROSS	GILA			0		SURFACE	<100	50
BIG BUCK GROUP	MFTBEL HNG+ EXPL.	GILA			0		UNDERGRO	100 - 1,000	50
BLACK BRUSH	WESTERN MINING +	GILA			0		UNDERGRO	<100	100
DONNA LEE	SMITH, H.C.	GILA			0		UNDERGRO	<100	50
FIRST CHANCE(GLO	BAKER + CLINE	GILA			0		UNDERGRO	<100	50
HOPE CLAIM	WYOMING MINERALS	GILA			0		UNDERGRO	1,000 - 100,000	150
MORSESHOE	HAUGHT, ALFRED	GILA			0		SURFACE	<100	0
JON	WYOMING MINERALS	GILA			0		UNDERGRO	1,000 - 100,000	50
LITTLE JOE	NICOLS+J.BANKS+	GILA			0		UNDERGRO	1,000 - 100,000	100
LUCKY BOY	STACY + EPPINGER	GILA			0		UNDERGRO	1,000 - 100,000	50
LUCKY STOP	WYOMING MINERALS	GILA			0		UNDERGRO	1,000 - 100,000	100
MELINDA GROUP	MORGAN+SHAW+HILB	GILA			0		UNDERGRO	1,000 - 100,000	50
RED BLUFF	WYOMING MINERALS	GILA			0		UNDERGRO	1,000 - 100,000	50
RED CLIFF 1	PATTERSON, ROY	GILA			0		SURFACE	<100	50
SUCKWHITE	TULSA MINERALS C	GILA			0		UNDERGRO	1,000 - 100,000	100
SUE	ARIZONA GLOBE UP	GILA			0		UNDERGRO	100 - 1,000	50
TOMATO JUICE	LEWIS,FRANK ETAL	GILA			0		UNDERGRO	100 - 1,000	100
MORFMAN CLAIM	NICHOLS, CHARLES	GILA			0		UNDERGRO	100 - 1,000	100
FLAT TIRE	MARALSON,A.M.JR.	GRAHAM			0		SURFACE	<100	50
B + M	BICKLE + MANLEY	MARICOPA			0		UNDERGRO	<100	50
MORSESHOE GROUP	JOHNSON + YOUNG	MARICOPA			0		UNDERGRO	<100	50
MALAPAI	TOMTO NATL FORRE	MARICOPA			0		SURFACE	<100	100
CHAPEL	KNIGHT,ANDRIE G	MOHAVE	25	33 N	10.0 W	14	UNDERGRO	<100	50
CERYL M 1	THOMPSON, L.A.	MOHAVE			0		UNDERGRO	<100	0
DEMO GROUP 24	SPARTAN MINING C	MOHAVE			0		UNDERGRO	<100	50
HACKS CANYON	WESTERN NUCLEAR	MOHAVE	26	37 N	5.0 W	14	UNDERGRO	1,000 - 100,000	150
RADDON 1	ANNESLY GRIFFITH	MOHAVE			0		UNDERGRO	<100	50
ALMA-SEEGIN	NAVAJO TRIBE	NAVAJO		41 N	19.0 E	14	UNDERGRO	1,000 - 100,000	200

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 7

MINES NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** ARIZONA (CONT'D) *****									
BIG CHIEF	NAVAJO TRIBE	NAVAJO	23	41 N	19.0 E	14	UNDERGRD	1,000 - 100,000	130
BIG FOUR 2	NAVAJO TRIBE	NAVAJO		41 N	19.0 E	14	UNDERGRD	1,000 - 100,000	130
BOOTHACK	NAVAJO TRIBE	NAVAJO	33	41 N	19.0 E	14	UNDERGRD	1,000 - 100,000	400
CARNOTITE CANYON	NAVAJO TRIBE	NAVAJO			0		SURFACE	<100	300
FERM	NAVAJO TRIBE	NAVAJO	34	41 N	19.0 E	14	UNDERGRD	1,000 - 100,000	100
FIRELIGHT 6-NASC	NAVAJO TRIBE	NAVAJO	36	41 N	19.0 E	14	UNDERGRD	1,000 - 100,000	200
GOOF 6	BERRY + HATFIELD	NAVAJO			0		SURFACE	<100	0
GOTHE	NAVAJO TRIBE	NAVAJO			0		UNDERGRD	<100	200
HANSEN	COSTON + BARTON	NAVAJO	1	18 N	19.0 E	14	UNDERGRD	100 - 1,000	50
J CITY 1	F. + M. MINING	NAVAJO	33	19 N	19.0 E	14	UNDERGRD	<100	0
KAY GROUP	DAVIS + HATCH	NAVAJO	20	18 N	23.0 E	14	SURFACE	100 - 1,000	100
LITTLE JOHN 1,2+	YOUNG, WEPRIILL	NAVAJO	2	17 N	23.0 E	14	SURFACE	<100	0
MAC 3	MAC KAHON, JOHN	NAVAJO	4	17 N	23.0 E	14	SURFACE	<100	0
MITCHELL BUTTE	NAVAJO TRIBE	NAVAJO			0		UNDERGRD	1,000 - 100,000	150
MITTEN 2	NAVAJO TRIBE	NAVAJO			0		SURFACE	1,000 - 100,000	50
MONUMENT 1	NAVAJO TRIBE	NAVAJO	17	41 N	20.0 E	14	UNDERGRD	1,000 - 100,000	50
MOONLIGHT	NAVAJO TRIBE	NAVAJO	14	41 N	19.0 E	14	UNDERGRD	>100,000	150
MORALE	NAVAJO TRIBE	NAVAJO	19	24 N	22.0 E	14	SURFACE	100 - 1,000	50
NAVAJO	HALL, JOE	NAVAJO	26	20 N	23.0 E	14	UNDERGRD	<100	0
NEW MEX + ARTZ 1	BAY SHORE MINING	NAVAJO			0 E	14	SURFACE	<100	0
RAINBOW SMITH 1+	RAINBOW URANIUM	NAVAJO	36	16 N	22.0 E	14	SURFACE	<100	0
RESERVATION	NAVAJO TRIBE	NAVAJO			0		SURFACE	<100	50
ROCK GARDEN 25	BAY SHORE MINING	NAVAJO	22	19 N	23.0 E	14	SURFACE	<100	50
SAIN R20E T19N +	PO-ELL, L. +	NAVAJO			0		SURFACE	<100	50
SALLY	NAVAJO TRIBE	NAVAJO			0		UNDERGRD	<100	50
SAM CHARLEE 1	NAVAJO TRIBE	NAVAJO			0		SURFACE	<100	150
SEC. 33-18N-23E	N. MEX. ARIZ. LAND C	NAVAJO	24	19 N	20.0 E	14	SURFACE	<100	50
SM 60-TRACT 11	NAVAJO TRIBE	NAVAJO			0		UNKNOWN	1,000 - 100,000	0
SPENCER 3 MINE	NAVAJO TRIBE	NAVAJO			0		UNDERGRD	100 - 1,000	200
STARLIGHT (EAST)	NAVAJO TRIBE	NAVAJO	16	41 N	19.0 E	14	UNDERGRD	1,000 - 100,000	150
SUNLIGHT	NAVAJO TRIBE	NAVAJO	22	41 N	19.0 E	14	UNDERGRD	1,000 - 100,000	400
SUNLIGHT SOUTH	NAVAJO TRIBE	NAVAJO	22	41 N	19.0 E	14	UNDERGRD	1,000 - 100,000	200
SUNRISE	BOYER + PETTIT	NAVAJO	4	17 N	23.0 E	14	SURFACE	<100	0
WINSLOW 6 + 7	TWIN STATES URAN	NAVAJO	32	20 N	17.0 E	14	UNDERGRD	<100	50
BLACK DYKE	HOLLER, J.	PIMA			0		UNDERGRD	<100	100
LINDA LEE 2	RUBERSON + PAYTAIN	PIMA			0		UNDERGRD	<100	50
DURANIUM 1	SANTA CRUZ URAN. C	SANTA CRUZ			0		UNDERGRD	100 - 1,000	50
MONTANA GROUP	MILLER, HUGO W.	SANTA CRUZ			0		SURFACE	<100	50
WHITE OAK	CLARKE, W. H.	SANTA CRUZ			0		SURFACE	<100	0
MOON BEAM	DAY + ADAMS	YAVAPAI			0		SURFACE	<100	0
SEVEN STARS	DUNNING, CHARLES	YAVAPAI			0		SURFACE	<100	0
UR-ARIZ(ANDERSON	MINERALS EXPL CO	YAVAPAI	9	11 N	10.0 W	14	SURFACE	1,000 - 100,000	300
F H BARNEY	HOUSLEY, GLEN	UNKNOWN			0		SURFACE	<100	0
***** CALIFORNIA *****									
CROWN MINES	CROWN MINING CO	IMPERIAL			0		UNDERGRD	<100	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 8

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MERID.	RANGE	TOWNSHIP	SEC.	COUNTY	CONTROLLER NAME	MINING METHOD
***** CALIFORNIA (CONT'D) *****									
GYP		50		0			IMPERIAL	ROBBINS, HAPRIET	SURFACE
COSO CLAIM		100	25	37.0 E	20 S	20	INYO	FED. RES-PNT-UP	SURFACE
BUCKEYE GROUP	100 - 1,000	100	17	32.0 E	27 S	20	KERN	MIRACLE MINING C	SURFACE
KERGON 1		150	20	32.0 E	27 S	20	KERN	GREAT LAKE OIL	SURFACE
LITTLE SPARKLEP		100	17	32.0 E	27 S	20	KERN	KERN URANIUM CO	UNDERGRO
NOB HILL	100 - 1,000	100	23	38.0 E	32 S	20	KERN	CASTLE BUTTE URA	UNDERGRO
OWEN 3		50	33	22.0 E	31 S	20	KERN	OWEN MINING CO.	SURFACE
PIONEER 1		0	26	35.0 E	27 S	20	KERN	GREAT SALT LAKE	SURFACE
CORNELIA C 2		0	7	18.0 E	23 N	20	LASSEN	BAKER + SONS	SURFACE
LOLA G	100 - 1,000	50		0			LASSEN	BAKER + MILLER	SURFACE
PATSY GROUP		50	33	24.0 E	6 S	20	MADERA	WAGNON, EARL M.	UNDERGRO
EMERALD 1		50		0			MONO	NEVADA GOLDFIELD	SURFACE
GROUP 102		50	13	20.0 E	6 S	25	RIVERSIDE	UNITED MNG+DEVEL	UNDERGRO
LUCKY 3		0		0			RIVERSIDE	HARDING J.L.	SURFACE
MELODY GROUP		50	7	21.0 E	6 S	25	RIVERSIDE	MINERAL EXPLOIT.	UNDERGRO
NORTHEAST GROUP	1,000 - 100,000	50	19	21.0 E	6 S	25	RIVERSIDE	SAFRANEK, C.+J.	UNDERGRO
WHITE DOME		0		0			RIVERSIDE	DAY + ADAMS	SURFACE
BIG HUNCH 1		50		0			SAN BERNARDINO	BIG HUNCH URAN 1	UNDERGRO
MINDTE		50	29	2.0 E	2 N	25	SAN BERNARDINO	MINDTE, P.A.	SURFACE
ROSE OF SHARON		100		0			SAN BERNARDINO	DAILESS+DKO,B+G	SURFACE
PINE CONE+SL STR		50		0			SIERRA	DOG VALLEY URAN.	UNDERGRO
JUNIPER(SHRA PS)	1,000 - 100,000	100	8	20.0 E	5 N	20	TUOLUMNE	PATHFINDER	SURFACE
***** COLORADO *****									
BLUE HORSE + NAN		0	13	19.0 W	44 N	22	BOULDER	DUNCAN HATF	SURFACE
FAIR DAY MINE		250	23	72.0 W	2 N	06	BOULDER	BUSBY, P.L.	UNDERGRO
KIPP LEASE	1,000 - 100,000	50		0			BOULDER	MTN STATES OIL +	SURFACE
MILL TAILINGS		0		0			BOULDER	PANORNEY OZARK	MISC.-PB
MILLER LEASE		50		0			BOULDER	MTN STATES OIL +	SURFACE
MOUNTAIN GOAT		50		0			BOULDER	FRONTIER MINING	SURFACE
BISK		50		0			BOULDER	ENIBETA CORP	UNDERGRO
VICTORY GROUP	100 - 1,000	50		0			BOULDER	JIMTOWN URANIUM	UNDERGRO
HIGHLANDER		100		0			CLEAR CREEK	UNCOMPAGNE URAN	SURFACE
LITTLE WARRIOR 1		50		0			CLEAR CREEK	SEACOL CUPP	UNDERGRO
MARTHA E.		150		0			CLEAR CREEK	MTN STATES URAN.	SURFACE
SPANISH BAR GROU		100		0			CLEAR CREEK	SPANISH BAR URAN	UNDERGRO
BECK MOUNTAIN LO		0		0			CUSTER	WEST, GEORGE A	SURFACE
KING MIDAS		50		0			CUSTER	KLINE, BILL+ASSOC	SURFACE
WALTERS RANCH		0		0			CUSTER	MUNDY, LEWIS C.	SURFACE
ARROWHEAD GROUP		50		0			DOLORES	ARROWHEAD MINING	SURFACE
BARLO+ CREEK		50	15	10.0 W	40 N	22	DOLORES	FOOTE MINERALS	UNDERGRO
BROKEN THUMB 2		100		0			DOLORES	PETTIGREW, HARLEY	SURFACE
GOOD HOPE-NEVADA		0		0			DOLORES	FM PINKERTON	UNDERGRO
RAINEY DAY	100 - 1,000	0		0			DOLORES	WILLIFORD, BEN J	UNDERGRO
SOUTH BARLOW		0		0			DOLORES	FOOTE MINERALS	UNDERGRO
ARROWHEAD 1	100 - 1,000	100		0			EAGLE	SKYLAND ENGR.CO.	SURFACE

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 9

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CON'D) *****									
DORADO	UNKNOWN CONTRLR	EAGLE			0		SURFACE	<100	0
LUCKY BEN LEAST	YOUNG, CLAYBORNE	EL PASO			0		SURFACE	100 - 1,000	50
BILL + HUD 2 + 4	HARVEY, W.P., JR.	FREMONT			0		UNDERGRO	<100	50
CAP ROCK 40	JOHNSON C.B., INC.	FREMONT	30	17 S	72.0 W	06	SURFACE	<100	50
CLAIM 2	BEN RIVER MNG	FREMONT	16	19 S	71.0 W	06	SURFACE	<100	0
DICKSON-SHOOPER	JUNIPER OIL + MNG	FREMONT	26	17 S	73.0 W	06	SURFACE	1,000 - 100,000	50
DILLEY RANCH	BEATON, JOHN	FREMONT			0		UNDERGRO	100 - 1,000	50
FIRST CHANCE (TAL)	HURD + ASSOC.	FREMONT			0		UNDERGRO	100 - 1,000	50
HIGH PARK	COTTER CORPORATE	FREMONT			0		SURFACE	<100	0
HONEST JOHN 1-15	STURBAUM, STANLEY	FREMONT			0		UNDERGRO	100 - 1,000	0
JOAN 2	SEACOL INC	FREMONT	14	17 S	73.0 W	22	SURFACE	1,000 - 100,000	150
KNOB HILL ORE #0	HEISEN, EARL	FREMONT	22	17 S	73.0 W	06	UNDERGRO	1,000 - 100,000	50
LAST CHANCE	HEISEN, CHARLES	FREMONT	31	17 S	72.0 W	06	UNDERGRO	1,000 - 100,000	100
LIGHTNING 2	REYNOLDS + PRADY	FREMONT			0		UNDERGRO	100 - 1,000	150
LITTLE ABNER 1	MURRIS RECOVERY	FREMONT	14	17 S	73.0 W	06	SURFACE	1,000 - 100,000	100
MARY L. 1	SMALLER, F.J.	FREMONT	13	17 S	73.0 W	06	UNDERGRO	1,000 - 100,000	50
MISERY MINE 7	MC CORMACK, ROBT.	FREMONT			0		UNDERGRO	<100	50
MOOSE RAINBOW	JUNIPER OIL + MNG	FREMONT			0		SURFACE	100 - 1,000	50
PICNIC TREE	CYPRUS MINES	FREMONT	26	17 S	73.0 W	06	SURFACE	1,000 - 100,000	100
RAINBOW DRF BODY	JUNIPER OIL + MNG	FREMONT			0		SURFACE	100 - 1,000	50
RED CLIFF 30	UNKNOWN CONTRLR	FREMONT			0		UNDERGRO	1,000 - 100,000	50
SAND CREEK 4	CURTIS + THORPE	FREMONT	1	16 S	71.0 W	06	SURFACE	<100	50
SEC. 36, 17S-73N-4E	US BANK-GRND JCT	FREMONT	36	17 S	73.0 W	06	UNDERGRO	1,000 - 100,000	150
SEC. 36, 17S-73N-4E	US BANK-GRND JCT	FREMONT	36	17 S	73.0 W	06	UNDERGRO	1,000 - 100,000	100
SPRING VALLEY	JONES + TAYLOR	FREMONT			0		SURFACE	<100	0
SUNSHINE GROUP	OLIVER GLEN + WED	FREMONT	30	17 S	72.0 W	06	SURFACE	1,000 - 100,000	100
THORNE 9+10	ATLAS MINERALS	FREMONT	26	17 S	73.0 W	06	UNDERGRO	1,000 - 100,000	100
ELK VAN TUNNELL	MASCON BROS + OLS	GARFIELD			0		SURFACE	<100	0
END OF TRAIL 1	DOBBS F.C.	GARFIELD			0		UNDERGRO	100 - 1,000	50
ENTERPRISE 1,2+3	ENTERPRISE MNG.	GARFIELD			0		SURFACE	<100	0
GARFIELD MINE	UNION CARBIDE CP	GARFIELD			0		UNDERGRO	1,000 - 100,000	300
HOMESTEAD	HAYES EXPL + URAN	GARFIELD			0		UNDERGRO	<100	0
INCORPORATED 1	BENNETTIS + ASSOC	GARFIELD			0		SURFACE	<100	0
LOTTI B	NEW CASTLE URAN.	GARFIELD			0		SURFACE	<100	50
MARHOLA LOSE 1-1	JEPSON + HOLT	GARFIELD			0		UNDERGRO	<100	0
RIFLE MINE	UNION CARBIDE CP	GARFIELD			0		UNDERGRO	>100,000	300
WARD GULCH	LEHR, VERNON L.	GARFIELD			0		SURFACE	<100	0
BOYANZA	RUTTER, ROBERT L	GILPIN			0		SURFACE	<100	50
ROOT RANCH LEASE	ROOT RANCH	GILPIN			0		SURFACE	<100	50
TWO SISTERS	UNITED MNG. + L.	GILPIN			0		SURFACE	<100	200
WOOD MINE	LYMAN, BOB	GILPIN			0		UNDERGRO	<100	600
ALASKA HUMES	HUMES, FRED	GRAND	21	5 S	82.0 W	06	UNDERGRO	100 - 1,000	50
HUME	PIONEER EXPL. CO.	GRAND			0		SURFACE	<100	0
LUCKY JACK	SCHNEITZLE, H.J.	GRAND			0		SURFACE	<100	50
UNDECIDED 4	U S DEPT WILDLIF	GRAND			0		UNDERGRO	100 - 1,000	50
BIG RED 27	TURNER + HACKNEY	GUNNISON	11	49 N	5.0 E	06	UNDERGRO	100 - 1,000	50
BETH 1-10	MULLINS + BEAUCHAMP	HINSDALE		44	5.0	22	UNDERGRO	<100	50
ANAL 1	SECURITY EXPL CO	HUERFANO	17	25 S	70.0 W	06	SURFACE	<100	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 10

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
STUMBLING STUD	BRISCOE, G.E.+ASSO	HUERFANO			0		UNDERGRO	100 - 1,000	50
ASCENSION MINE	FERR MC GEE	JEFFERSON	24	3 S	71.0 W	06	UNDERGRO	1,000 - 100,000	200
AUBREY LAD-IG LS	RESERVE OIL+MINES	JEFFERSON			0		UNDERGRO	1,000 - 100,000	50
FORK PROSPECT	ROCKY MTN. ENERGY	JEFFERSON			0		UNDERGRO	<100	50
GRAPE VINE 1	E.E.LF=IS, INC.	JEFFERSON	29	4 S	79.0 W	06	SURFACE	1,000 - 100,000	100
LADWIG 2 PROSPEC	ENERGY FUELS MUC	JEFFERSON	19	3 S	70.0 W	06	UNDERGRO	1,000 - 100,000	100
LEYDEN MINE	MORFNO URAN, CORP	JEFFERSON			0		UNDERGRO	100 - 1,000	50
MANN RANCH	J L MANN URANIUM	JEFFERSON	12	5 S	70.0 W	06	UNDERGRO	1,000 - 100,000	100
MENA 1	UNKNOWN CONTROLR	JEFFERSON	26	2 S	71.0 W	06	UNDERGRO	1,000 - 100,000	50
OHMAN MINE	OHMAN, LEIGH D.	JEFFERSON			0		UNDERGRO	<100	50
PALLAORO LEASE	FOUR CORNERS OIL	JEFFERSON			0		UNDERGRO	100 - 1,000	150
QUATMAN LEASE	UNKNOWN CONTROLR	JEFFERSON			0		SURFACE	<100	50
STONE PLACER 7	CONTIN, URAN, WYO.	JEFFERSON			0		SURFACE	1,000 - 100,000	100
WRIGHT LEASE	COTTER COPPORATA	JEFFERSON			0		UNDERGRO	1,000 - 100,000	400
BLACKHAWK	BARNETT MINING C	LA PLATA			0		UNDERGRO	<100	0
LUCKY LEPRACON	SUN-SFT URAN, INC.	LA PLATA			0		SURFACE	<100	0
SHORTY LODGE	STEARNS, I.C.	LA PLATA	5	36 N	11.0 W	22	UNDERGRO	<100	50
BLACK HAWK MINE	CHEROKEE MINES C	LARIMER			0		UNDERGRO	100 - 1,000	150
BOULDER ROCK	MC LAUGHLIN, E.	LARIMER			0		SURFACE	<100	0
EUREKA	NEW WAR URANIUM	LARIMER			0		SURFACE	<100	50
RED HILL 1	ROBINSON, E.C.	LARIMER			0		UNDERGRO	100 - 1,000	50
WAKKETA LEASE	MORTON, ELBERT	LARIMER			0		UNDERGRO	<100	50
AJAX 1	AJAX MINING OIL	MESA	19	31 N	19.0 W	22	UNDERGRO	100 - 1,000	0
ARROWHEAD 4	ATLAS-AMAX	MESA	2	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
ARROWHEAD 4	ATLAS-AMAX	MESA	2	50 N	18.0 W	22	UNDERGRO	<100	50
ARROWHEAD 1 + 7	ATLAS-AMAX	MESA	10	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
ARROWHEAD 10	ATLAS-AMAX	MESA	11	50 N	18.0 W	22	UNDERGRO	100 - 1,000	50
ARROWHEAD 11	ATLAS-AMAX	MESA	10	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
ARROWHEAD 13	ATLAS-AMAX	MESA	10	50 N	18.0 W	22	UNDERGRO	<100	100
ARROWHEAD 14	ATLAS-AMAX	MESA	11	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
ARROWHEAD 17	ATLAS-AMAX	MESA	2	50 N	18.0 W	22	UNDERGRO	100 - 1,000	50
ARROWHEAD 18	ATLAS-AMAX	MESA	2	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
ARROWHEAD 19	ATLAS-AMAX	MESA	2	50 N	18.0 W	22	UNDERGRO	100 - 1,000	50
ARROWHEAD 20	ATLAS-AMAX	MESA	7	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
ARROWHEAD 25	ATLAS-AMAX	MESA	3	50 N	18.0 W	22	UNDERGRO	100 - 1,000	100
ARROWHEAD 27	ATLAS-AMAX	MESA	10	50 N	18.0 W	22	UNDERGRO	100 - 1,000	0
ARROWHEAD 28	ATLAS-AMAX	MESA	10	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
ARROWHEAD 29	ATLAS-AMAX	MESA	10	50 N	18.0 W	22	UNDERGRO	100 - 1,000	150
ARROWHEAD INC 1+	ATLAS-AMAX	MESA	2	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
ARROWHEAD INC 12	ATLAS-AMAX	MESA	3	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
ARROWHEAD INC 24	ATLAS-AMAX	MESA	11	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
ARROWHEAD INC 4	ATLAS-AMAX	MESA			0		UNDERGRO	1,000 - 100,000	50
AT-05-1-36	UNION CARBIDE CP	MESA	4	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
ATLAS-LOVE MESA	DUNKLE, DALE	MESA	6	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
AUSTIN + AUSTIN	WOODARD, CHARLES	MESA			0		UNDERGRO	100 - 1,000	250
BANCO 1	DAVIS, RALPH M.	MESA	17	51 N	18.0 W	22	SURFACE	<100	50
BELMONT 1 + 2	CARPICO, J.M.	MESA	3	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
BOSSIE GROUP	MC MANUS, JOHN U	MESA	36	50 N	18.0 W	22	SURFACE	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 11

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
BIG INDIAN LEASE	ARROWHEAD URAN.	MESA	19	50 N	18.0 W	22	SURFACE	<100	0
BIG HAVFRICK	KELLEY, DAN E.	MESA	20	50 N	18.0 W	22	UNDERGRO	<100	0
BIG MEVEY	ATLAS-AMAX	MESA	32	50 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BLACK MESA	WILLIAMS, GROVER	MESA			0		UNDERGRO	100 - 1,000	0
BLACK ROCK 2	BEHDER, E.L.	MESA	27	50 N	19.0 W	22	UNDERGRO	<100	0
BLACK STREAK	UNION CARBIDE	MESA	31	50 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
BLUE CRFCK	BLUE CREEK MNG.	MESA	19	50 N	17.0 W	22	SURFACE	<100	0
BLUE MESA VIEW	BLUE CREEK MININ	MESA	30	50 N	17.0 W	22	SURFACE	<100	0
BLUE RIBBON 1	ATLAS-AMAX	MESA	3	50 N	18.0 W	22	UNDERGRO	100 - 1,000	100
BLUE RIBBON 3	ATLAS-AMAX	MESA	10	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
BLUE RIBBON 17	ATLAS-AMAX	MESA	17	50 N	18.0 W	22	UNDERGRO	<100	0
BLUE RIBBON 32	ATLAS-AMAX	MESA	16	50 N	18.0 W	22	UNDERGRO	<100	0
BLUE RIBBON GRDU	ATLAS-AMAX	MESA	3	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
BLUEBIRD	ATLAS-AMAX	MESA	19	50 N	17.0 W	22	UNDERGRO	100 - 1,000	0
BLUEBIRD DUMP	ATLAS-CLIMAX	MESA	19	50 N	17.0 W	22	DUMPS	100 - 1,000	0
BONNIE	STEWART, WESLEY	MESA			0		SURFACE	<100	0
BUD 1	MADE, NAY	MESA			0		SURFACE	<100	0
BUICK	SCHUMACHER, J.I.	MESA	31	51 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
C-G-27 (BONNIE 1	U.S. GOVT.	MESA	13	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
CALAMITY MINEST	HENDRICKSON, W.B.H.	MESA	35	50 N	18.0 W	22	UNDERGRO	<100	0
CALAMITY MESA DU	ATLAS MINERALS	MESA	10	50 N	18.0 W	22	DUMPS	100 - 1,000	50
CALCO	HAMMICK-ORTMAYER	MESA	23	50 N	18.0 W	22	UNDERGRO	100 - 1,000	100
CAT TRACK	SHIPROCK LTD.	MESA			0		UNKNOWN	100 - 1,000	100
CAVE CANYON	ALJAY MINING & OIL	MESA	9	50 N	19.0 W	22	UNDERGRO	100 - 1,000	0
CHICO + C FRACT	GRAHAM, F.R.	MESA	36	51 N	19.0 W	22	UNDERGRO	100 - 1,000	0
CLIFF DELLER	ZANETT, FRED	MESA	22	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
CLIMAX	UNKNOWN CONTROL	MESA			0		UNDERGRO	<100	0
CLIMAX RESIDUE	UNION CARBIDE CP	MESA			0		MISC.-PB	1,000 - 100,000	0
CLIMAX YD CLEANU	KFOGH+SHUMWAY	MESA			0		MISC.-PB	1,000 - 100,000	0
COAL IDAN-CITATI	VAN URANIUM	MESA	6	50 N	18.0 W	22	SURFACE	<100	0
COTTONWOOD 3+5	UNION CARBIDE	MESA	22	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
COVE 1 ADIT	UNKNOWN	MESA	4	50 N	19.0 W	22	UNDERGRO	<100	0
CRESCENT	GRIFE + BAILEY	MESA	25	50 N	18.0 W	22	UNDERGRO	<100	0
CROWN NEST	ATLAS-AMAX	MESA	11	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
CUR (GATEWAY)	NICKERSON ALBERT	MESA	5	50 N	18.0 W	22	UNDERGRO	<100	50
DALILU-YELLOW-BIR	BURNSIDE, GEORGE	MESA	4	25 E	26.0 E	24	UNDERGRO	100 - 1,000	50
DEAL GROUP	DEAL MNG CO. INC.	MESA	4	49 N	17.0 W	22	SURFACE	<100	0
DEPRESSION 2+3	AMERICAN WST MTL	MESA	18	50 N	18.0 W	22	UNDERGRO	100 - 1,000	100
DEPRESSION 4 + 5	NUCLEAR FUELS IN	MESA	19	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
DEPRESSION 6	CONTINENTAL MATL	MESA	17	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
DEPRESSION GROUP	MATTERHORN MNG.	MESA	18	50 N	18.0 W	22	UNDERGRO	100 - 1,000	0
DRUM DUST	LATON BROS. DRUM	MESA			0		MISC.-PB	100 - 1,000	0
DURANGO 2	BENHAM + TUNNEL	MESA	34	51 N	19.0 W	22	UNDERGRO	<100	50
ELIZABETH 7-10	ATLAS-AMAX	MESA	4	49 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
EMERSON	ATLAS-AMAX	MESA	19	50 N	17.0 W	22	UNDERGRO	100 - 1,000	0
FLAT TOP	BURNETT, WILLIAM	MESA	17	50 N	18.0 W	22	UNDERGRO	<100	0
FOUNTAIN OF YOUT	MAYOUTH MINING	MESA	31	51 N	18.0 W	22	UNDERGRO	100 - 1,000	0
FRACTION	BEAVER MESA	MESA	12	50 N	18.0 W	22	UNDERGRO	100 - 1,000	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 12

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
GATEWAY TAILINGS	ALABASTER PRODUC	MESA	10	51 N	19.0 W	22	MISC.-PB	1,000 = 100,000	0
GILMORE LODGE	GATEWAY MNG+DEV	MESA	36	51 N	20.0 W	22	UNDERGRO	100 = 1,000	0
GLADYS 1	RIGGLE+HARTOUGH	MESA	19	51 N	18.0 W	22	UNDERGRO	<100	50
HANSON-NEGUS	ATLAS-APAX	MESA	17	50 N	17.0 W	22	UNDERGRO	1,000 = 100,000	150
HARVEY 1	FOSTER + FOSTER	MESA	28	51 N	18.0 W	22	UNDERGRO	<100	50
HARVEY-PICK+SHOV	BROWN, RAY	MESA	28	51 N	18.0 W	22	UNDERGRO	100 = 1,000	0
HOLE 24	UNKNOWN CONTROLR	MESA			0		SURFACE	<100	0
HOPE 1 TO 4	ATLAS+FOOTE	MESA	29	50 N	17.0 W	22	UNDERGRO	100 = 1,000	50
HUMDINGER	BROWN, RAY	MESA	5	50 N	18.0 W	22	UNDERGRO	<100	0
J.W. LEWIS	J.W. LEWIS	MESA			0		UNKNOWN	<100	300
J.W.L.FRACTION	UNION CARBIDE CP	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 = 100,000	350
JEAN 1 + 2	AMERICAN Leduc II	MESA			0		UNDERGRO	<100	0
JODY GROUP	BOND, JACK	MESA			0		UNDERGRO	1,000 = 100,000	50
JOE	E.E. LEWIS, INC.	MESA	31	51 N	18.0 W	22	UNDERGRO	1,000 = 100,000	100
JOHN BROWN	GATEWAY MNG+DEV	MESA	1	50 N	20.0 W	22	SURFACE	<100	0
JOHN BROWN 14 +	UNION CARBIDE CP	MESA			0		UNDERGRO	1,000 = 100,000	300
JOHNNY MAE 3	NEW IDRIA MNG+CH	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 = 100,000	250
JU DEE 1	IP+IN, WAYNE A.	MESA			0		SURFACE	<100	0
JUMBO 1	ATLAS-APAX	MESA	19	50 N	17.0 W	22	UNDERGRO	100 = 1,000	0
KANARADU 3	WARREN, R.V.	MESA	33	50 N	18.0 W	22	SURFACE	<100	0
KARNS INCLINE	UNION CARBIDE CP	MESA	12	50 N	20.0 W	22	UNDERGRO	1,000 = 100,000	300
KING SOLOMON	CANYON URAN CO I	MESA	24	48 N	18.0 W	22	UNDERGRO	100 = 1,000	50
KLONDIKE	BLUE CREK MININ	MESA	25	50 N	18.0 W	22	UNDERGRO	<100	0
LA PLAZA 1	CARNOTITE EXPL C	MESA	22	50 N	19.0 W	22	UNDERGRO	100 = 1,000	0
LA SAL	CEN PAC MNG CO	MESA			0		UNDERGRO	1,000 = 100,000	150
LA SAL 1 + 2	WOODARD, CHARLES	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 = 100,000	300
LA SAL 2,4,6+25	ATLAS-APAX	MESA			0		UNKNOWN	100 = 1,000	0
LA SAL 5 + 7	SHIPMAN MNG+EXPL	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 = 100,000	300
LA SAL GROUP	SHIPMAN MNG+EXPL	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 = 100,000	450
LA SALLE GROUP	FLANDERS MINING	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 = 100,000	150
LEE 1-6	HAMPICK +LINSOT	MESA	21	50 N	18.0 W	22	UNDERGRO	<100	0
LEGAL + LUCKY DA	NARDINE, PAUL	MESA	7	50 N	19.0 W	22	SURFACE	<100	0
LEVADA	MONTGOMERY, JACK	MESA	23	50 N	18.0 W	22	UNDERGRO	<100	0
LINERTY BELL 2	HALL, RIO B.	MESA			0		UNDERGRO	<100	0
LINCOLN	SHIPROCK, LTD	MESA	31	51 N	18.0 W	22	UNDERGRO	1,000 = 100,000	200
LITTLE JOHNNY	SHIPROCK LTD	MESA	31	51 N	18.0 W	22	UNDERGRO	1,000 = 100,000	50
LITTLE PAYERICK	WARREN, R.V.	MESA	21	50 N	18.0 W	22	UNDERGRO	<100	0
LOCUS 1,2+3	MITCHELL, C.E.	MESA	28	50 N	18.0 W	22	SURFACE	<100	0
LODE CLAIM	BUCAR MINES	MESA	33		0		UNDERGRO	<100	0
LOG CABIN	ZUFELT, LYNN	MESA	35	46 N	18.0 W	22	UNDERGRO	100 = 1,000	0
LONE PEAK	STRODE, EMORY	MESA	27	51 N	19.0 W	22	SURFACE	<100	0
LOOK OUT	KEELINE OIL+URAN	MESA	19	50 N	18.0 W	22	SURFACE	<100	50
LOST DUTCHMAN	PIONEER URAN INC	MESA	26	51 N	20.0 W	22	SURFACE	1,000 = 100,000	250
LUCKY BOY	UNION CARBIDE CP	MESA			0		UNDERGRO	1,000 = 100,000	50
LUCKY DAY	AJAX MINING +OIL	MESA	7	50 N	19.0 W	22	UNDERGRO	100 = 1,000	0
LUCKY HOLE	GULLETT, ORA R.	MESA			0		SURFACE	<100	0
LUCKY PINE 2	CORDREY+O. CONNOR	MESA	28	24 S	26.0 E	24	UNDERGRO	<100	0
LUCKY STRIKE	BUTLER, DEAN K.	MESA	32	50 N	18.0 W	22	UNDERGRO	<100	450

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 13

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
LUCKY STRIKE 7	BARONE + FOGEMAN	MESA	32	50 N	18.0 W	22	SURFACE	<100	0
LUMSDEN 2 + 6	UNION CARBIDE CP	MESA	36	51 N	20.0 W	22	UNDERGRO	1,000 - 100,000	200
MAMMOTH	SEYMOUR+MYERS	MESA	31	51 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
MAMMOTH-LINCOLN	MAMMOTH MINING	MESA	31	51 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
MARK 2	UNION CARBIDE CP	MESA	1	50 N	20.0 W	22	UNDERGRO	1,000 - 100,000	200
MARY 3	UNION CARBIDE CP	MESA	27	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	200
MAVERICK	JONES + BOWLES	MESA	3	50 N	18.0 W	22	SURFACE	<100	0
MAVERICK 6	ATLAS-AMAX	MESA	3	50 N	18.0 W	22	UNDERGRO	<100	100
MESA 5 (BEVR MSA	RALPH FOSTER+S	MESA			0		UNDERGRO	1,000 - 100,000	100
MESA 8	FOSTER + SONS	MESA	12	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
MESA CREEK	CHRISTMAS, W. MNG	MESA			0		UNDERGRO	<100	0
MILL CLFAY UP	ATLAS MINERALS	MESA			0		MISC.-PB	1,000 - 100,000	50
MILL SITE LODGE	PALMER, JACK	MESA			0		UNDERGRO	<100	0
MINERAL CHANNEL	ATLAS-AMAX	MESA	12	50 N	18.0 W	22	UNDERGRO	100 - 1,000	0
MINERAL CHANNEL	ATLAS-AMAX	MESA	12	50 N	18.0 W	22	UNDERGRO	<100	50
MINING LEASE 34	ROSPA + RUBY F	MESA	31	44 N	18.0 W	22	SURFACE	<100	0
MLB-C-G-26	RAJAH VNTR-GOVTL	MESA			0		UNDERGRO	100 - 1,000	100
MLB-C-G-27A	PIONR URAY GVTLS	MESA			0		UNKNOWN	1,000 - 100,000	400
MONROE 18	AMERICAN LEDUC U	MESA	36	50 N	18.0 W	22	UNDERGRO	100 - 1,000	50
MONTLEZUMA	NIELSON, HAROLD I.	MESA	10	50 N	19.0 W	22	UNDERGRO	<100	0
NEILSON	DULANEY MINING C	MESA	10	50 N	19.0 W	22	SURFACE	<100	0
NEILSON MOTHER D	DULANEY MINING C	MESA	10	50 N	19.0 W	22	UNDERGRO	<100	0
NEWHEISEL	WODDARD, CHARLES	MESA	31	51 N	19.0 W	22	UNDERGRO	1,000 - 100,000	250
OUTLAW ECONOMY	FOSTER + SONS	MESA	12	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
PAY LODGE	KELLEY, DAN E.	MESA	19	51 N	18.0 W	22	UNDERGRO	100 - 1,000	0
PAYDAY 1 THRU 7	BLACK CAT EXPL,C	MESA	2	50 N	19.0 W	22	SURFACE	<100	0
PAYROCK GROUP	DOYLE M K	MESA	17	51 N	18.0 W	22	UNDERGRO	100 - 1,000	50
PEACH 10 INC.1+2	AMERICAN LEDUC U	MESA	25	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
PIF FACE 1	CHAPIN, LOUIS	MESA	2	46 N	17.0 W	22	UNDERGRO	100 - 1,000	0
PPT,CONCFNTRATE	ALARASTER PRODUCE	MESA	10	51 N	19.0 W	22	MISC.-PB	<100	0
PROTECTOR	SCHUMACHER, J.I.	MESA	31	51 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
PURPLE HEART	PETITTI, JOHN J.	MESA			0		SURFACE	<100	0
RADIUM 7	ATLAS-AMAX	MESA	9	50 N	18.0 W	22	UNDERGRO	<100	50
RAE MARIE 3	LEECO GAS + OIL	MESA			0		UNDERGRO	100 - 1,000	200
RAE MARIE GROUP	L. S. DAWSON	MESA	33	24 S	26.0 E	24	UNDERGRO	1,000 - 100,000	200
RAINBOW	BROWN + WILLIAMS	MESA	18	50 N	18.0 W	22	UNDERGRO	100 - 1,000	0
RAINY DAY	FOSTER + SONS	MESA	35	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
RAJAH 1	UNION CARBIDE CP	MESA	36	51 N	20.0 W	22	UNDERGRO	100 - 1,000	550
RAJAH 11+ 63	UNION CARBIDE	MESA	35	51 N	20.0 W	22	UNDERGRO	1,000 - 100,000	550
RAJAH 72	UNION CARBIDE CP	MESA	36	51 N	20.0 W	22	UNDERGRO	100 - 1,000	450
RANCH VIEW	BLUE CREEK MINIM	MESA	30	50 N	17.0 W	22	SURFACE	<100	0
RAVEN 3	UNION CARBIDE	MESA	32	50 N	17.0 W	22	UNDERGRO	100 - 1,000	100
REMA	DEL RIO MNG CO.	MESA	19	51 N	18.0 W	22	UNDERGRO	100 - 1,000	150
RONNIE 2	BARNETT, GRANT T	MESA	12	50 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
RUDOT 1	BENHAM + TUNNEL	MESA	34	51 N	19.0 W	22	UNDERGRO	100 - 1,000	50
SALUTE 3	AMERICAN LEDUC U	MESA	25	50 N	18.0 W	22	UNDERGRO	<100	0
SAMPLE REJECT	SMITH, J. FARL	MESA			0		MISC.-PB	<100	0
SCOTT 2	HAYRICK + LINSBOT	MESA	27	50 N	18.0 W	22	SURFACE	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 14

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.
***** COLORADO (CONT'D) *****									
SURFACE	<100	0	SHELBY DEAN 2	STULLER + BOWEN	MESA	34	51 N	19.0 W	22
MISC.-PB	<100	0	SILVER MOON	BFCR, L.C.	MESA	10	1 N	3.0 W	22
UNDERGRO	1,000 = 100,000	0	SMALL SPOT	ZIMMERMAN, BEN	MESA	9	50 N	18.0 W	22
UNDERGRO	1,000 = 100,000	50	SNOW SHOE	FOSTER + SONS	MESA	12	50 N	18.0 W	22
UNDERGRO	100 = 1,000	0	SOLDIER BOY	PETRO NUCLEAR	MESA	11	47 N	20.0 W	22
UNDERGRO	1,000 = 100,000	100	SPRING	ATLAS-AMAX	MESA	12	50 N	18.0 W	22
SURFACE	<100	100	STORMY TREASURE	CHESAPEAKE+ COLO	MESA	35	50 N	18.0 W	22
SURFACE	<100	0	STRODE 1	STRODE, EMORY	MESA			0	
UNDERGRO	100 = 1,000	50	SUN	FOSTER + SONS	MESA	12	50 N	18.0 W	22
UNDERGRO	1,000 = 100,000	150	SUN SPOT-CLOUD 1	UNION CARBIDE CP	MESA	32	51 N	19.0 W	22
SURFACE	<100	0	SUPPLY 11	MONTGOMERY, JACK	MESA	35	50 N	18.0 W	22
SURFACE	<100	50	SUPPLY 14	AMERICAN LEADUC U	MESA	35	50 N	18.0 W	22
UNDERGRO	100 = 1,000	0	SURPRISE	MUVOKIV MNG.CO.	MESA			0	
SURFACE	<100	0	TENDERFOOT GROUP	GRAHAM, FRED	MESA	19	51 N	18.0 W	22
UNDERGRO	100 = 1,000	0	THE DUKE	UNION CARBIDE CP	MESA	32	51 N	19.0 W	22
MISC.-PB	<100	0	TODITLO	EPDA	MESA			0	
UNDERGRO	100 = 1,000	0	TROJAN 18 + 20	UNION CARBIDE CP	MESA	23	50 N	18.0 W	22
SURFACE	<100	0	VANADIUM KING 1	WRIGHT, WARREN E.	MESA	19	51 N	18.0 W	22
UNDERGRO	<100	0	VANADIUM KING 2	WRIGHT, WARREN E.	MESA	19	51 N	18.0 W	22
UNDERGRO	<100	250	VIRGEN	CLEGHORN, DOUGLAS	MESA			0	
UNDERGRO	100 = 1,000	100	WASP	CONTINENTAL MAIL	MESA	18	50 N	18.0 W	22
UNDERGRO	<100	0	WRAY MESA	UTIDA URAN.CO.	MESA	30	47 N	19.0 W	22
SURFACE	<100	0	YELLOW CAT	WRIGHT, WARREN E.	MESA	21	24 S	26.0 E	24
UNDERGRO	1,000 = 100,000	0	YELLOW JACKET 15	ATLAS-AMAX	MESA	3	50 N	18.0 W	22
UNDERGRO	100 = 1,000	0	YELLOW JACKET 9	ATLAS-AMAX	MESA	3	50 N	18.0 W	22
UNDERGRO	1,000 = 100,000	100	YELLOW JACKET 14	ATLAS-AMAX	MESA	3	50 N	18.0 W	22
SURFACE	100 = 1,000	100	BESSIE 9 + 10	UNION CARBIDE CP	MOFFAT			0	
UNDERGRO	100 = 1,000	50	BLUE MOUNTAINS 4	SKULL CREEK COAL	MOFFAT			0	
SURFACE	<100	0	BREARLINE	JONES, R.V.	MOFFAT			0	
SURFACE	<100	50	BUFFALO HEAD	BUFFALO HEAD MNG	MOFFAT			0	
SURFACE	1,000 = 100,000	50	BUTLER LEASE 1	CONOCO-PIONEER	MOFFAT	30	8 N	91.0 W	06
SURFACE	<100	0	CEDARS 1	HENDERSON, COUNTS	MOFFAT			0	
SURFACE	<100	0	CLAIM 1	BASHAW + ULSH	MOFFAT			0	
SURFACE	<100	0	CLETA GROUP	BESSELL + KAYE	MOFFAT			0	
SURFACE	100 = 1,000	50	DOC ARMOUR MINE	UNION CARBIDE CP	MOFFAT			0	
SURFACE	>100,000	50	GERIPUDE	UNION CARBIDE CP	MOFFAT	8	7 N	94.0 W	06
SURFACE	<100	0	GLORY BEE	ELLER + CALDWELL	MOFFAT			0	
SURFACE	1,000 = 100,000	50	JOHNSON M.C.B.	UNION CARBIDE CP	MOFFAT	9	7 N	94.0 W	06
UNDERGRO	1,000 = 100,000	50	LITTLE STAR	POWERS, LLOYD	MOFFAT	33	5 N	00.0 W	06
UNDERGRO	<100	0	LUCKY BOY	RANGER URAN.CO.	MOFFAT			0	
SURFACE	>100,000	200	RON-ROLLO	UNION CARBIDE CP	MOFFAT			0	
SURFACE	100 = 1,000	50	SEC.16, 64-94	TONY WILLIAMS CO	MOFFAT	16	6 N	94.0 W	06
SURFACE	100 = 1,000	50	SUGARLOAF	MRS.G.I.TOWER	MOFFAT			0	
SURFACE	100 = 1,000	50	THREE SISTERS	LEVKULICH, BILL	MOFFAT			0	
UNDERGRO	<100	100	BLUE EAGLE 1	YOUNG, ELFEE	MONTEZUMA			0	
SURFACE	<100	0	KARLE KAY	FOUR CORNERS OIL	MONTEZUMA			0	
UNDERGRO	100 = 1,000	50	ROBERTA JEAN	HOTZ, EARL	MONTEZUMA			0	
SURFACE	<100	50	SWALLOW 1	BLIGAR, GENE	MONTEZUMA			0	

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 15

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
VEACH	BRITTAIN + HALL	MONTEZUMA			0		SURFACE	<100	0
VIRGINIA AVE	BUCKEYE MINING C	MONTEZUMA			0		UNDERGRO	<100	0
1ST NATIONAL BAN	UNION CARBIDE CP	MONTEPOSE	21	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
2ND NATIONAL BAN	UNION CARBIDE CP	MONTEPOSE	28	47 N	17.0 W	22	UNDERGRO	100 - 1,000	0
30-30	HADDEN, T.A.	MONTEPOSE			0		UNDERGRO	1,000 - 100,000	0
45-90	UNION CARBIDE CP	MONTEPOSE	10	47 N	17.0 W	22	UNDERGRO	100 - 1,000	0
ABAJO 1-5	UNION CARBIDE CP	MONTEPOSE	19	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
ALL STARS EYEVIEW	UNION CARBIDE CP	MONTEPOSE	28	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
ALTA	MONOGRAM MINING	MONTEPOSE	23	48 N	18.0 W	22	UNDERGRO	100 - 1,000	50
ALTAR, CAPELLA, V	UNION CARBIDE CP	MONTEPOSE	13	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
AMERICAN EAGLE 4	WEBB RESOURCES	MONTEPOSE	3	45 N	19.0 W	22	UNDERGRO	1,000 - 100,000	50
AMERICAN EAGLE G	MAYFIELD, JFPRY	MONTEPOSE	10	45 N	19.0 W	22	UNDERGRO	1,000 - 100,000	150
ANCHOR	UNION CARBIDE CP	MONTEPOSE	3	48 N	18.0 W	22	UNDERGRO	100 - 1,000	0
ANGLE	GRAPPLICH SAM V	MONTEPOSE	1	47 N	20.0 W	22	UNDERGRO	<100	0
ANNA MAY 1	UNION CARBIDE CP	MONTEPOSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
ANNA MAY 1 DUMPS	UNION CARBIDE CP	MONTEPOSE	18	46 N	17.0 W	22	DUMPS	1,000 - 100,000	50
ANNEX	NEESMA, GLEN	MONTEPOSE	7	47 N	17.0 W	22	SURFACE	<100	0
ARCTURUS	UNION CARBIDE CP	MONTEPOSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
ARROWHEAD (URAVAN	UNION CARBIDE CP	MONTEPOSE	28	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
AUSTIN (DOLORES)	UNION CARBIDE CP	MONTEPOSE	20	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
AZTEC	FOOTE MINERALS	MONTEPOSE	33	46 N	19.0 W	22	UNDERGRO	100 - 1,000	50
AZTEC	UNION CARBIDE CP	MONTEPOSE	20	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
B P 1	DAVIS, RAY	MONTEPOSE			0		SURFACE	<100	50
BABE RUTH	FOOTE MINERALS	MONTEPOSE	14	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
BABY FAH	FOOTE MINERALS	MONTEPOSE	6	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BADGER	UNION CARBIDE CP	MONTEPOSE	19	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BADGER 1	UNION CARBIDE CP	MONTEPOSE	19	47 N	17.0 W	22	UNDERGRO	100 - 1,000	0
BADGER 2	UNION CARBIDE CP	MONTEPOSE	26	48 N	18.0 W	22	UNDERGRO	100 - 1,000	0
BADGER DUMP	UNION CARBIDE CP	MONTEPOSE	19	47 N	17.0 W	22	DUMPS	100 - 1,000	50
BADGER-CROWN PRY	MICRO COPPER	MONTEPOSE	26	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
BAGGER	UNION CARBIDE CO	MONTEPOSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BALL POINT	MONTGOMERY, JACK	MONTEPOSE	11	46 N	18.0 W	22	UNDERGRO	<100	0
BANNER	UNION CARBIDE CP	MONTEPOSE	12	46 N	18.0 W	22	UNDERGRO	<100	0
BEAVER	UNION CARBIDE CP	MONTEPOSE	32	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
BED ROCK	UNKNOWN CONTROLR	MONTEPOSE			0		SURFACE	<100	0
BENCH	LYLE FRANCIS	MONTEPOSE			0		SURFACE	<100	0
BERNARD	UNION CARBIDE CP	MONTEPOSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
BERTIES BEAUTY C	KRESS + FISCHER	MONTEPOSE	8	46 N	18.0 W	22	UNDERGRO	<100	0
BETTER B 7	MATERNHORN MNG.	MONTEPOSE	31	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
BETTY JEAN	MONOGRAM MINING	MONTEPOSE	22	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
BIG BULL	FOOTE MINERALS	MONTEPOSE	12	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
BIG DICK	UNION CARBIDE CP	MONTEPOSE	19	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
BIG MITT	UNION CARBIDE CP	MONTEPOSE	21	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BIG ROCK	SIMPLOT, J.R. CO	MONTEPOSE	33	46 N	18.0 W	22	UNDERGRO	100 - 1,000	0
BIG SHOT	MOSLANDER, GEORGE	MONTEPOSE	6	47 N	17.0 W	22	SURFACE	<100	0
BILL BADDY-LUCKYR	UNION CARBIDE CP	MONTEPOSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BIRTHDAY 1	GREEN, ARTHUR	MONTEPOSE	7	46 N	17.0 W	22	SURFACE	<100	0
BISHMARK	UNION CARBIDE CP	MONTEPOSE	29	47 N	17.0 W	22	UNDERGRO	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 16

WINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
BITTER CREEK	UNION CARBIDE	MONTROSE	12	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
BLACK DINAH	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BLACK DINAH DUMP	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	DUMPS	100 - 1,000	150
BLACK EAGLE	COCKRUM, F.V.	MONTROSE	4	45 N	16.0 W	22	SURFACE	<100	0
BLACK GNAT	MALEY MINING CO.	MONTROSE			0		SURFACE	<100	0
BLACK HAWK	WRIGHT, GEORGIA T	MONTROSE	27	49 N	18.0 W	22	SURFACE	<100	0
BLACK JACK	MAGIC URANIUM CO	MONTROSE	2	45 N	18.0 W	22	UNDERGRO	<100	50
BLACK PRINCE	BINDER, F. V.	MONTROSE	33	46 N	19.0 W	22	UNDERGRO	100 - 1,000	100
BLACK ROCK	UNION CARBIDE	MONTROSE	23	48 N	18.0 W	22	UNDERGRO	100 - 1,000	100
BLACK TOM	UNION CARBIDE CP	MONTROSE	20	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
BLACKFOOT RATTLE	SANDRA MARIE MIN	MONTROSE	10	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
BLISS	UNION CARBIDE C	MONTROSE			0		UNDERGRO	1,000 - 100,000	350
BLONDA	AYERS, EVERETT	MONTROSE			0		SURFACE	<100	0
BLONDY	GRIFE, W.F.	MONTROSE	26	48 N	18.0 W	22	UNDERGRO	<100	50
BLUE BELL	UNION CARBIDE CP	MONTROSE	35	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
BLUE BIRD	UNION CARBIDE CP	MONTROSE	25	48 N	18.0 W	22	UNDERGRO	100 - 1,000	100
BLUE BIRD DUMP	UNION CARBIDE CP	MONTROSE	25	48 N	18.0 W	22	DUMPS	100 - 1,000	50
BOB 6,7,8	UNION CARBIDE CP	MONTROSE			0		UNDERGRO	1,000 - 100,000	400
BOB 9	BELL, A.L. + B.L.	MONTROSE	5	45 N	17.0 W	22	SURFACE	<100	0
BOB CAT	UNION CARBIDE CP	MONTROSE	19	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BOB CAT	GRAPLICH SAM V	MONTROSE	1	47 N	20.0 W	22	SURFACE	<100	0
BONANZA	SECURITY URANIUM	MONTROSE			0		UNDERGRO	<100	0
BONITA 1	BONITA URANIUM C	MONTROSE	32	49 N	17.0 W	22	UNDERGRO	<100	0
BONACIN BETTY	JONES + THOMPSON	MONTROSE			0		SURFACE	<100	0
BPOORE 1	MAYFIELD, JERRY	MONTROSE	10	45 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
BROOMSTICK	FOOTE MINERALS	MONTROSE	11	45 N	18.0 W	22	UNDERGRO	<100	0
BROWN DERSY	FAFLEY, EMERY	MONTROSE			0		SURFACE	<100	0
BRUSHY BASIN	SCHUMACHER J. I.	MONTROSE	28	28 S	26.0 E	24	UNDERGRO	100 - 1,000	0
BUMBLES	COCHRAN URANIUM	MONTROSE			0		UNDERGRO	<100	0
BUCKEYE 4	SMITH-JENSEN+DAY	MONTROSE			0		SURFACE	<100	0
BUCKHORN 1	CURTIS, CLYDE	MONTROSE	21	48 N	18.0 W	22	SURFACE	<100	0
BUCKSHOT	UNION CARBIDE CP	MONTROSE	31	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
BUCKSKIN	JARMAN + HADDEN	MONTROSE	35	46 N	18.0 W	22	UNDERGRO	<100	250
BUCKSKIN GROUP	F V BINDER	MONTROSE	35	46 N	18.0 W	22	UNDERGRO	<100	100
BUTTERFLY	UNION CARBIDE	MONTROSE	10	47 N	17.0 W	22	UNDERGRO	100 - 1,000	0
BUTTERFLY	UNION CARBIDE	MONTROSE	14	47 N	20.0 W	22	UNDERGRO	1,000 - 100,000	100
C F C	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	100 - 1,000	150
CABIN VIEW	CAMDORSE URANIUM	MONTROSE	33	46 N	18.0 W	22	UNDERGRO	<100	0
CALVERT 2	UNION CARBIDE CP	MONTROSE	10	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
CAMEL (LEIGHTON)	FOOTE MINERALS	MONTROSE	11	45 N	18.0 W	22	UNDERGRO	100 - 1,000	200
CANOPUS	UNION CARBIDE CP	MONTROSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
CANYON 2	PATTERSON, PAT	MONTROSE			0		UNDERGRO	<100	0
CANYON VIEW	YACKEL, CAPL	MONTROSE	4	45 N	19.0 W	22	UNDERGRO	100 - 1,000	0
CARPATIA	UNION CARBIDE CP	MONTROSE	18	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
CARPENTER RIDGE	UNKNOWN CONTROL	MONTROSE			0		SURFACE	<100	0
CASHIN MILL	KARPISCH D.C.	MONTROSE	22	47 N	18.0 W	22	SURFACE	<100	0
CEDAR RIDGE	AYERS, EVERETT	MONTROSE	4	45 N	19.0 W	22	UNDERGRO	100 - 1,000	0
CHECKER	PATTERSON+ NEILS	MONTROSE	30	47 N	17.0 W	22	SURFACE	<100	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 17

MIKE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
CHESTERFIELD TPE	UNION CARBIDE CP	MONTROSE			0		UNDERGRO	1,000 - 100,000	0
CHILLI 5	UNION CARBIDE CP	MONTROSE	25	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
CHIPMUNK 1	AYERS, EVERETT	MONTROSE	4	45 N	19.0 W	22	UNDERGRO	<100	0
CHRISTIE	STOCKS, R.H., SISK	MONTROSE			0		SURFACE	<100	0
CLIFOWELLER DUMP	UNION CARBIDE CP	MONTROSE			0		DUMPS	100 - 1,000	0
CLUB 2	UNION CARBIDE CP	MONTROSE	20	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
CLUB GROUP	GLOSSER, RAYMOND	MONTROSE	32	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
CLUB GROUP	UNION CARBIDE	MONTROSE	29	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
COLUMBUS	FOOTE MINERALS	MONTROSE	2	46 N	17.0 W	22	UNDERGRO	<100	0
CONFUSION	SAV V. GRAMMICH	MONTROSE	12	47 N	20.0 W	22	UNDERGRO	100 - 1,000	0
COPPER JACK	MAKVEL MNG CO	MONTROSE	19	48 N	18.0 W	22	UNDERGRO	100 - 1,000	0
CORPORATION	UNION CARBIDE	MONTROSE	16	47 N	17.0 W	22	UNDERGRO	100 - 1,000	250
CORRECT	UNION CARBIDE CP	MONTROSE			0		UNDERGRO	<100	0
COTTONWOOD 1,2,3	PATTERSON, PAT	MONTROSE	33	46 N	19.0 W	22	UNDERGRO	100 - 1,000	0
COUGAR	DOOLEY T.T.	MONTROSE			0		SURFACE	<100	0
CRIPPLE CREEK	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
CRIPPLE CREEK DU	UNION CARBIDE CP	MONTROSE	21	47 N	17.0 W	22	DUMPS	1,000 - 100,000	100
CRIPPLE CRK 2 DU	UNION CARBIDE CP	MONTROSE	21	47 N	17.0 W	22	DUMPS	1,000 - 100,000	100
CUE BALL	UNION CARBIDE CP	MONTROSE	3	47 N	17.0 W	22	UNDERGRO	<100	0
D + D 3	ROGERS + HUNT	MONTROSE			0		UNDERGRO	<100	0
D + D 5	DOWELL H.L.	MONTROSE			0		UNDERGRO	<100	0
DADS	MICRO COPPER CO	MONTROSE	14	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
DAN PATCH	UNION CARBIDE CP	MONTROSE	35	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
DAWN	STORY, OREN	MONTROSE			0		SURFACE	<100	0
DEER, JULY, SLIM	UNION CARBIDE CP	MONTROSE	28	46 N	17.0 W	22	UNDERGRO	>100,000	650
DIANA	FOOTE MINERALS	MONTROSE	14	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	200
DOAGY 2-LAST DOL	UNION CARBIDE CP	MONTROSE	20	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
DOLORES 2	MAIRD+SNYDER MNG	MONTROSE	30	44 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
DOLORES MINE	UNION CARBIDE CP	MONTROSE	19	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
DONALD L DUMP	UNION CARBIDE CP	MONTROSE			0		DUMPS	100 - 1,000	0
DONNA K	FOOTE MINERALS	MONTROSE	31	47 N	16.0 W	22	UNDERGRO	1,000 - 100,000	200
DOROTHY	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
DOROTHY E.	ST REGIS URANIUM	MONTROSE	18	48 N	18.0 W	22	UNDERGRO	<100	0
DOUBLE JACK	KYLE, JAMES A.	MONTROSE	30	47 N	16.0 W	22	UNDERGRO	100 - 1,000	0
DUCHES 2 + 3	GRIFF W.E.	MONTROSE	34	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
DUSTY DUMP	UNION CARBIDE CP	MONTROSE			0		DUMPS	100 - 1,000	0
EAGLE ROCK 1	DOWELL H.L.	MONTROSE			0		UNDERGRO	<100	0
EDITH IRENE	STEWART, JAMES	MONTROSE	11	45 N	19.0 W	22	UNDERGRO	<100	50
EDNA MAE	JOHANNSEN E.J.	MONTROSE	21	45 N	18.0 W	22	UNDERGRO	100 - 1,000	0
EIGHT BALL	UNION CARBIDE CP	MONTROSE	3	47 N	17.0 W	22	UNDERGRO	100 - 1,000	0
EIGHT O CLOCK	FINCH, TOM	MONTROSE	33	46 N	18.0 W	22	SURFACE	<100	0
ELIZARETH ANN 1	HARDWICK + LOVLESS	MONTROSE			0		UNDERGRO	<100	0
EVENING STAR	PETRO NUCLEAR	MONTROSE	11	47 N	20.0 W	22	UNDERGRO	1,000 - 100,000	50
EXPECTANT 1	LONDON, ROBERT	MONTROSE	3	45 N	19.0 W	22	SURFACE	<100	0
FAERY QUEEN	UNION CARBIDE	MONTROSE	10	47 N	17.0 W	22	UNDERGRO	100 - 1,000	50
FARMER BOY	KINSEY + MALICK	MONTROSE			0		SURFACE	<100	0
FAULTLESS	MAYFIELD+VICKERS	MONTROSE	24	45 N	19.0 W	22	UNDERGRO	<100	0
FAWN SPRINGS 11	UNION CARBIDE CP	MONTROSE	6	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 18

WINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLOGRADO (CONT'D) *****									
FAWN SPRINGS 12	UNION CARBIDE CP	MONTROSE	31	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
FAWN SPRINGS 13	UNION CARBIDE CP	MONTROSE	6	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
FAWN SPRINGS 15	UNION CARBIDE CP	MONTROSE	6	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
FAWN SPRINGS 18	UNION CARBIDE CP	MONTROSE	6	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
FAWN SPRINGS 21	UNION CARBIDE CP	MONTROSE	31	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
FAWN SPRINGS 29	FOOTE MINERALS	MONTROSE	7	45 N	17.0 W	22	UNDERGRO	100 - 1,000	100
FAWN SPRINGS 3	UNION CARBIDE CP	MONTROSE	31	46 N	17.0 W	22	UNDERGRO	100 - 1,000	300
FAWN SPRINGS 30	UNION CARBIDE CP	MONTROSE	6	45 N	17.0 W	22	UNDERGRO	100 - 1,000	350
FAWN SPRINGS 4+1	UNION CARBIDE CP	MONTROSE	6	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
FAWN SPRINGS 5 T	UNION CARBIDE CP	MONTROSE	6	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
FIFTH NATIONAL B	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	100 - 1,000	150
FIREBIRD	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
FIPECRAKER	UNION CARBIDE CP	MONTROSE	35	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	300
FLAT TOP	SAN MIGUEL MINES	MONTROSE			0		SURFACE	<100	0
FLORENCE NELLIE	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
FOSSIL	FOOTE MINERALS	MONTROSE	7	45 N	17.0 W	22	UNDERGRO	100 - 1,000	50
FOURTH JULY + K	UNION CARBIDE CP	MONTROSE	24	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
FOURTH NAIL BANK	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	100 - 1,000	100
FOX	UNION CARBIDE CP	MONTROSE	30	48 N	17.0 W	22	UNDERGRO	100 - 1,000	150
FOX CISTERN	UNION CARBIDE CP	MONTROSE	19	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
FRACTION + FRAC	UNION CARBIDE CP	MONTROSE	19	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
GILBERT	UNION CARBIDE CP	MONTROSE	32	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
GNOME	UNION CARBIDE CP	MONTROSE	17	47 N	17.0 W	22	UNDERGRO	<100	0
GOLDEN EAGLE 14+	ALBANO, CLAIR	MONTROSE	9	45 N	19.0 W	22	UNDERGRO	100 - 1,000	150
GOOD HOPE PFD FO	FAGEN + FAGEN	MONTROSE	10	47 N	17.0 W	22	SURFACE	<100	0
GRAND DAD	MAL PAIS MNG.	MONTROSE			0		SURFACE	<100	50
GRANDVIEW	H.D.+JIM RUTT	MONTROSE			0		UNDERGRO	1,000 - 100,000	0
GRASS ROOTS	UNION CARBIDE CP	MONTROSE	34	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
GRASS ROOTS DUMP	UNION CARBIDE CP	MONTROSE	27	48 N	17.0 W	22	DUMPS	100 - 1,000	100
GRAY	UNION CARBIDE CP	MONTROSE	13	46 N	18.0 W	22	SURFACE	1,000 - 100,000	50
GRAY DUMP	UNION CARBIDE CP	MONTROSE	13	46 N	18.0 W	22	DUMPS	<100	0
GRAY FOX	BELL, PIRL W.	MONTROSE	11	47 N	17.0 W	22	UNDERGRO	100 - 1,000	0
GREAT WESTERN	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
GREAT WESTERN DU	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	DUMPS	100 - 1,000	50
GREENBACK	UNION CARBIDE CP	MONTROSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
GROUNDHOG	UNION CARBIDE CP	MONTROSE	17	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
GROUNDHOG	OLIVER + RUSS	MONTROSE	9	48 N	19.0 W	22	UNDERGRO	100 - 1,000	0
GYP LEASE	FOOTE MINERALS	MONTROSE	10	45 N	19.0 W	22	UNDERGRO	1,000 - 100,000	50
HAPPY	UNION CARBIDE CP	MONTROSE	21	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
HAPPY JOE	MILLER+BLACKBURN	MONTROSE	2	45 N	19.0 W	22	UNDERGRO	<100	0
HAPPY THOUGHT	UNION CARBIDE CP	MONTROSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
HARD LUCK	LONG, ARTHUR	MONTROSE			0		UNDERGRO	<100	0
HARDROCK	FOOTE MINERALS	MONTROSE	18	46 N	18.0 W	22	UNDERGRO	100 - 1,000	150
HAROLD	UNION CARBIDE CP	MONTROSE	32	46 N	17.0 W	22	UNDERGRO	100 - 1,000	150
HENRY CLAY	UNION CARBIDE CP	MONTROSE	29	4 N	1.7 W	22	UNDERGRO	1,000 - 100,000	50
HENRY CLAY DUMPS	UNION CARBIDE CP	MONTROSE	29	47 N	17.0 W	22	DUMPS	1,000 - 100,000	250
HIDDEN BASIN	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
HIGH BALL 5	SHUMWAY+ DADE	MONTROSE	26	46 N	18.0 W	22	UNDERGRO	100 - 1,000	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 19

MIKE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
HOMESTEAD	BALL, ALBERT	MONTROSE			0		SURFACE	<100	0
HONEYMOON	UNION CARRIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
HONEYMOON DUMPS	UNION CARRIDE CP	MONTROSE	20	47 N	17.0 W	22	DUMPS	1,000 - 100,000	50
HORSEHAIR 1	URRALRURU-HOLZ	MONTROSE	1	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
HORSEHAIR GROUP	GUIRE, HELEN W.	MONTROSE	2	45 N	18.0 W	22	UNDERGRO	100 - 1,000	0
HOT ROCK	MEYER, KARL	MONTROSE	12	46 N	17.0 W	22	SURFACE	<100	0
HOT SPOT	PRICE + LEDLOW	MONTROSE			0		UNDERGRO	<100	0
HOWLING COYOTE	DANVERS, DON	MONTROSE	33	47 N	19.0 W	22	UNDERGRO	<100	0
HUMMER	UNION CARRIDE CP	MONTROSE	21	46 N	17.0 W	22	UNDERGRO	>100,000	200
HUMMER DUMPS	UNION CARRIDE CP	MONTROSE	21	46 N	17.0 W	22	DUMPS	100 - 1,000	150
IENA	SCHUMACHER, J I	MONTROSE			0		UNDERGRO	<100	50
INDEX	LA RUE, O.C.	MONTROSE	2	48 N	18.0 W	22	UNDERGRO	100 - 1,000	0
IOLA	UNION CARRIDE CP	MONTROSE			0		UNDERGRO	100 - 1,000	0
IRENE	UNION CARRIDE CP	MONTROSE	29	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
ISLAND VIEW 1-7	CHAPMAN + FRANKS	MONTROSE	11	45 N	19.0 W	22	SURFACE	<100	100
J.B.GROUP	UNION CARRIDE CP	MONTROSE	3	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
J.J.	UNION CARRIDE CP	MONTROSE	26	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
JACK RABBIT	UNKNOWN CONTROLR	MONTROSE	11	45 N	18.0 W	22	UNDERGRO	<100	0
JACKPOT GROUP	BEE HIVE MINING	MONTROSE	2	48 N	20.0 W	22	SURFACE	<100	0
JCEP	PETERSON, FREDDIE	MONTROSE	18	47 N	17.0 W	22	SURFACE	<100	0
JITTENBUG	MARVEL MNG.CO.	MONTROSE	21	48 N	18.0 W	22	UNDERGRO	100 - 1,000	0
JO ANNE GROUP	CHERIGOS, HARRY P	MONTROSE	25	48 N	18.0 W	22	UNDERGRO	100 - 1,000	100
JOE	UNION CARRIDE CP	MONTROSE			0		UNDERGRO	1,000 - 100,000	200
JOE DANDY	UNION CARRIDE CP	MONTROSE	21	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
JOE DANDY DUMPS	UNION CARRIDE CP	MONTROSE	21	46 N	17.0 W	22	UNDERGRO	<100	150
JOE RIVERSIDE	UNION CARRIDE CP	MONTROSE	29	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
JOHN Z.	SMART, D.E.	MONTROSE	10	46 N	18.0 W	22	UNDERGRO	100 - 1,000	50
JOKER	PLATFAU URANIUM	MONTROSE	24	48 N	18.0 W	22	UNDERGRO	100 - 1,000	50
JOKER	BLACK HAWK URAN.	MONTROSE	10	45 N	19.0 W	22	UNDERGRO	100 - 1,000	50
JOKER	ELDER, FRANK	MONTROSE	34	46 N	18.0 W	22	UNDERGRO	100 - 1,000	0
JUDY ANN	ST REGIS URANIUM	MONTROSE	18	48 N	18.0 W	22	UNDERGRO	<100	50
JUMBO	BRINK, AL	MONTROSE	11	45 N	19.0 W	22	UNDERGRO	<100	0
JUNE BUG	AYERS, EVERETT	MONTROSE	30	28 S	26.0 E	24	UNDERGRO	<100	0
JUNGLE BASIN	FOOTE MINERALS	MONTROSE	35	45 N	17.0 W	22	SURFACE	<100	0
JUST RIGHT	MENDISCO, FELIX	MONTROSE	31	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
KING	E.E. LEWIS, INC.	MONTROSE			0		UNDERGRO	<100	50
KING OF LODS	UNION CARRIDE CP	MONTROSE	19	47 N	17.0 W	22	UNDERGRO	100 - 1,000	0
LARK 7 + 8	UNION CARRIDE CP	MONTROSE	11	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	250
LAST CHANCE 1	UNION CARRIDE CP	MONTROSE	9	48 N	19.0 W	22	UNDERGRO	1,000 - 100,000	50
LAST HOPE	DYNATEK ENTERPRISE	MONTROSE	22	48 N	18.0 W	22	UNDERGRO	<100	0
LAST LOAD	SNYDER, FRED JR.	MONTROSE	8	48 N	19.0 W	22	SURFACE	<100	0
LAZY THREE	D. + D. URAN + EXPL	MONTROSE	25	48 N	18.0 W	22	UNDERGRO	100 - 1,000	50
LEVI	UNION CARRIDE CP	MONTROSE	9	45 N	18.0 W	22	UNDERGRO	100 - 1,000	0
LITTLE BASIN	UNION CARRIDE CP	MONTROSE	27	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
LITTLE BUCKHORN	ATLAS-AMAX	MONTROSE	10	48 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
LITTLE CHIEF	MALEY, LEO	MONTROSE	13	47 N	18.0 W	22	SURFACE	<100	0
LITTLE DICK	UNION CARRIDE CP	MONTROSE	30	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
LITTLE DICK DUMP	UNION CARRIDE CP	MONTROSE	30	48 N	17.0 W	22	DUMPS	<100	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 20

MINING	TOTAL PRODUCTION	DEPTH	MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	METHOD	(TONS AS OF 01/01/79)	(FT.)
			***** COLORADO	(CONT'D)	*****							
			LITTLE JEWEL	FOOTE MINERALS	MONTROSE	3	45 N	19.0 W	22	UNDERGRO	100 - 1,000	0
			LITTLE JOE	MALEY LEO C	MONTROSE	14	48 N	18.0 W	22	SURFACE	<100	0
			LITTLE SLIP 1	HIEIT, MARION J.	MONTROSE			0		SURFACE	<100	0
			LO HIGH	SULLIVAN+MOORE+	MONTROSE			0		UNDERGRO	<100	0
			LOG CABIN	NICPO COPPER CO	MONTROSE	35	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
			LOMI	JOHANNSEN E.J.	MONTROSE	17	46 N	17.0 W	22	UNDERGRO	<100	0
			LONE CEDAR	GALYEAN, JAMES F	MONTROSE	34	46 N	17.0 W	22	UNDERGRO	<100	0
			LONE PINE	CECIL BUNKER	MONTROSE	5	48 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
			LONG PARK 1	UNION CARBIDE CP	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
			LONG PARK 10	UNION CARBIDE	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
			LONG PARK 10 DUM	UNION CARBIDE CP	MONTROSE			0		DUMPS	<100	0
			LONG PARK 11	UNION CARBIDE CP	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	100 - 1,000	0
			LONG PARK 12	UNION CARBIDE CP	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
			LONG PARK 2	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
			LONG PARK 3	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
			LONG PARK 4	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
			LONG PARK 5	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	100 - 1,000	100
			LONG PARK 6	UNION CARBIDE CP	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
			LONG PARK 6 DUM	UNION CARBIDE CP	MONTROSE	27	47 N	17.0 W	22	DUMPS	1,000 - 100,000	50
			LONG PARK 9	UNION CARBIDE CP	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	650
			LONG PARK GROUP	SUTHERLAND MNG.	MONTROSE			0		UNDERGRO	100 - 1,000	0
			LUCK DAY	COUCH + TRONE	MONTROSE	2	45 N	18.0 W	22	UNDERGRO	<100	0
			LUCKY BLUNDER	UNION CARBIDE CP	MONTROSE	18	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
			LUCKY DOG	UNION CARBIDE CP	MONTROSE	29	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
			LUCKY MAPX	RIDENOUR + ROSS	MONTROSE			0		UNDERGRO	<100	0
			LUCKY STRIKE	HEITZEL, LFE	MONTROSE	29	48 N	17.0 W	22	SURFACE	<100	50
			LUCKY STRIKE 4	WRIGHT, BILL	MONTROSE			0		SURFACE	<100	0
			LYNX	UNION CARBIDE CP	MONTROSE	29	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	450
			MAGGIE C	UNION CARBIDE	MONTROSE	21	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
			MAGGIE C DUM	UNION CARBIDE CP	MONTROSE			0		DUMPS	100 - 1,000	0
			MARGIE 2	FLANDERS MINING	MONTROSE	30	48 N	17.0 W	22	SURFACE	<100	0
			MARGIE GROUP	FOOTE MINERALS	MONTROSE	30	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
			MARJORIE ANN	PETRO NUCLEAR	MONTROSE	11	47 N	20.0 W	22	UNDERGRO	1,000 - 100,000	100
			MARTHA BELLE	UNION CARBIDE CP	MONTROSE	6	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	300
			MARY ANN 4-DOOROT	UNION CARBIDE CP	MONTROSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
			MARY JANE	E, JOHANNSEN	MONTROSE			0		UNKNOWN	100 - 1,000	0
			MAUDE	UNION CARBIDE	MONTROSE	14	47 N	20.0 W	22	UNDERGRO	100 - 1,000	50
			MAYBE DUMPS	UNION CARBIDE CP	MONTROSE	35	46 N	17.0 W	22	DUMPS	100 - 1,000	400
			MEDIA	UNION CARBIDE	MONTROSE	34	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
			MERRY CHRISTMAS	CHRISTMAS, M. MNG	MONTROSE	2	48 N	18.0 W	22	SURFACE	<100	0
			MERRY WIDOW	UNION CARBIDE	MONTROSE	10	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
			MESA	CARE, ORVAL	MONTROSE	33	46 N	19.0 W	22	UNDERGRO	1,000 - 100,000	50
			MESA 2	PATTERSON, PAT	MONTROSE	34	46 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
			MESA 3	HOPKINS + SMITH	MONTROSE	21	36 N	29.0 E	14	SURFACE	<100	0
			MIDAB	SMITH, ED	MONTROSE			0		SURFACE	<100	0
			MIKE 1	LAMBERT, JAMES R	MONTROSE	36	44 N	17.0 W	22	SURFACE	<100	0
			MILL 4	UNION CARBIDE CP	MONTROSE	4	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
			MINERAL PARK 2	ATLAS-AMAX	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 21

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
MINERAL PARK 3	ATLAS-AMAX	MONTROSE	27	47 N	17.0 W	22	UNDERGRO	<100	200
MINING LEASE 10	LA SALLE MINING	MONTROSE	5	47 N	17.0 W	22	UNDERGRO	>100,000	250
MINING LEASE 11	UNION CARBIDE CP	MONTROSE	5	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
MINING LEASE 13	BUNKER + CO	MONTROSE	31	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
MINING LEASE 14	UNION CARBIDE CP	MONTROSE	22	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	400
MINING LEASE 15	BRITO, BEN	MONTROSE	21	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
MINING LEASE 23	ALBANO, CLAIR	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
MINING LEASE 24	BARKLEY + CO	MONTROSE	5	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
MINING LEASE 37	BRATTUCK DEVM MG	MONTROSE	6	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	500
MINING LEASE 47	WORCESTER MINES	MONTROSE	24	48 N	18.0 W	22	UNDERGRO	>100,000	600
MLB-C-JD-7	COTTER-GOVTLEASE	MONTROSE			0		UNDERGRO	100 - 1,000	250
MLB-C-SR-10	TRIO IND CVTLE	MONTROSE			0		UNDERGRO	1,000 - 100,000	150
MLB-C-SR-15	COUGAR VENTURES	MONTROSE			0		UNDERGRO	1,000 - 100,000	400
MOORE	SEARS, RALPH	MONTROSE	33	46 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
MONOGRAH 12	MONOGRAH MINING	MONTROSE	27	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
MONOGRAH 5-FRM G	MONOGRAH MNG C	MONTROSE	22	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	200
MOONBEAM	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
MORN,STAR-MOONLI	E.E. LEWIS, INC.	MONTROSE	28	46 N	19.0 W	22	UNDERGRO	1,000 - 100,000	50
MORNING GLORY 2	TINTIC URANIUM	MONTROSE	28	47 N	19.0 W	22	UNDERGRO	<100	0
MOVIE STAR	UNION CARBIDE CP	MONTROSE	22	48 N	17.0 W	22	UNDERGRO	100 - 1,000	100
MUCKER	UNION CARBIDE CP	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
MUM	PIONEER UPAY INC	MONTROSE	36	47 N	17.0 W	24	UNDERGRO	100 - 1,000	200
MUSTARD	UNION CARBIDE CP	MONTROSE	30	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
NAT GROUP	TRANS MONID URAN	MONTROSE			0		SURFACE	<100	100
NATURITA 24	GRIPE, A.F.	MONTROSE	29	47 N	16.0 W	22	UNDERGRO	<100	50
NATURITA 4	GRIPE, ALLEN L.	MONTROSE	33	47 N	16.0 W	22	SURFACE	<100	50
NAVAJO	TEMPLETON, CLIFF	MONTROSE	19	45 N	18.0 W	22	UNDERGRO	<100	0
NAVAJO	BCHUMACHER, J I	MONTROSE	11	45 N	18.0 W	22	UNDERGRO	100 - 1,000	200
NEW CAMP BIRD	MONTGOMERY, JACK	MONTROSE	33	46 N	18.0 W	22	SURFACE	<100	0
NIL 2 DUMP	UNION CARBIDE CP	MONTROSE	26	46 N	17.0 W	22	DUMPS	100 - 1,000	400
NORTH STAR DUMP	UNION CARBIDE CP	MONTROSE	14	48 N	18.0 W	22	DUMPS	100 - 1,000	200
NORTH STAR UNAME	UNION CARBIDE	MONTROSE	14	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	200
NUCLA	UNION CARBIDE	MONTROSE	24	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
NUCLA	UNION CARBIDE CP	MONTROSE	24	48 N	18.0 W	22	UNDERGRO	100 - 1,000	50
OLD CRGW	CAMOOSE URANIUM	MONTROSE	4	45 N	18.0 W	22	UNDERGRO	<100	0
OLD GRANDAD	FOOTE MINERALS	MONTROSE	6	45 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
OLD QUAKER	UNION CARBIDE CP	MONTROSE	18	45 N	17.0 W	22	UNDERGRO	100 - 1,000	0
OLD SALT LICK	UNION CARBIDE CP	MONTROSE	29	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
OLD STAG I	ENSLEY+OSTROWSKI	MONTROSE			0		SURFACE	<100	0
OPERA BOX	UNION CARBIDE CP	MONTROSE	20	46 N	17.0 W	22	UNDERGRO	>100,000	250
OPHIR BLUEBIRD	UNION CARBIDE CP	MONTROSE	24	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
OPHIR DUMP	UNION CARBIDE CP	MONTROSE	24	48 N	18.0 W	22	DUMPS	1,000 - 100,000	50
OREGON	UNION CARBIDE CP	MONTROSE	11	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
OVERSIGHT	UNION CARBIDE CP	MONTROSE	21	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
PABLO 4 + 5	CLEGHORN+WASHBRN	MONTROSE			0		UNDERGRO	100 - 1,000	200
PAIN-OBNOXIOUS	UNION CARBIDE CP	MONTROSE	32	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
PARADOX 4,5+6	UNION CARBIDE CP	MONTROSE	15	46 N	17.0 W	22	UNDERGRO	>100,000	350
PARADOX BELLE	UNION CARBIDE CP	MONTROSE	30	47 N	17.0 W	22	UNDERGRO	<100	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 22

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
PARADOX C	UNION CARBIDE CP	MONTROSE	22	46 N	17.0 W	22	UNDERGRD	1,000 - 100,000	150
PARADOX D	UNION CARBIDE CP	MONTROSE	21	46 N	17.0 W	22	UNDERGRD	>100,000	250
PARADOX VIEW	BELL, GEORGE L.	MONTROSE	2	46 N	17.0 W	22	UNDERGRD	100 - 1,000	0
PATTERSON GROUP	HOLMAN, CLAY	MONTROSE	18	46 N	18.0 W	22	SURFACE	<100	0
PATTFERSON SEEP	WRIGHT, BILL	MONTROSE	18	46 N	18.0 W	22	SURFACE	<100	0
PATTY 4	MICRO COPPER CO	MONTROSE	34	46 N	18.0 W	22	UNDERGRD	100 - 1,000	0
PATTY 5	MICRO COPPER CO	MONTROSE	3	45 N	18.0 W	22	UNDERGRD	1,000 - 100,000	50
PAYDAY	MONOGRAM MINING	MONTROSE	23	48 N	19.0 W	22	UNDERGRD	1,000 - 100,000	50
PAYDAY	BLUEBIRD MINING	MONTROSE	34	45 N	18.0 W	22	UNDERGRD	<100	0
PEG LEG 2	UNION CARBIDE CP	MONTROSE	19	47 N	17.0 W	22	UNDERGRD	100 - 1,000	150
PINION-CEGAR GRO	UNKNOWN CONTRL	MONTROSE	24	18 N	19.0 W	22	SURFACE	<100	0
PLUTO-SATURN	UNION CARBIDE CP	MONTROSE	12	46 N	18.0 W	22	UNDERGRD	1,000 - 100,000	250
POINT-EMPIRE	MONOGRAM MINING	MONTROSE	23	48 N	19.0 W	22	UNDERGRD	1,000 - 100,000	50
POOCH + POOCH 1	BLUE BIRD MINING	MONTROSE	9	45 N	19.0 W	22	UNDERGRD	<100	0
POOP POY	STRODF, EMORY	MONTROSE	3	48 N	18.0 W	22	SURFACE	<100	0
PRAYER 8 + 9	UNION CARBIDE CP	MONTROSE	14	47 N	20.0 W	22	UNDERGRD	1,000 - 100,000	350
PRINCESS PAT	UNION CARBIDE CP	MONTROSE			0		UNDERGRD	<100	0
PROBABLE	ROBINSON, CLAUDE	MONTROSE	4	45 N	19.0 W	22	UNDERGRD	<100	50
PRODUCTION	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRD	1,000 - 100,000	100
PRODUCTION DUMPS	UNION CARBIDE CP	MONTROSE			0		DUMPS	<100	100
PROHIBITION	AYERS + AYERS	MONTROSE	8	48 N	18.0 W	22	UNDERGRD	100 - 1,000	0
QUAKREL	MICRO COPPER	MONTROSE	12	45 N	18.0 W	22	UNDERGRD	100 - 1,000	0
QUO VADIS	DOALL + ASSOC.	MONTROSE	3	48 N	19.0 W	22	SURFACE	<100	0
R.A.L. 1	GREAT WESTERN UR	MONTROSE	23	48 N	18.0 W	22	UNDERGRD	1,000 - 100,000	0
R.A.M.	UNION CARBIDE CP	MONTROSE	33	48 N	17.0 W	22	UNDERGRD	>100,000	150
R.A.M. DUMP	UNION CARBIDE CP	MONTROSE	33	48 N	17.0 W	22	DUMPS	1,000 - 100,000	50
RABBIT FOOT 2	UNION CARBIDE CP	MONTROSE	2	45 N	18.0 W	22	UNDERGRD	<100	0
RADAR-EARLY MORN	KRAUSF + MILLER	MONTROSE	24	45 N	19.0 W	22	SURFACE	<100	50
RADIUM CYCLE	BEE HIVE MINING	MONTROSE	8	48 N	19.0 W	22	UNDERGRD	1,000 - 100,000	0
RADIUM HILL 10	UNION CARBIDE	MONTROSE	10	45 N	18.0 W	22	UNDERGRD	1,000 - 100,000	350
RADIUM HILL 31	FOOTE MINERALS	MONTROSE	10	45 N	18.0 W	22	UNDERGRD	1,000 - 100,000	150
RADIUM HILL 50	UNION CARBIDE	MONTROSE			0		UNDERGRD	100 - 1,000	150
RADIUM HILL 7	FOOTE MINERALS	MONTROSE	16	45 N	18.0 W	22	UNDERGRD	1,000 - 100,000	400
RADIUM KING(URAY	UNION CARBIDE CP	MONTROSE	1	46 N	17.0 W	22	UNDERGRD	<100	0
RADIUM QUEEN 13	MAYVEL PEG CO	MONTROSE	20	48 N	18.0 W	22	UNDERGRD	1,000 - 100,000	50
RAINBOW	TOMPES, DAVID	MONTROSE	27	47 N	19.0 W	22	SURFACE	<100	0
RAJAH DUMP ONE	COL COL DRILLING	MONTROSE	7	46 N	19.0 W	22	UNDERGRD	<100	100
RAJAH-BIG CHIEF	FOOTE MINERALS	MONTROSE	6	48 N	19.0 W	22	UNDERGRD	1,000 - 100,000	0
RAMBLER	UNION CARBIDE CP	MONTROSE	33	48 N	17.0 W	22	UNDERGRD	100 - 1,000	350
RAMBLER DUMPS	UNION CARBIDE CP	MONTROSE	33	48 N	17.0 W	22	DUMPS	1,000 - 100,000	50
RATX	HIDDEN SPLENDOR	MONTROSE	7	47 N	17.0 W	22	SURFACE	<100	0
RATTLER 1	BRUCE, FERRIS	MONTROSE			0		UNDERGRD	<100	0
RATTLESLAKE 2	CRANDELL, MARION	MONTROSE	33	45 N	18.0 W	22	SURFACE	<100	0
RATTLESLAKE-DAVI	B + L MNG.	MONTROSE	2	47 N	20.0 W	22	UNDERGRD	1,000 - 100,000	100
RATTLESLAKE-TURNO	YEADON + KEMPF	MONTROSE	3	47 N	17.0 W	22	UNDERGRD	1,000 - 100,000	50
RAVEN	UNION CARBIDE CP	MONTROSE	30	48 N	17.0 W	22	UNDERGRD	1,000 - 100,000	50
RED BEDS	RADIUM HILL URAN	MONTROSE	23	48 N	19.0 W	22	UNDERGRD	<100	0
RED BIRD 1 + 2	OLIVER + BUSS	MONTROSE	9	48 N	19.0 W	22	UNDERGRD	1,000 - 100,000	150

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 23

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
RED BIRD 20	MCGHEE, L.	MONTROSE	5	48 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
RED COB	UNION CARBIDE CP	MONTROSE	2	48 N	18.0 W	22	UNDERGRO	<100	0
RED HEAD 1	DOYLE M.K.	MONTROSE			0		SURFACE	<100	0
RED ROCK 2	MICRO COPPER	MONTROSE	29	48 N	18.0 W	22	UNDERGRO	100 - 1,000	150
RED ROCK 5	MICRO COPPER	MONTROSE	28	48 N	18.0 W	22	UNDERGRO	100 - 1,000	250
RED SOX-YANKEES	UNION CARBIDE	MONTROSE	23	48 N	16.0 W	22	UNDERGRO	1,000 - 100,000	250
REDBIRD-YELLOWBI	CLAXTON, W.D., + ASO	MONTROSE	33	46 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
REMAINT 1	SOUTHWESTERN OIL	MONTROSE	7	48 N	18.0 W	22	SURFACE	<100	150
RENEGADE GROUP	CLEGHORN+WASHBURN	MONTROSE	12	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
REPUBLICAN DUMP	UNION CARBIDE CP	MONTROSE			0		DUMPS	1,000 - 100,000	0
REPUBLICAN-DUSTY	UNION CARBIDE	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
RES. BLOCK 2 NL, 4	GARDNER + BROWN	MONTROSE	33	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
RIGEL	UNION CARBIDE CP	MONTROSE	13	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
RIM CLAIMS	GRIPE W.E.	MONTROSE	23	48 N	18.0 W	22	UNDERGRO	<100	0
RIMROCK BLUES 15+1	UNION CARBIDE CP	MONTROSE	36	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
RIMROCK	MICRO COPPER CO	MONTROSE	15	48 N	19.0 W	22	UNDERGRO	100 - 1,000	0
RIMROCK BLUES 20	UNION CARBIDE CP	MONTROSE			0		UNDERGRO	100 - 1,000	100
RIMROCK BLUES 5	UNION CARBIDE CP	MONTROSE	9	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
RIMROCK BLUES 6+	UNION CARBIDE CP	MONTROSE	1	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
RIMROCK BLUES 6E	UNION CARBIDE CP	MONTROSE	1	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
RIMROCK BLUES 9	UNION CARBIDE CP	MONTROSE	1	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
ROCK RAVEN	UNION CARBIDE CP	MONTROSE	35	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
RODDY 1	ALBERT PELL	MONTROSE			0		UNDERGRO	<100	0
ROSEBUD	FOOTE MINERALS	MONTROSE	13	48 N	18.0 W	22	UNDERGRO	<100	0
ROYAL OAK	UNION CARBIDE CP	MONTROSE	10	45 N	18.0 W	22	UNDERGRO	<100	0
RUBADALE	AMARILLO MNG	MONTROSE	28	47 N	17.0 W	22	UNDERGRO	100 - 1,000	50
RUSTY 5	BEERS, SIDNEY J.	MONTROSE	6	47 N	17.0 W	22	SURFACE	<100	0
SAM	UNION CARBIDE CP	MONTROSE	36	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
SANDY	UNION CARBIDE CP	MONTROSE	20	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
SAUCER BASIN GPD	NEESHAN, GLEN	MONTROSE	6	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
SCHOOL YARM	FOOTE MINERALS	MONTROSE	2	45 N	18.0 W	22	UNDERGRO	100 - 1,000	50
SEC. 20, 46V-17E N	ATOMIC ENERGY CO	MONTROSE	20	46 N	17.0 W	22	UNDERGRO	<100	250
SEGO LILY 1	DANVENS, DON	MONTROSE	32	47 N	19.0 W	22	UNDERGRO	<100	0
SHADOW	MONTGOMERY, W.W.	MONTROSE	20	48 N	18.0 W	22	SURFACE	<100	0
SHADOW ROCK	BLEAK, CHARLES	MONTROSE	1	48 N	19.0 W	22	UNDERGRO	100 - 1,000	0
SHAMROCK	UNION CARBIDE CP	MONTROSE	29	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
SHAMROCK	FOOTE MINERALS	MONTROSE	35	46 N	18.0 W	22	UNDERGRO	<100	50
SHARKEY	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
SHOOTING STAR	DOWELL, MC ILVIR	MONTROSE	25	48 N	18.0 W	22	SURFACE	<100	50
SHRIVER	UNION CARBIDE CP	MONTROSE			0		UNDERGRO	100 - 1,000	0
SLIM CHANCE	SCHARF, T.J.	MONTROSE			0		UNDERGRO	<100	0
SMOXY	CLYDE URAN. CORP.	MONTROSE	8	47 N	17.0 W	22	SURFACE	<100	150
SOCKET	MONTGOMERY, W.W.	MONTROSE	1	48 N	19.0 W	22	SURFACE	<100	0
SPINNER OF LATER	NICHOLS, CARL L.	MONTROSE			0		UNDERGRO	100 - 1,000	0
SPHINX	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
SPHINX DUMP	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	DUMPS	100 - 1,000	50
ST PATRICK	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
STAR 13 + 14	UNION CARBIDE C	MONTROSE			0		UNDERGRO	1,000 - 100,000	250

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 24

MINES NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
STAR 3 + 4	UNION CARBIDE CP	MONTROSE	28	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
STAR 3 DUMP	UNION CARBIDE CP	MONTROSE	28	48 N	17.0 W	22	DUMPS	<100	150
STAR 5 + 6	UNION CARBIDE CP	MONTROSE	28	48 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
STARLIGHT	SIMPLOT, J.R. CO	MONTROSE	33	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
STARLIGHT 1	F A SITTON	MONTROSE	33	46 N	18.0 W	22	UNDERGRO	100 - 1,000	0
STARLIGHT 2	SIMPLOT, J.R. CO	MONTROSE	33	46 N	18.0 W	22	UNDERGRO	<100	0
STARLIGHT 4	SIMPLOT, J.R. CO	MONTROSE	32	46 N	18.0 W	22	SURFACE	<100	0
STARLIGHT 8	SIMPLOT, J.R. CO	MONTROSE	33	46 N	18.0 W	22	UNDERGRO	100 - 1,000	0
STEER 1-8	FOOTE MINERALS	MONTROSE	14	4 N	1.8 W	22	UNDERGRO	1,000 - 100,000	100
STRAIGHT ARROW	GRINA, RUSSELL D	MONTROSE	3	46 N	17.0 W	22	UNDERGRO	<100	0
SUMNER	UNION CARBIDE CP	MONTROSE	2	47 N	20.0 W	22	SURFACE	<100	0
SUNFLOWER 2	UNION CARBIDE CP	MONTROSE	18	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	200
SUNNYSIDE	UNION CARBIDE CP	MONTROSE	24	46 N	18.0 W	22	UNDERGRO	<100	0
SUNRISE GROUP	MICRO COPPER	MONTROSE	34	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
SUNSET	CONSOLIDATED URA	MONTROSE	32	46 N	18.0 W	22	UNDERGRO	100 - 1,000	0
SURPRISE	FOOTE MINERALS	MONTROSE	4	47 N	17.0 W	22	UNDERGRO	<100	0
SWINDLER DUMP	UNION CARBIDE CP	MONTROSE			0		DUMPS	<100	0
SYLVEYS POCKET	PATTENSON, PAT	MONTROSE			0		SURFACE	<100	0
TANGO	UNION CARBIDE CO	MONTROSE			0		UNDERGRO	1,000 - 100,000	100
TEAPOT DOME 2 +	UNION CARBIDE CP	MONTROSE			0		UNDERGRO	1,000 - 100,000	100
TEEPER POLE	BLUE CREEK MINIA	MONTROSE	36	50 N	18.0 W	22	SURFACE	<100	0
THREE JACKS	UNION CARBIDE	MONTROSE	14	47 N	20.0 W	22	UNDERGRO	1,000 - 100,000	100
THREE MUSKETEERS	HOIZ, EARL	MONTROSE	8	48 N	18.0 W	22	UNDERGRO	<100	50
THUNDERBOLT	UNION CARBIDE CP	MONTROSE	23	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
TNT 2	UNION CARBIDE	MONTROSE	21	47 N	17.0 W	22	UNDERGRO	100 - 1,000	100
TNT 3	UNION CARBIDE	MONTROSE	21	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
TOD HIGH	PIONEER URAN INC	MONTROSE	36	47 N	20.0 W	22	UNDERGRO	1,000 - 100,000	100
TOP NOTCH	COX, MARION	MONTROSE			0		SURFACE	<100	0
TOWN HOUSE	UNION CARBIDE CP	MONTROSE	20	44 N	17.0 W	22	UNDERGRO	1,000 - 100,000	4950
TOWNSITE(1)	UNION CARBIDE	MONTROSE			0		UNDERGRO	1,000 - 100,000	100
TRAMP 2	UNION CARBIDE CP	MONTROSE	6	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
TRAMP DUMPS	UNION CARBIDE CP	MONTROSE	6	47 N	17.0 W	22	DUMPS	1,000 - 100,000	50
TRIANGULATION	UNION CARBIDE CP	MONTROSE	34	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	100
TRIPPOD	UNION CARBIDE CP	MONTROSE	20	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	250
TRIPPOD LOW GRADE	UNION CARBIDE CP	MONTROSE			0		LOW GRADE	1,000 - 100,000	50
TRUSCOTT	UNION CARBIDE CP	MONTROSE	28	48 N	17.0 W	22	UNDERGRO	100 - 1,000	50
TWILIGHT	UNION CARBIDE CP	MONTROSE	2	48 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
TWIN SISTERS	MARVEL MNG CO	MONTROSE	19	48 N	18.0 W	22	UNDERGRO	100 - 1,000	0
TWO BITS	MAUPIN, ED	MONTROSE	34	46 N	18.0 W	22	UNDERGRO	<100	150
TWO SHOVEL	UNION CARBIDE CP	MONTROSE	18	47 N	17.0 W	22	UNDERGRO	<100	0
U.S. GRANT	UNION CARBIDE CP	MONTROSE	8	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	150
UNCLE SAM (URAVAN)	UNION CARBIDE CP	MONTROSE	13	48 N	18.0 W	22	UNDERGRO	100 - 1,000	350
URANIUM GIRL INC	URANIUM GIRL INC	MONTROSE	14	47 N	20.0 W	22	UNDERGRO	1,000 - 100,000	250
URANUS-DOROTHY J	UNION CARBIDE CP	MONTROSE	18	46 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
URAVAN 2	SCHUMACHER, J.I.	MONTROSE	4	47 N	17.0 W	22	UNDERGRO	1,000 - 100,000	50
UREKA	MC GEHEE, J.P. + L	MONTROSE	10	48 N	19.0 W	22	UNDERGRO	100 - 1,000	0
VADEN VIEW	FOOTE MINERALS	MONTROSE	13	46 N	18.0 W	22	UNDERGRO	1,000 - 100,000	200
VALENTINE(MS-202)	UNION CARBIDE CP	MONTROSE	24	47 N	18.0 W	22	UNDERGRO	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 25

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	NEPID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
VALLEY VIEW	E L LEWIS, INC	MONTROSE	8	48 N	19,0 W	22	UNDERGRO	1,000 = 100,000	0
VALLEY VIEW-4, ST	UNION CARBIDE CP	MONTROSE	20	47 N	17,0 W	22	UNDERGRO	1,000 = 100,000	50
VAN	UNION CARBIDE	MONTROSE	12	46 N	17,0 W	22	UNDERGRO	100 = 1,000	0
VANABLEND 47	SILVER STATE URA	MONTROSE	29	48 N	16,0 W	22	SURFACE	<100	100
VANADITE	UNION CARBIDE	MONTROSE	29	47 N	17,0 W	22	UNDERGRO	1,000 = 100,000	0
VANADIUM KING	UNION CARBIDE	MONTROSE	22	47 N	17,0 W	22	UNDERGRO	1,000 = 100,000	350
VENTURE LODGE	RICE + WILLIAMS	MONTROSE	31	46 N	17,0 W	22	SURFACE	<100	0
VICTORY 2	KELLY, MARK	MONTROSE			0		SURFACE	<100	50
VIRGIN 3	CLEGHORN, DOUGLAS	MONTROSE	22	47 N	17,0 W	22	UNDERGRO	1,000 = 100,000	350
VISTA GRANDE	BAH V GRAMMICH	MONTROSE	2	47 W	20,0 W	22	UNDERGRO	<100	0
VONNIE 5	SCOUTER URANIUM	MONTROSE			0		SURFACE	<100	0
WATCHMAN	FOOTE MINERALS	MONTROSE	34	46 N	19,0 W	22	UNDERGRO	<100	100
WATERLOO	MARVEL MNG CO	MONTROSE	20	48 N	18,0 W	22	SURFACE	<100	0
WEDGE 1	UNION CARBIDE	MONTROSE	10	47 N	17,0 W	22	UNDERGRO	1,000 = 100,000	250
WEDGE-PI WARREN	UNION CARBIDE	MONTROSE	2	47 N	20,0 W	22	UNDERGRO	1,000 = 100,000	100
WEDNESDAY + THUR	UNION CARBIDE CP	MONTROSE	22	47 N	17,0 W	22	UNDERGRO	1,000 = 100,000	600
WEST	MAGIC URANIUM CO	MONTROSE	35	46 N	18,0 W	22	UNDERGRO	<100	250
WEST LODGE	WILLIAMS, D.L.	MONTROSE			0		UNDERGRO	100 = 1,000	150
WEST MARTHA BELL	UNION CARBIDE CP	MONTROSE	31	49 N	17,0 W	22	UNDERGRO	1,000 = 100,000	200
WHITE COX	COLE+MITCHELL+WM	MONTROSE	1	45 N	20,0 W	22	UNDERGRO	<100	0
WHITE CROW	FOSTER, LYMAN	MONTROSE	3	45 N	19,0 W	22	SURFACE	<100	50
WHITNEY	UNION CARBIDE CP	MONTROSE	35	47 N	17,0 W	22	UNDERGRO	1,000 = 100,000	400
WILD CAT 2	BURNETT, WILLIAM	MONTROSE	28	48 N	18,0 W	22	SURFACE	<100	0
WILD CAT 8	MICRO COPPER	MONTROSE	27	49 W	18,0 W	22	UNDERGRO	100 = 1,000	0
WILD HORSE-COLOR	UNION CARBIDE CP	MONTROSE	11	48 N	18,0 W	22	UNDERGRO	1,000 = 100,000	100
WILDCAT 3	MICRO COPPER	MONTROSE	28	48 N	18,0 W	22	UNDERGRO	100 = 1,000	0
WINDY DAY	UNION CARBIDE	MONTROSE	3	47 N	17,0 W	22	UNDERGRO	100 = 1,000	50
WOODCHUCK	MOAB TREASURY UP	MONTROSE	23	48 N	19,0 W	22	SURFACE	<100	0
WOODWARD	WELLS ENGLISH+ST	MONTROSE	1	45 N	18,0 W	22	SURFACE	<100	0
WRIGHT	UNION CARBIDE CP	MONTROSE	27	48 N	17,0 W	22	UNDERGRO	1,000 = 100,000	150
YELLOW JACKET	UNION CARBIDE CP	MONTROSE	33	48 N	17,0 W	22	UNDERGRO	1,000 = 100,000	0
ZEBRA	UNION CARBIDE CP	MONTROSE	31	46 N	17,0 W	22	UNDERGRO	100 = 1,000	0
ZELLA GROUP	JOHNSON, HUGO W.	MONTROSE	16	48 N	18,0 W	22	UNDERGRO	100 = 1,000	100
GEN DANDY 3	MARTIN QUAYLE BR	PARK			0		UNDERGRO	100 = 1,000	50
LADY ELY 1	ARMINE, JOHN	PARK			0		UNDERGRO	<100	50
LAST CHANCE	WARD + FEELEY	PARK			0		SURFACE	<100	50
LUCKY JM 1,2+3	FOUR STAR EXPL.C	PARK	26	10 S	75,0 W	06	UNDERGRO	100 = 1,000	50
MAC GEORGE 4	YELLOW QUEEN URA	PARK			0		UNDERGRO	100 = 1,000	50
REDSKIN MINES	REDSKIN MINES IN	PARK			0		UNDERGRO	<100	50
SHIRLEY MAE	GADSDIS MINING	PARK	16	11 S	78,0 W	06	SURFACE	100 = 1,000	0
FRYING PAN GROUP	ASPEN MINING CO.	PITKIN			0		UNDERGRO	<100	100
EVERY RANCH	CLIFF + CREEK UR	PUEBLO	6	18 S	66,0 W	06	UNDERGRO	1,000 = 100,000	100
ALLEN	E. + H. LEASING CO	RIO BLANCO			0		SURFACE	<100	0
BROWN 2	DEVEREAUX BROS.	RIO BLANCO			0		UNDERGRO	100 = 1,000	200
BROWN 5	DEVERFAUX BROS.	RIO BLANCO			0		SURFACE	<100	200
MURRELL 1,2+3	LYLE FRANCIS	RIO BLANCO			0		SURFACE	1,000 = 100,000	150
BURRELL 5	MINIER, DOROTHY	RIO BLANCO			0		SURFACE	100 = 1,000	100
BUTTERFLY GROUP	JRJ MNG.	RIO BLANCO			0		UNDERGRO	1,000 = 100,000	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 26

WINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
CHRIS	PHILLIPS, CLAYTON	RIO BLANCO			0		SURFACE	<100	0
COAL CREEK 1	DEVEREAUX BROS.	RIO BLANCO			0		UNDERGRO	1,000 - 100,000	100
COLUMBINE 1	DEVEREAUX BROS.	RIO BLANCO			0		UNDERGRO	100 - 1,000	50
EVENING STAR	DEEM TRAIL MINER	RIO BLANCO			0		UNDERGRO	<100	100
FRYING PAN 1	PAYTON, NORMAN	RIO BLANCO			0		SURFACE	<100	0
LAST DAY	MC ALESTER FUELS	RIO BLANCO			0		UNDERGRO	1,000 - 100,000	0
LUCIL 106	JONES W.S.	RIO BLANCO			0		SURFACE	<100	0
M.+G.	IOWA URANIUM CO	RIO BLANCO			0		SURFACE	<100	0
MARVINE VIEW 10	DEVEREAUX BROS.	RIO BLANCO			0		UNDERGRO	100 - 1,000	0
MIDNIGHT GROUP	HARRP, HARRY H. JR	RIO BLANCO			0		UNDERGRO	1,000 - 100,000	100
MIDNIGHT NH	HARRP, HARRY H. JR	RIO BLANCO			0		UNDERGRO	1,000 - 100,000	350
NAOMI ANN	MC ALESTER FUELS	RIO BLANCO			0		UNDERGRO	100 - 1,000	50
RIO BLANCO	RIO BLANCO CORP	RIO BLANCO			0		UNDERGRO	<100	150
S.+G. 4	MARCY EXPL. + MNG	RIO BLANCO			0		UNDERGRO	<100	100
SHYLO GROUP	DEVEREAUX BROS.	RIO BLANCO			0		UNDERGRO	1,000 - 100,000	100
ST. LUE	SMITH, LOUISE	RIO BLANCO			0		SURFACE	<100	0
T.B.	TUNGSTEN URANIUM	RIO BLANCO			0		UNDERGRO	<100	0
TWIN STAR 500	TWIN STAR MINING	RIO BLANCO			0		SURFACE	<100	0
UTE GROUP	MINIER, CORROTHY	RIO BLANCO			0		UNDERGRO	<100	50
WINDY POINT 1	GENTRY W.W.	RIO BLANCO			0		UNDERGRO	100 - 1,000	0
BEGINNERS LUCK 3	MIKE MIZOKAMI	SAGUACHE			0		UNDERGRO	<100	50
MORCAT LODGE	PHIPPS, O.F.	SAGUACHE			0		UNDERGRO	<100	50
RONITA GROUP	D + J URANIUM	SAGUACHE			0		UNDERGRO	100 - 1,000	100
LA RUE 2	CANADIAN MOLLY	SAGUACHE	30	47 N	2.0 E	22	UNDERGRO	<100	50
LITTLE INDIAN 36	HOMESTEAK MNG CO	SAGUACHE		48 N	6.0 E	22	UNDERGRO	1,000 - 100,000	100
LOOKOUT 22	MONARCH EXPL.CO.	SAGUACHE	27	48 N	6.0 E	22	UNDERGRO	100 - 1,000	50
LOS OCHOS	HOMESTEAK MNG CO	SAGUACHE	33	48 N	2.0 E	22	UNDERGRO	>100,000	350
MARSHALL PASS 5	UNCOMPAGNE EXPL	SAGUACHE		48 N	6.0 E	22	SURFACE	<100	0
MERCURY 1	COL TEX URANIUM	SAGUACHE			0		UNDERGRO	<100	50
MOCKING RIDG	GARDNER, JOE	SAGUACHE			0		UNDERGRO	<100	50
PAM LODGE	CALIF, UTAH PETRO	SAGUACHE			0		SURFACE	<100	0
RAM + RAM 1	PIKE MINING CO.	SAGUACHE			0		SURFACE	<100	50
SEC. 3 NH NEQ T-	GUNNISON MINING	SAGUACHE	3	47 N	2.0 E	22	SURFACE	1,000 - 100,000	100
ELK PARK MINE	GADDIS MINING	SAN JUAN	18	40 N	7.0 W	22	UNDERGRO	<100	50
GRAYSILL	FOOTE MINERALS	SAN JUAN	28	40 N	9.0 W	22	UNDERGRO	1,000 - 100,000	0
ADA RELL	HARMON J.J.	SAN MIGUEL	16	43 N	19.0 W	22	UNDERGRO	<100	0
ALCHEMIST	FUJII K + HALL R	SAN MIGUEL	4	43 N	10.0 W	22	UNDERGRO	100 - 1,000	0
APRIL	B + M MINING CO	SAN MIGUEL			0		UNDERGRO	1,000 - 100,000	0
AVA JAY	FRITZ ERICKSON M	SAN MIGUEL	25	44 N	20.0 W	22	UNDERGRO	1,000 - 100,000	100
BABE 1-4	WILLIAMS, RAY L.	SAN MIGUEL			0		UNDERGRO	<100	50
BACHELOR	HERRING, JAMES	SAN MIGUEL	11	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	300
BAUD EAGLE	SPENCER MINING C	SAN MIGUEL	30	44 N	16.0 W	22	UNDERGRO	1,000 - 100,000	50
BAY MULE	UNKNOWN CONTROLR	SAN MIGUEL			0		SURFACE	<100	0
BEAN 10	ATLAS-FOOTE	SAN MIGUEL	32	44 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
BEAN 15, 16, + 17	ATLAS-FOOTE	SAN MIGUEL	31	44 N	19.0 W	22	UNDERGRO	<100	100
BEAN 2 + 3	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
BEAN 4 + 5	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
BEAN 6	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	100 - 1,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DCE, GRAND JUNCTION, COLORADO

PAGE 27

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.
***** COLORADO (CONT'D) *****									
UNDERGRD	1,000 - 100,000	100	BEAN PATCH	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22
UNDERGRD	1,000 - 100,000	0	BEAR CREEK	FOOTE MINERALS	SAN MIGUEL	4	42 N	10.0 W	22
SURFACE	<100	0	BETTY JANE 2	UNKNOWN CONTROL	SAN MIGUEL			0	
SURFACE	<100	0	BIG BUCK 1	CLOSSON, DANIEL S	SAN MIGUEL	26	44 N	20.0 W	22
UNDERGRD	100 - 1,000	0	RIG CHIEF	UNION CARBIDE CP	SAN MIGUEL	29	43 N	18.0 W	22
UNDERGRD	100 - 1,000	0	BIG MEDICINE	EMTESS OIL & UPA	SAN MIGUEL	23	45 N	19.0 W	22
SURFACE	<100	0	BIG 3	DUNCAN & SANCHEZ	SAN MIGUEL	32	44 N	19.0 W	22
UNDERGRD	<100	50	BLACK BIRD	ATLAS-FOOTE	SAN MIGUEL	8	43 N	19.0 W	22
UNDERGRD	100 - 1,000	0	BLACK FOX	DUNCAN R L MININ	SAN MIGUEL	5	43 N	18.0 W	22
UNDERGRD	1,000 - 100,000	100	BLACK JACK	ATLAS-FOOTE	SAN MIGUEL	28	43 N	19.0 W	22
SURFACE	<100	0	BLACK KING 5	BACHELOR CORP.	SAN MIGUEL			0	
UNDERGRD	1,000 - 100,000	150	BLK SPID.-RED AN	C.L. STEWART	SAN MIGUEL	30	43 N	19.0 W	22
UNDERGRD	100 - 1,000	0	BLUE MOON	DAVIS & GOFORTH	SAN MIGUEL	16	45 N	18.0 W	22
SURFACE	<100	0	BLUEBIRD	DULANEY MNG.	SAN MIGUEL	24	43 N	20.0 W	22
UNDERGRD	100 - 1,000	0	BLUFF	UNION CARBIDE CP	SAN MIGUEL	35	45 N	18.0 W	22
SURFACE	<100	150	BRETTON & MRCOT	BRETTON & MRCOT	SAN MIGUEL			0	
UNDERGRD	<100	0	BROWN MULE	BLAIR, FLOYD	SAN MIGUEL	7	43 N	16.0 W	22
UNDERGRD	100 - 1,000	0	BUCKHORN	BROWN, E.R.	SAN MIGUEL	34	43 N	18.0 W	22
UNDERGRD	100 - 1,000	0	BUGWINE	BALORA, HERMAN M.	SAN MIGUEL	31	45 N	19.0 W	22
UNDERGRD	<100	0	BULL MOOSE	THOMPSON, T.E.	SAN MIGUEL	32	43 N	19.0 W	22
UNDERGRD	100 - 1,000	850	BULL SHARK GROUP	URANIUM PROCESSE	SAN MIGUEL	12	42 N	18.0 W	22
UNDERGRD	1,000 - 100,000	50	BURRO POINT	SCHUMACHER, J I	SAN MIGUEL	16	45 N	18.0 W	22
UNKNOWN	1,000 - 100,000	0	CANYON	KEOGH & SHUMWAY	SAN MIGUEL			0	
UNDERGRD	100 - 1,000	100	CANYON VIEW	DUNCAN R L MININ	SAN MIGUEL	5	43 N	18.0 W	22
SURFACE	<100	0	CAPE MAIRS	BARRETT & BROWN	SAN MIGUEL			0	
SURFACE	<100	0	CEDAR RIDGE GROU	BILL MINING CO	SAN MIGUEL			0	
UNDERGRD	100 - 1,000	0	CHARLES T. 1	CANFIELD, ARTHUR	SAN MIGUEL	10	43 N	19.0 W	22
UNDERGRD	1,000 - 100,000	0	CHARLES T. 2A	CANFIELD, ARTHUR	SAN MIGUEL			0	
UNDERGRD	<100	50	CHAPLOTTE 1	LLOYD, PERLE D.	SAN MIGUEL			0	
UNDERGRD	100 - 1,000	0	CHESTA	NEW DEAL MNG CO	SAN MIGUEL	26	45 N	18.0 W	22
UNDERGRD	1,000 - 100,000	200	CHIEF 1-3	FRITZ ERICKSON P	SAN MIGUEL			0	
SURFACE	<100	0	CHINMESE	WILLIAMS+MC GEHE	SAN MIGUEL			0	
UNDERGRD	100 - 1,000	0	CHIPMUNK	PETRO NUCLEAR	SAN MIGUEL	16	45 N	18.0 W	22
UNDERGRD	100 - 1,000	200	CHIPMUNK 1	AZTEC MINING CO.	SAN MIGUEL	35	44 N	19.0 W	22
SURFACE	<100	0	CLEAR CREEK	UNKNOWN CONTROL	SAN MIGUEL			0	
UNDERGRD	<100	0	CLEAR VIEW	LYENS, W.E.	SAN MIGUEL	6	42 N	17.0 W	22
UNDERGRD	100 - 1,000	0	CLIFF DWELLER	SKALLA, A.F.	SAN MIGUEL	9	44 N	17.0 W	22
SURFACE	<100	0	COLORADO CAT	SHYDER, LEE	SAN MIGUEL	15	44 N	19.0 W	22
UNDERGRD	1,000 - 100,000	150	CONE 1-6	ATLAS-FOOTE	SAN MIGUEL	30	44 N	19.0 W	22
UNDERGRD	<100	0	COWHAND 2	BOLDRA, HERMAN M.	SAN MIGUEL	31	45 N	19.0 W	22
UNDERGRD	<100	0	CRUCIBLE	NIELSON, ELMER	SAN MIGUEL	7	43 N	10.0 W	22
UNDERGRD	<100	0	CUP 1	RISENHOOVER, MATH	SAN MIGUEL			0	
UNDERGRD	100 - 1,000	0	CURTIS	SAN JUAN LEASING	SAN MIGUEL	28	43 N	18.0 W	22
SURFACE	<100	0	CUSCO	LUBBOCK MNG.CO.	SAN MIGUEL			0	
SURFACE	<100	0	CYCLE 3	SEARS, RALPH	SAN MIGUEL			0	
UNDERGRD	1,000 - 100,000	250	DELUXE & MASTER	BITTON & BITTON	SAN MIGUEL			0	
DUMPS	1,000 - 100,000	50	DEREMO DUMPS	UNION CARBIDE CP	SAN MIGUEL			0	
UNDERGRD	<100	0	DICKIE 1 & 3	B.E.K. CORPORATIO	SAN MIGUEL			0	

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 28

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.
***** COLORADO (CONT'D) *****									
UNDERGRO	<100	0	DOLORES RIVER	FOOTE MINERALS	SAN MIGUEL			0--	--
UNDERGRO	1,000 - 100,000	250	DOLORES RIVER	DOLORES RIVER BM	SAN MIGUEL			0	
UNDERGRO	<100	0	DONALD HILL	UNION CARBIDE CP	SAN MIGUEL	15	44 N	18.0 W	22
UNDERGRO	<100	0	DONEGAN LEASE	WELSON + LANGTON	SAN MIGUEL	8	43 N	10.0 W	22
UNDERGRO	100 - 1,000	0	DRAGON	ALMONT MINES INC	SAN MIGUEL			0	
UNDERGRO	<100	0	DUNCAN	UNION CARBIDE CP	SAN MIGUEL	26	45 N	19.0 W	22
UNDERGRO	<100	0	DURANGO + L ANIM	OLIVER BROS	SAN MIGUEL	33	44 N	19.0 W	22
UNDERGRO	1,000 - 100,000	0	EARLY MORN GROUP	D.C. BUNKER MINE	SAN MIGUEL	24	44 N	17.0 W	22
UNDERGRO	<100	0	ECLIPSE	UNION CARBIDE CP	SAN MIGUEL	10	44 N	18.0 W	22
SURFACE	<100	0	EMPIRE GROUP	ORTHMEYER MINING	SAN MIGUEL			0	
SURFACE	<100	0	FAIR VIEW	UNKNOWN CONTROLP	SAN MIGUEL			0	
UNDERGRO	1,000 - 100,000	200	FALL CREEK GROUP	FOOTE MINERALS	SAN MIGUEL	7	43 N	10.0 W	22
UNDERGRO	1,000 - 100,000	100	FIREFLY 3	E.E. LEVINS, INC.	SAN MIGUEL			0	
UNDERGRO	<100	50	FIVE POINTS	FIVE POINTS URAN	SAN MIGUEL			0	
UNDERGRO	100 - 1,000	0	FLORENCE	HAMILTON, VERL	SAN MIGUEL			0	
UNDERGRO	<100	100	FOX GROUP	BURWELL MINING	SAN MIGUEL			0	
UNDERGRO	100 - 1,000	100	FOX GROUP	MILLING JACK	SAN MIGUEL			0	
UNDERGRO	100 - 1,000	0	FRACTION	DULANEY MINING C	SAN MIGUEL	8	43 N	19.0 W	22
UNDERGRO	100 - 1,000	0	FRACTION 1	DULANEY MNG	SAN MIGUEL	10	43 N	19.0 W	22
UNDERGRO	1,000 - 100,000	50	FRANCE	FRITZ ERICKSON M	SAN MIGUEL	20	43 N	19.0 W	22
UNDERGRO	<100	50	FRANCES	UNION CARBIDE	SAN MIGUEL			0	
UNDERGRO	100 - 1,000	0	FRANKLIN 1 + 2	TORRES, DAVID	SAN MIGUEL	33	46 N	19.0 W	22
UNDERGRO	100 - 1,000	0	FRAZIER	BENNETT + ROSE	SAN MIGUEL	24	43 N	11.0 W	22
UNDERGRO	1,000 - 100,000	50	FRENCHY 2	ORTHMEYER MINING	SAN MIGUEL	29	43 N	19.0 W	22
UNDERGRO	1,000 - 100,000	50	FULL MOON GROUP	UNION CARBIDE C	SAN MIGUEL	15	43 N	18.0 W	22
SURFACE	<100	0	G.M.D. 1	DOVELL H.L.	SAN MIGUEL			0	
SURFACE	<100	0	GAP	REED, EDGAR J.	SAN MIGUEL	16	43 N	16.0 W	22
UNDERGRO	100 - 1,000	100	GERALD T.	MATTERHORN MNG.	SAN MIGUEL	19	44 N	18.0 W	22
UNDERGRO	100 - 1,000	0	GIANT	UNION CARBIDE CP	SAN MIGUEL	24	45 N	19.0 W	22
UNDERGRO	100 - 1,000	0	GLEN 27	BOSKY, HENRY	SAN MIGUEL	28	44 N	18.0 W	22
SURFACE	<100	0	GOFORTH HOMESTEAD	DAVIS MINING CO	SAN MIGUEL	15	43 N	19.0 W	22
UNDERGRO	100 - 1,000	0	GOLDEN ROD 1	FAHRION + BAGRILL	SAN MIGUEL	11	43 N	19.0 W	22
UNDERGRO	1,000 - 100,000	50	GOLDEN ROD 2	ATOMIC ENRGY CO	SAN MIGUEL	14	43 N	19.0 W	22
UNDERGRO	1,000 - 100,000	100	GOLDEN ROD 4	JOHNSON + HARGRAVE	SAN MIGUEL	14	43 N	19.0 W	22
UNDERGRO	1,000 - 100,000	150	GOPHER	SCHUMACHER, J I	SAN MIGUEL	21	45 N	18.0 W	22
UNDERGRO	100 - 1,000	150	GOVERNOR MINE	CAPITOL SLABOARD	SAN MIGUEL	29	43 N	19.0 W	22
UNDERGRO	1,000 - 100,000	200	GRASS FLAT	BRUNWAY + DADF	SAN MIGUEL	32	44 N	18.0 W	22
UNDERGRO	<100	0	GRASSY HILL	COPPER OIL + MNG	SAN MIGUEL	24	45 N	19.0 W	22
SURFACE	<100	0	GREEN ARROW	UNKNOWN CONTROLP	SAN MIGUEL			0	
UNDERGRO	1,000 - 100,000	300	GROUND HOG	PICKENS, CHARLES	SAN MIGUEL	21	45 N	18.0 W	22
DUMPS	1,000 - 100,000	50	GROUND HOG DUMP	FOUR COMPYRS OIL	SAN MIGUEL	21	45 N	18.0 W	22
SURFACE	<100	0	GRUB STAKE	BYODEN, C.F. + SONS	SAN MIGUEL	23	44 N	19.0 W	22
UNDERGRO	1,000 - 100,000	0	GYP SUM HOMESTEAD	BELL, PIPL W.	SAN MIGUEL	33	45 N	18.0 W	22
UNDERGRO	<100	0	HALLOWEEN	NICHOLS + CRECELI	SAN MIGUEL			0	
UNDERGRO	100 - 1,000	150	HAPPY JACK	FOOTE MINERALS	SAN MIGUEL	4	45 N	19.0 W	22
UNDERGRO	1,000 - 100,000	100	HAWK-FRANKIE	ATLAS-FOOTE	SAN MIGUEL	16	43 N	19.0 W	22
UNDERGRO	100 - 1,000	0	HAYMAKER-SUNSET	UNION CARBIDE CP	SAN MIGUEL	29	45 N	18.0 W	22
UNDERGRO	100 - 1,000	0	HAZEL	DAVIS, TRENTON	SAN MIGUEL			0	

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 29

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
HOGBACK	DAVIS, TRENTON	SAN MIGUEL	19	43 N	19.0 W	22	UNDERGRO	100 - 1,000	50
HORSESHOE 1	BROOKS MINERALS	SAN MIGUEL	6	42 N	17.0 W	22	UNDERGRO	100 - 1,000	0
HORSESHOE 2	BROOKS MINERALS	SAN MIGUEL	6	42 N	17.0 W	22	UNDERGRO	100 - 1,000	150
HORSESHOE 3	BROOKS MINERALS	SAN MIGUEL	6	42 N	17.0 W	22	UNDERGRO	100 - 1,000	150
HORSESHOE 4	BROOKS MINERALS	SAN MIGUEL	6	42 N	17.0 W	22	SURFACE	<100	0
HORSESHOE 5	BROOKS MINERALS	SAN MIGUEL	6	42 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
HORSESHOE 6	BROOKS MINERALS	SAN MIGUEL	6	42 N	17.0 W	22	UNDERGRO	100 - 1,000	150
HORSESHOE 7	CENTURY MNG+DEVE	SAN MIGUEL	6	42 N	17.0 W	22	UNDERGRO	100 - 1,000	150
HORSESHOE REND 1	CENTURY MNG+DEVE	SAN MIGUEL			0		UNDERGRO	1,000 - 100,000	150
HOT DRILL 11	UNITED URAN CORP	SAN MIGUEL	11	42 N	18.0 W	22	UNDERGRO	<100	850
HOT SHOT	SCHUMACHER, J I	SAN MIGUEL	16	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
HOT SPOT	AYERS, EVERETT	SAN MIGUEL			0		UNDERGRO	<100	0
HOYMAN LEASE	DAWSON, L.S.	SAN MIGUEL			0		UNDERGRO	1,000 - 100,000	800
I.V.	DULANEY MINING C	SAN MIGUEL	15	43 N	19.0 W	22	UNDERGRO	<100	0
INDEPENDENCE	GOWFF + GUPWELL	SAN MIGUEL	17	43 N	19.0 W	22	SURFACE	<100	0
INSPIRATION 1	LEECO GAS + OIL	SAN MIGUEL	33	45 N	18.0 W	22	UNDERGRO	100 - 1,000	0
INSPIRATION 15	SITTON F A	SAN MIGUEL	33	45 N	18.0 W	22	SURFACE	<100	0
J.J. 2	SMITH + PETERS	SAN MIGUEL			0		SURFACE	<100	0
J.V. EAVENSON LSE	ALABADO MNG.CO,	SAN MIGUEL	27	43 N	19.0 W	22	UNDERGRO	<100	0
JACK KNIFE 3	SCHUMACHER, J I	SAN MIGUEL	16	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
JACK-O-LANTERN	DALE DILLON	SAN MIGUEL	32	44 N	18.0 W	22	SURFACE	<100	0
JACKIE L.	DULANEY MINING C	SAN MIGUEL	8	43 N	19.0 W	22	UNDERGRO	<100	0
JACKIE WALLS 3	FRITZ EPICKSON M	SAN MIGUEL	36	44 N	20.0 W	22	UNDERGRO	1,000 - 100,000	50
JACKKNIFE 3 W	SCHUMACHER J I	SAN MIGUEL	16	45 N	18.0 W	22	UNDERGRO	100 - 1,000	150
JACYPOT GROUP	BROWN + WRIGHT	SAN MIGUEL	16	44 N	17.0 W	22	UNDERGRO	100 - 1,000	0
JIM 2	DAVIS + GOFORTH	SAN MIGUEL	11	42 N	18.0 W	22	UNDERGRO	100 - 1,000	0
JOE BUSH GROUP	THOMPSON, E.	SAN MIGUEL	26	43 N	19.0 W	22	UNDERGRO	100 - 1,000	0
JOE DANDY	NEILSON, C.	SAN MIGUEL	8	43 N	10.0 W	22	UNDERGRO	100 - 1,000	0
JUPITER	DUNCAN WALTER	SAN MIGUEL	32	43 N	19.0 W	22	UNDERGRO	100 - 1,000	150
KATE MEYERS	BLIXT, ALFRED	SAN MIGUEL			0		UNDERGRO	<100	0
KATIE	DALPEZ, MARIO	SAN MIGUEL	24	43 N	19.0 W	22	UNDERGRO	<100	50
KEYSTONE	FAHPTON +SAGRILL	SAN MIGUEL	14	43 N	19.0 W	22	UNDERGRO	100 - 1,000	0
KLONDIKE	LAKALUCRE MINES	SAN MIGUEL	1	43 N	17.0 W	22	SURFACE	<100	50
LA SALLE	ATLAS-FOOTE	SAN MIGUEL	30	44 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
LAST CHANCE	GREENRIVER OIL +	SAN MIGUEL	23	44 N	19.0 W	22	UNDERGRO	<100	100
LAST HOPE	JIM BUTT	SAN MIGUEL	24	43 N	20.0 W	22	UNDERGRO	100 - 1,000	0
LATRICIA	DOYLE M A	SAN MIGUEL	20	43 N	19.0 W	22	UNDERGRO	100 - 1,000	200
LEE MOCK	KANHOLZ, ROY	SAN MIGUEL	1	43 N	19.0 W	22	SURFACE	<100	0
LEOPARD VANADIUM	BENNETT +CORNETT	SAN MIGUEL	26	44 N	11.0 W	22	UNDERGRO	100 - 1,000	0
LIBERTY BELL	MALICK, ELI	SAN MIGUEL			0		SURFACE	<100	0
LITTLE CHIEF	BALORA, HERMAN M.	SAN MIGUEL			0		SURFACE	<100	0
LITTLE HELFA	GOFORTH, WM.	SAN MIGUEL	18	43 N	18.0 W	22	UNDERGRO	<100	0
LITTLE MARIE	THOMPSON+GRIBBLE	SAN MIGUEL	23	44 N	19.0 W	22	SURFACE	<100	0
LITTLE MAX	UNION CARBIDE CP	SAN MIGUEL	28	43 N	18.0 W	22	UNDERGRO	100 - 1,000	0
LITTLE ROY	SNYDER, C.F.+SONS	SAN MIGUEL	28	43 N	19.0 W	22	UNDERGRO	<100	0
LIZZIE G	C NEILSON	SAN MIGUEL	8	43 N	10.0 W	22	UNDERGRO	100 - 1,000	0
LOVE PEAK 1-3	PIONEER URAN INC	SAN MIGUEL	32	43 N	18.0 W	22	UNDERGRO	100 - 1,000	100
LOVE STAR GROUP	J + J MNG.	SAN MIGUEL	4	43 N	19.0 W	22	UNDERGRO	100 - 1,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 30

WINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
LONESOME 14	PIONEER URAN INC	SAN MIGUEL			0		UNDERGRO	1,000 - 100,000	0
LONG RIDGE+L.R.	UNION CARBIDE CP	SAN MIGUEL	23	44 N	17,0 W	22	UNDERGRO	1,000 - 100,000	0
LOOKOUT	PETRO NUCLEAR	SAN MIGUEL	23	44 N	19,0 W	22	UNDERGRO	1,000 - 100,000	0
LOST	SEARS, RALPH	SAN MIGUEL	33	45 N	18,0 W	22	UNDERGRO	100 - 1,000	0
LOST BROTHERS	CROWN URANIUM CO	SAN MIGUEL	27	45 N	18,0 W	22	UNDERGRO	100 - 1,000	50
LOST DOG	JAYFIELD, JERRY	SAN MIGUEL	16	43 N	19,0 W	22	UNDERGRO	100 - 1,000	50
LSE. 875, S. 16, NW4	ST. OF COLORADO	SAN MIGUEL			0		SURFACE	<100	0
LUCKY 1 + JOE RA	UNITED URAN CORP	SAN MIGUEL	8	43 N	18,0 W	22	UNDERGRO	<100	200
LUCKY 8	PIKES PEAK URAN	SAN MIGUEL			0		UNDERGRO	<100	0
LUCKY DAY	DALPEZ, MAPIO	SAN MIGUEL	24	43 N	19,0 W	22	UNDERGRO	1,000 - 100,000	100
MAC INTYRE CLAIM	ORTHMYER MNG CO	SAN MIGUEL			0		SURFACE	<100	0
MAGPIE	DOYLE W K	SAN MIGUEL	36	45 N	18,0 W	22	UNDERGRO	1,000 - 100,000	100
MAINSTREET	PETRO NUCLEAR	SAN MIGUEL	16	45 N	18,0 W	22	UNDERGRO	1,000 - 100,000	100
MARGARET C. 1-6	TORRES, DAVID	SAN MIGUEL	7	43 N	16,0 W	22	UNDERGRO	<100	0
MARIE	TRE JOINT VENTUR	SAN MIGUEL	19	43 N	19,0 W	22	UNDERGRO	100 - 1,000	0
MARIE 1	TRE JOINT VENTUR	SAN MIGUEL	29	43 N	19,0 W	22	UNDERGRO	100 - 1,000	0
MARVE GROUP	FOOTE MINERALS	SAN MIGUEL	32	43 N	18,0 W	22	UNDERGRO	1,000 - 100,000	100
MARTHA BUE	SNYDER, FRANK	SAN MIGUEL	11	43 N	19,0 W	22	SURFACE	<100	0
MARY 2	ORTHMYER MINING	SAN MIGUEL			0		SURFACE	<100	0
MARY JANE-BROAD	HOPKINS MNG CO	SAN MIGUEL	22	45 N	18,0 W	22	UNDERGRO	1,000 - 100,000	0
MARY M.	FRONTIER VAN+URA	SAN MIGUEL			0		SURFACE	<100	0
MAY DAY	BALLS, E.M.	SAN MIGUEL	32	43 N	18,0 W	22	SURFACE	<100	0
MAYRE 1 + 2	ROGERS, W.A.	SAN MIGUEL	26	46 N	17,0 W	22	UNDERGRO	<100	100
MAYFLOWER	NIELSON + SONS	SAN MIGUEL	33	44 N	19,0 W	22	UNDERGRO	<100	0
MC KEE GROUP	RICE, JOE	SAN MIGUEL	22	45 N	18,0 W	22	UNDERGRO	<100	0
MC MILLAN	UNKNOWN CONTROLA	SAN MIGUEL			0		SURFACE	<100	0
MEMPHIS 1 + 2	LEE + SHALL	SAN MIGUEL	29	45 N	18,0 W	22	UNDERGRO	<100	100
MERCANTILE	DULANEY MNG CO	SAN MIGUEL	18	43 N	19,0 W	22	UNDERGRO	1,000 - 100,000	0
MERCANTILE 1	SITTON + DULANEY	SAN MIGUEL	18	43 N	19,0 W	22	UNDERGRO	100 - 1,000	0
MERCANTILE 2	SITTON + DULANEY	SAN MIGUEL	18	43 N	19,0 W	22	UNDERGRO	<100	0
MERCANTILE 3	SITTON + DULANEY	SAN MIGUEL	18	43 N	19,0 W	22	UNDERGRO	<100	0
MERCANTILE BETH	DULANEY MINING C	SAN MIGUEL	19	43 N	19,0 W	22	UNDERGRO	<100	0
MERCANTILE BRICH	DULANEY MINING C	SAN MIGUEL	17	43 N	19,0 W	22	UNDERGRO	100 - 1,000	0
MESA 7	DEVEREAUX BROS.	SAN MIGUEL	31	44 N	17,0 W	22	UNDERGRO	<100	0
MESA HILL	SNYDER, C.F. + SONS	SAN MIGUEL			0		UNDERGRO	100 - 1,000	0
MICKY 3	BAIRD-SNYDER MNG	SAN MIGUEL	20	43 N	19,0 W	22	SURFACE	<100	0
MIDNIGHT	URANIUM METALS	SAN MIGUEL	27	44 N	19,0 W	22	UNDERGRO	100 - 1,000	0
MINERAL MINES	WRIGHT, L.B.	SAN MIGUEL			0		UNDERGRO	100 - 1,000	0
MINERAL MOUNTAIN	WRIGHT, L.B.	SAN MIGUEL			0		UNDERGRO	<100	0
MINERAL MOUNTAIN	WRIGHT, KEITH	SAN MIGUEL			0		UNDERGRO	100 - 1,000	0
MINERAL MOUNTAIN	WRIGHT, L.B.	SAN MIGUEL			0		UNDERGRO	<100	0
MINERAL MOUNTAIN	WRIGHT, L.B.	SAN MIGUEL			0		UNDERGRO	100 - 1,000	0
MINERAL MOUNTAIN	WRIGHT, L.	SAN MIGUEL			0		UNDERGRO	100 - 1,000	0
MINERAL MOUNTAIN	WRIGHT, CLYDE	SAN MIGUEL			0		UNDERGRO	<100	0
MINERAL MOUNTAIN	WRIGHT, L.B.	SAN MIGUEL			0		UNDERGRO	100 - 1,000	0
MINING LEASE 1	SKIDMORE, T.M.	SAN MIGUEL	28	43 N	19,0 W	22	UNDERGRO	1,000 - 100,000	0
MINING LEASE 17	TURNER, JACK C.	SAN MIGUEL	30	44 N	18,0 W	22	UNDERGRO	1,000 - 100,000	200
MINING LEASE 18	MC CORMICK, WM.P	SAN MIGUEL	28	43 N	19,0 W	22	UNDERGRO	1,000 - 100,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 31

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CCNT'D) *****									
MINING LEASE 20	SAN JUAN LEASING	SAN MIGUEL	26	44 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
MINING LEASE 21	BAIRD+ROBINETTE+	SAN MIGUEL	5	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
MINING LEASE 27	GAMBLIN + FLOWLES	SAN MIGUEL	36	44 N	19.0 W	22	UNDERGRO	100 - 1,000	0
MINING LEASE 28	DUNCAN + RANCHEZ	SAN MIGUEL	32	44 N	18.0 W	22	UNDERGRO	1,000 - 100,000	0
MINING LEASE 29	ORTMEYER MNG CO	SAN MIGUEL	29	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
MINING LEASE 30	HARGROVE J.W.	SAN MIGUEL	31	44 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
MINING LEASE 32	BLUF CRFER MININ	SAN MIGUEL			0		UNDERGRO	1,000 - 100,000	150
MINING LEASE 4	DULANEY MNG CO	SAN MIGUEL	14	43 N	19.0 W	22	UNDERGRO	>100,000	0
MINING LEASE 42	BLIXT, ALFRED + OS	SAN MIGUEL	30	44 N	18.0 W	22	SURFACE	<100	0
MINING LEASE 43	KAMHOLZ, ROY	SAN MIGUEL	10	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
MINING LEASE 44	ATOMIC ENERGY CO	SAN MIGUEL	5	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
MINING LEASE 45	CROWLEY J.R.	SAN MIGUEL	15	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
MINING LEASE 46	GAYNO MINING CO	SAN MIGUEL	30	44 N	18.0 W	22	UNDERGRO	1,000 - 100,000	450
MINING LEASE 6	HOLLING, HENRY	SAN MIGUEL	23	44 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
MITCHEL+CUCHER G	FOGIE MINERALS	SAN MIGUEL	14	45 N	18.0 W	22	UNDERGRO	100 - 1,000	0
MONTEZUMA	DWENEY, H.R.	SAN MIGUEL	19	43 N	18.0 W	22	SURFACE	<100	0
MONUMENT 4	TRISTY + RYFRLY	SAN MIGUEL			0		SURFACE	<100	0
MOQUI JUG	UNION CARPIDE CP	SAN MIGUEL	29	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
MORTGAGE LIFTER	WILLIAMS + CONN	SAN MIGUEL	4	43 N	19.0 W	22	UNDERGRO	<100	0
MUCHO GRND(DOL.R	BROOKS MINERALS	SAN MIGUEL	1	42 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
MULE GROUP	HILL+DUNCAN+BUFF	SAN MIGUEL			0		UNDERGRO	<100	0
MULESHOE 6	HILL+DUNCAN+BUFF	SAN MIGUEL	22	43 N	18.0 W	22	UNDERGRO	<100	0
MYSTERY	BARPETT + BREWER	SAN MIGUEL	26	43 N	18.0 W	22	UNDERGRO	100 - 1,000	0
NATIONAL	CRISTEA MNG.CO,	SAN MIGUEL			0		SURFACE	<100	0
NELLIF GRAY	MADDOX,C.N.	SAN MIGUEL			0		SURFACE	<100	0
NEW DEAL	E H JOHNSSEN	SAN MIGUEL			0		UNDERGRO	100 - 1,000	0
NIGGER HEAD	MOTLEY, R.C.	SAN MIGUEL	36	44 N	20.0 W	22	SURFACE	<100	0
NORMA JEAN 1 + 2	UNION CARPIDE CP	SAN MIGUEL	28	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
NORTH CON MILL	TURNER, ACE	SAN MIGUEL			0		SURFACE	<100	0
NORTH SLOPE 2	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
NORTHER 5 + 6	UNION CARPIDE CP	SAN MIGUEL	22	44 N	19.0 W	22	UNDERGRO	1,000 - 100,000	550
NORTHERN LIGHT	MOTZ, EARL	SAN MIGUEL	19	44 N	18.0 W	22	UNDERGRO	100 - 1,000	200
OMEGA	UNKNOWN	SAN MIGUEL	35	44 N	11.0 W	22	SURFACE	<100	0
OWENSBY	UNKNOWN CONTROLLER	SAN MIGUEL	16	43 N	19.0 W	22	UNDERGRO	<100	0
PAINTED ROCK	PRETTY BLUE MNG	SAN MIGUEL	31	44 N	18.0 W	22	SURFACE	<100	0
PARROT GROUP	BROOKS MINERALS	SAN MIGUEL	26	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
PAYSTREAK 3	UNION CARPIDE CP	SAN MIGUEL	30	43 N	18.0 W	22	UNDERGRO	100 - 1,000	0
PFCAS 1	UNITED URAN CORP	SAN MIGUEL			0		SURFACE	<100	0
PENIGAL	DALPEZ, MARIO	SAN MIGUEL	24	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
PENJU	JACK THOMPSON	SAN MIGUEL	7	43 N	16.0 W	22	UNDERGRO	100 - 1,000	0
PHILLIPS 66 1	PHILLIPS, DORR	SAN MIGUEL	24	44 N	17.0 W	22	UNDERGRO	100 - 1,000	0
PIKE BUG	TURNER + SMITH	SAN MIGUEL	29	43 N	19.0 W	22	SURFACE	<100	0
PITCHFORK	BOKUM CORPORATION	SAN MIGUEL	32	44 N	16.0 W	22	UNDERGRO	1,000 - 100,000	100
POLAPIS 1	WAYERLY MINING C	SAN MIGUEL			0		SURFACE	<100	0
POUR OFF	UNKNOWN CONTROLLER	SAN MIGUEL			0		SURFACE	<100	0
PROSPECTORS FORT	ROBERTS RUTH	SAN MIGUEL	23	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
QUEEN OF SPADES	THUNDERBIRD URAN	SAN MIGUEL	18	43 N	18.0 W	22	UNDERGRO	<100	0
RADIUM	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 32

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
RADIUM 1	SITTON F A	SAN MIGUEL	5	43 N	19.0 W	22	SURFACE	<100	0
RADIUM 12	ATLAS-FOOTE	SAN MIGUEL	4	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	150
RADIUM 19	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	100 - 1,000	100
RADIUM 22	ATLAS-FOOTE	SAN MIGUEL	37	44 N	19.0 W	22	UNDERGRO	<100	100
RADIUM 24	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
RADIUM 25	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	SURFACE	<100	0
RADIUM 26 + 27	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	100 - 1,000	100
RADIUM 29	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	150
RADIUM 3	ATLAS-FOOTE	SAN MIGUEL	4	43 N	19.0 W	22	UNDERGRO	100 - 1,000	0
RADIUM 4	ATLAS-FOOTE	SAN MIGUEL	4	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
RADIUM 5 + 6	ATLAS-FOOTE	SAN MIGUEL	8	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
RADIUM 6	ATLAS-FOOTE	SAN MIGUEL	9	43 N	19.0 W	22	UNDERGRO	<100	0
RADIUM 7	ATLAS-FOOTE	SAN MIGUEL	8	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	150
RADIUM 8	ATLAS-FOOTE	SAN MIGUEL	9	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
RAINY DAY	UNION CARBIDE CP	SAN MIGUEL	35	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
RAMBLER	UNION CARBIDE CP	SAN MIGUEL	9	44 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
FAT HOLE	ROGERS, JACK	SAN MIGUEL					SURFACE	<100	0
RATTLESNAKE	RUTHERFORD, R.F.	SAN MIGUEL					SURFACE	<100	0
RATTLESNAKE	KNIGHT, N.B.	SAN MIGUEL	19	43 N	18.0 W	22	SURFACE	<100	0
RATTLESNAKE 1	CARKHUFF, CAMPBELL	SAN MIGUEL	4	45 N	19.0 W	22	SURFACE	<100	100
RED ANT	KNUCKLES, W.T.	SAN MIGUEL	30	43 N	19.0 W	22	UNDERGRO	<100	100
RED HORSE	UNION CARBIDE CP	SAN MIGUEL	2A	45 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
RED ROCK 4 + 5	BROOKS MINERALS	SAN MIGUEL	35	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	250
RED ROCK 5	BROOKS MINERALS	SAN MIGUEL	35	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	100
RED SNAKE	JF-ELL, LOREN	SAN MIGUEL					SURFACE	<100	50
RED TOP	SULLIVAN, BILL	SAN MIGUEL	6	42 N	17.0 W	22	UNDERGRO	<100	0
RED TOP	JOHANSON, HUGO W.	SAN MIGUEL	9	44 N	18.0 W	22	UNDERGRO	<100	50
RED WING	SALES, BURT	SAN MIGUEL	23	44 N	17.0 W	22	SURFACE	<100	0
RED CLAIM	REED, EDGAR J.	SAN MIGUEL					UNDERGRO	<100	0
RIM	DRINKARD, JOHN	SAN MIGUEL	1	42 N	18.0 W	22	SURFACE	<100	0
RIVERVIEW	UNION CARBIDE CP	SAN MIGUEL	33	45 N	19.0 W	22	UNDERGRO	100 - 1,000	50
ROBERT M	MARLANG, J. + C.	SAN MIGUEL	16	44 N	17.0 W	22	UNDERGRO	<100	0
ROBERTA JEAN	HAMPTON R.J.	SAN MIGUEL					UNDERGRO	<100	0
ROSE JUMP	MARLANG, FRED	SAN MIGUEL	16	44 N	17.0 W	22	UNDERGRO	<100	0
ROY LEE	BROOKS MINERALS	SAN MIGUEL	34	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
S.R. GROUP	DYER, GUY B.	SAN MIGUEL	11	42 N	18.0 W	22	UNDERGRO	1,000 - 100,000	4950
SAGE 11	SHUMWAY, DADE	SAN MIGUEL	32	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	300
SAN MIGUEL	ATLAS-FOOTE	SAN MIGUEL	8	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
SANTA MARIA	SKIDMORE, T.H.	SAN MIGUEL	14	43 N	19.0 W	22	UNDERGRO	<100	0
SARA M.	FOOTE MINERALS	SAN MIGUEL	26	43 N	18.0 W	22	UNDERGRO	100 - 1,000	0
SARAH ELLEN	GRIPE R.	SAN MIGUEL	26	43 N	18.0 W	22	UNDERGRO	100 - 1,000	0
SARAH JANE	TORRES, DAVID	SAN MIGUEL					SURFACE	<100	0
SCHLEY	SNYDER, ROBERT A.	SAN MIGUEL	26	43 N	19.0 W	22	SURFACE	<100	0
SIBLEY	DULANEY MINING C	SAN MIGUEL	17	43 N	19.0 W	22	UNDERGRO	100 - 1,000	0
SINGLE JACK	CRISTEA MNG. CO.	SAN MIGUEL	26	44 N	11.0 W	22	UNDERGRO	<100	0
SLICK RIM	ROSENQUIST, HOMER	SAN MIGUEL	16	45 N	18.0 W	22	UNDERGRO	<100	50
SLICK ROCK MILL	SKYLAND SALES, IN	SAN MIGUEL					UNDERGRO	100 - 1,000	0
SPRING	UNION CARBIDE CP	SAN MIGUEL	9	44 N	18.0 W	22	UNDERGRO	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 33

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** COLORADO (CONT'D) *****									
SPUD PATCH	UNION CARBIDE	SAN MIGUEL	29	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	50
STAR	OLIVER + RUSS	SAN MIGUEL	30	43 N	18.0 W	22	SURFACE	<100	0
SUNDAY	LEE H. MORRIS	SAN MIGUEL	29	43 N	18.0 W	22	UNDERGRO	100 - 1,000	0
SUNDOWN	LUBROCK MNG.CO.	SAN MIGUEL	27	44 N	19.0 W	22	UNDERGRO	<100	0
SUNDOWN GROUP	R + B MINING CO.	SAN MIGUEL	25	45 N	19.0 W	22	UNDERGRO	<100	200
SUNNYSIDE	BURNETT + CO	SAN MIGUEL	32	44 N	18.0 W	22	UNDERGRO	100 - 1,000	0
SUNRISE	ATLAS-FOOTE	SAN MIGUEL	5	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	350
SUNRISE + PATENT	ATLAS-FOOTE	SAN MIGUEL	5	44 N	19.0 W	22	UNDERGRO	1,000 - 100,000	350
SUNRISE 1	ATLAS-FOOTE	SAN MIGUEL	8	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	200
SUNRISE 3, 4, +	ATLAS-FOOTE	SAN MIGUEL	8	43 N	19.0 W	22	UNDERGRO	100 - 1,000	100
SUNSHINE 6	AYERS, EVERETT	SAN MIGUEL	21	44 N	17.0 W	22	UNDERGRO	<100	0
TAILMONT	CHARLES PICKENS	SAN MIGUEL	27	44 N	19.0 W	22	UNDERGRO	1,000 - 100,000	350
TERRIBLE	MAYFIELD, JERRY	SAN MIGUEL	10	45 N	19.0 W	22	UNDERGRO	1,000 - 100,000	0
TEXAS CHIEF 1	KUYFENDALL + ELMO	SAN MIGUEL			0		UNDERGRO	<100	0
TINY	UNION CARBIDE CP	SAN MIGUEL	30	44 N	16.0 W	22	UNDERGRO	1,000 - 100,000	100
TORNADO	UNION CARBIDE CP	SAN MIGUEL	3	45 N	19.0 W	22	UNDERGRO	100 - 1,000	350
TWO BAR	UNION CARBIDE CP	SAN MIGUEL	24	44 N	17.0 W	22	UNDERGRO	1,000 - 100,000	0
UNTAN 2 LODE	TUCHSCHER, C.L.	SAN MIGUEL	28	43 N	19.0 W	22	UNDERGRO	<100	0
UNCLE SAM	ALTMAN+PELLER+JA	SAN MIGUEL	6	42 N	17.0 W	22	UNDERGRO	100 - 1,000	0
UNCLE SAM(GYP VA	UNION CARBIDE CP	SAN MIGUEL	10	44 N	18.0 W	22	UNDERGRO	1,000 - 100,000	150
UTE 4	CRANDELL, MARION	SAN MIGUEL	4	45 N	19.0 W	22	UNDERGRO	<100	0
VALLEY VIEW	FOSTER, LYMAN	SAN MIGUEL	19	45 N	18.0 W	22	UNDERGRO	<100	0
VAN	TORRES, DAVID	SAN MIGUEL	12	46 N	17.0 W	22	SURFACE	<100	0
VAN,QUEEN(SLK RK	SHIPROCK,LTO	SAN MIGUEL	17	44 N	20.0 W	22	UNDERGRO	1,000 - 100,000	1500
VANADIUM	HATTERS, M.R.	SAN MIGUEL	31	44 N	18.0 W	22	UNDERGRO	100 - 1,000	0
VANADIUM 7	UNION CARBIDE CP	SAN MIGUEL	23	44 N	17.0 W	22	UNDERGRO	<100	0
VANADIUM QUEEN G	MC DOUGALD+WRIGH	SAN MIGUEL	12	44 N	20.0 W	22	SURFACE	<100	0
VANUPE CLAIMS	SHORT, ELBERT	SAN MIGUEL	11	44 N	18.0 W	22	UNDERGRO	<100	100
VICTOR	REED, EDGER J.	SAN MIGUEL	7	43 N	16.0 W	22	SURFACE	<100	0
VICTOR 2	BISHOP CANYON UP	SAN MIGUEL	20	43 N	19.0 W	22	UNDERGRO	100 - 1,000	50
VIRGINIA	JOHANNSEN E.J.	SAN MIGUEL			0		SURFACE	<100	0
WALLY-DOUBLE JAC	BERTOCH, F.M.	SAN MIGUEL	14	43 N	18.0 W	22	UNDERGRO	1,000 - 100,000	200
WHIZ BANG	FOOTE MINERALS	SAN MIGUEL	3	45 N	19.0 W	22	UNDERGRO	<100	0
WINDY DAY-BELL	HUNT, V.F.	SAN MIGUEL	8	43 N	16.0 W	22	UNDERGRO	100 - 1,000	50
WYOMING	DRINKARD, JOHN	SAN MIGUEL	15	30 S	24.0 E	24	UNDERGRO	<100	0
YELLOW GIPL	LYLE FRANCIS	SAN MIGUEL			0		UNDERGRO	100 - 1,000	150
YELLOWBIRD-RADIU	ATLAS-FOOTE	SAN MIGUEL	4	43 N	19.0 W	22	UNDERGRO	1,000 - 100,000	100
YORKTON	DAWSON, W.B.	SAN MIGUEL	19	45 N	18.0 W	22	SURFACE	<100	50
ZEBPA	ROCKWELL, ALVA A	SAN MIGUEL			0		UNDERGRO	<100	0
ZEBHA 1	GARNER, WILLARD	SAN MIGUEL			0		UNDERGRO	<100	100
ABRIL 2,4+8	CURTIS + THORPE	TELLER	30	15 S	70.0 W	22	SURFACE	<100	50
GENEVIEVE LODE	TRANS MTN.URAN+G	TELLER			0		UNDERGRO	<100	100
SEC. 36 SWO NEO	COTTER CORPORATION	TELLER	36	15 S	70.0 W	06	SURFACE	100 - 1,000	100

***** FLORIDA *****

W. R. GRACE	W. P. GRACE	POLK	0		MISC.-PA	<100	0
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INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 34

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** IDAHO *****									
COAL CREEK GROUP	LAUB, DON	CUSTER	17	11 N	14,0 E	08	UNDERGRO	1,000 - 100,000	50
EAST BASIN CREEK	EVANS, E.F.	CUSTER			0		SURFACE	1,000 - 100,000	100
ELK 1-DEERSTRIKE	LAUB, DON	CUSTER	15	11 N	14,0 E	08	UNDERGRO	1,000 - 100,000	100
LIGHTNING GROUP	UNKNOWN CONTROLR	CUSTER			0		UNDERGRO	100 - 1,000	50
SHORTY GROUP	UNKNOWN CONTROLR	CUSTER	6	11 N	14,0 F	08	UNDERGRO	1,000 - 100,000	50
DONNA LOU 1	UNKNOWN CONTROLR	LEMHI			0		SURFACE	<100	0
***** MISSOURI *****									
ST LOU ARPT PSD	COMMERCIAL D1	ST. LOUIS			0		MISC.-PB	1,000 - 100,000	50
***** MONTANA *****									
COBBAN LEASE	UNKNOWN CONTROLR	BROADWATER	20	9 N	1,0 E	23	SURFACE	<100	50
BOR 6	HIDDEN SPLENDOR	CARBON	17	9 S	27,0 E	23	SURFACE	100 - 1,000	50
BUCKHORN 2	MIDWEST EXPLRATH	CARBON	26	9 S	28,0 F	23	UNDERGRO	<100	50
DANDY-MARIE-PERC	BICE MINING CO	CARBON	8	9 S	27,0 E	23	SURFACE	1,000 - 100,000	50
FRAN	UNKNOWN CONTROLR	CARBON			0		UNDERGRO	100 - 1,000	50
KEY	BIG HORN MNG CO.	CARBON			0		SURFACE	<100	0
OLD GLORY	PRYOR MINING CO	CARBON	32	9 S	27,0 E	23	UNDERGRO	100 - 1,000	50
PEARL	BICE MINING CO	CARBON			0		SURFACE	<100	50
PEARC GROUP	MICHAUD, I.L.	CARBON	9	9 S	27,0 E	23	UNDERGRO	100 - 1,000	100
SANDRA	MIDLAND MNG. CO.	CARBON			0		UNDERGRO	<100	50
SNAIL 2	COLORED URANIUM	CARBON			0		UNDERGRO	100 - 1,000	50
SWAMP FROG	STOICK, JAMES J.	CARBON			0		UNDERGRO	100 - 1,000	50
WALDRON	WALDRON, P.	FALLON	19	10 N	61,0 E	05	SURFACE	<100	50
UNIT 3	TEXAS INSTRUMENT	HILL			0		SURFACE	<100	50
FREE ENTERPRISE	ELKHORN MINING	JEFFERSON			0		UNDERGRO	100 - 1,000	0
HAYNES HOMESTEAD	LENHART, PAYMOND	JEFFERSON			0		SURFACE	100 - 1,000	50
LONE EAGLE GROUP	EISELEIN, AM ETA	JEFFERSON	31	8 N	4,0 W	23	SURFACE	<100	150
RUN GROUP	POWELL, W.W.	MADISON	26	7 S	4,0 W	23	UNDERGRO	<100	50
***** NEVADA *****									
FIRST CHANCE	BUSHMAN, WILLIAM	CLARK	30	18 S	71,0 E	20	SURFACE	<100	50
GREEN MONSIFR	SMITH + COMPS	CLARK			0		SURFACE	<100	0
RACETPACK	TOM WHITE +ASSOC	ELKO			0		SURFACE	1,000 - 100,000	50
RIKROCK	BOGDANICH DEVEL.	ELKO			0		SURFACE	100 - 1,000	50
SOUTH FORY	VALLEY ENGR+DEV.	ELKO			0		UNDERGRO	1,000 - 100,000	50
MOONLIGHT	TRANS WORLD URAN	HUMBOLDT			0		UNDERGRO	100 - 1,000	100
EARLY DAY	APEX MINERALS	LANDER	2	18 N	43,0 E	20	UNDERGRO	1,000 - 100,000	100
LOW BOY	VALLEY VIEW URAN	LANDER	24	18 N	44,0 W	20	UNDERGRO	1,000 - 100,000	50
BLUE BIRD CLAIM	MULBE,PARKS+HEZZ	LINCOLN			0		UNDERGRO	100 - 1,000	50
WHITE CLOUD	LEE, LESTER E.	LINCOLN			0		SURFACE	<100	50
GLACIER KING CLA	MANCINI, FRANK	LYON			0		SURFACE	<100	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 31

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** NEVADA (CONT'D) *****									
RIVER ROAD	MOUNT, LLOYD	LYON			0		SURFACE	<100	100
BROKEN ROW GROUP	HOUCK, EDWARD	MINERAL			0		UNDERGRO	<100	0
CAROL R	BIG M URANIUM CO	MINERAL			0		UNDERGRO	100 = 1,000	50
BLACK RONANZA	FLAMINGO MINING	NYE			0		UNDERGRO	<100	50
ARMSTRONG I	ARMSTRONG, ARDEN	WASHOE			0		UNDERGRO	<100	50
BUCKHORN	DELAVERA, T.R.	WASHOE			0		SURFACE	100 = 1,000	50
FLAGG SECTION	NORTH AMERICAN A	WASHOE			0		UNDERGRO	<100	50
LOWERY CLAIMS	LOWERY URAN CO.	WASHOE			0		SURFACE	100 = 1,000	50
RED BLUFF	DE LONGCHAMPS F.	WASHOE			0		UNDERGRO	100 = 1,000	50
TICK CANYON	COURVOISIER, CHAS	WASHOE			0		UNDERGRO	<100	50
***** NEW JERSEY *****									
CHARLOTTE	BEMCO INC	UNION			0		UNDERGRO	<100	50
***** NEW MEXICO *****									
BABY MINE	THREE ELEVEN MNG	CATRON			0		UNDERGRO	<100	0
MIDNIGHT (MCPHAU)	GULF MINERAL	CATRON	11	2 N	11.0 W	22	SURFACE	100 = 1,000	50
RED BASIN	HAYSTACK MTH, DEV	CATRON	19	2 N	10.0 W	22	SURFACE	<100	0
SEC. 21, 34-164 NE	STATE OF NEW MEX	CATRON	21	3 N	16.0 W	22	UNDERGRO	<100	0
BLUE STAR	LEACH & LEACH	DOÑA ANA			0		SURFACE	<100	0
FLOYD COLLINS	LEACH, A.A.	GRANT			0		UNDERGRO	100 = 1,000	50
IMEZ-SEC 24-205-1	WESTERN EXPLORAT	GRANT			0		UNDERGRO	100 = 1,000	50
SEC. 21, 1RS-154	STATE OF NEW MEX	GRANT			0		SURFACE	<100	0
POLITA 2, 17N-29E	PACHECO, J. AND B.	HARDING	5	17 N	29.0 F	22	SURFACE	<100	50
NAPANE SEC 25 N1	BAKER WILDER SYN	HIDALGO			0		UNDERGRO	<100	0
29-14-09 MINE WA	REKERR MCGEE	MCKINLEY			0		WHATPROD	<100	0
ALTA (5+6-11-4)	CITRUS CNTY LAND	MCKINLEY	6	14 N	11.0 W	22	UNDERGRO	1,000 = 100,000	200
B HILL 18+23(SEC	CITRUS CNTY LAND	MCKINLEY	20	13 N	9.0 W	22	UNDERGRO	1,000 = 100,000	150
BARB J 2(30,13-1	SMITH DEVELOPMNT	MCKINLEY	30	13 N	9.0 W	22	UNDERGRO	1,000 = 100,000	250
BARB J 3 (30-13-	MID-CONT MNG. CORP	MCKINLEY	30	13 N	9.0 W	22	UNDERGRO	>100,000	50
BARB J 1 (30-13-	MID-CONT MNG. CORP	MCKINLEY	30	13 N	9.0 W	22	UNDERGRO	1,000 = 100,000	400
BG-GRP(20-13-9)	UNITED NUCLEAR	MCKINLEY	20	13 N	9.0 W	22	UNDERGRO	1,000 = 100,000	200
BILLY THE FID	HENRY ANDREWS	MCKINLEY	19	14 N	11.0 W	22	UNDERGRO	1,000 = 100,000	50
BLACK JACK 2	INDIAN ALLOTEE	MCKINLEY	18	15 N	13.0 W	22	UNDERGRO	>100,000	250
BLACK JACK 1	INDIAN ALLOTEE	MCKINLEY	12	15 N	13.0 W	22	UNDERGRO	>100,000	700
BLUE PEAK(24-13-	GARCIA & ANDREWS	MCKINLEY	24	13 N	10.0 W	22	UNDERGRO	1,000 = 100,000	0
BOBCAT NEQ24-13-	HYDE ESTATE	MCKINLEY	24	13 N	9.0 W	22	UNDERGRO	<100	0
C D-5 (15-16-17)	SANTA FE PACIFIC	MCKINLEY	35	16 N	17.0 W	22	UNDERGRO	<100	50
CHAPLOTTE(33-13-	SANTA FE PACIFIC	MCKINLEY	33	13 N	9.0 W	22	UNDERGRO	100 = 1,000	0
CHILL WILLS(RALT	MARQUEZ, NABOR	MCKINLEY	24	13 N	9.0 W	22	UNDERGRO	1,000 = 100,000	450
CHURCHROCK (8+17	UNITED NUCLEAR	MCKINLEY	17	16 N	16.0 W	22	UNDERGRO	>100,000	650
CLIFFSIDE BFC 36	KERR-MCGEE CORP.	MCKINLEY			0		LOWGRADE	1,000 = 100,000	0
DAKOTA(4-13N-104	STATE OF NEW MEX	MCKINLEY	4	13 N	10.0 W	22	UNDERGRO	1,000 = 100,000	0
DAVENPORT(20-13-	HYDE ESTATE	MCKINLEY	20	13 N	9.0 W	22	UNDERGRO	<100	150

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 36

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.
***** NEW MEXICO (CONT'D) *****									
UNDERGRO	1,000 - 100,000	300	DIAMOND 2 LARGO	SHIPROCK, LTD	MCKINLEY	33	15 N	17.0 W	22
UNDERGRO	>100,000	400	DYSART 1 SEC 11	HOMESTEAK-U.M.	MCKINLEY	11	14 N	10.0 W	22
UNDERGRO	>100,000	450	DYSART 2	UNITED N. MISTKE	MCKINLEY			0	
UNDERGRO	1,000 - 100,000	200	E. MALPAIS 20-13-	FOUR CORNERS EXP	MCKINLEY	20	13 N	9.0 W	22
UNDERGRO	<100	50	ELKINS (24-14-17)	ELKINS, LAWRENCE	MCKINLEY			0	
UNDERGRO	1,000 - 100,000	50	EVFLYA (9-14-11)	CITRUS CNTY LAND	MCKINLEY	9	14 N	11.0 W	22
UNDERGRO	1,000 - 100,000	100	FAITH (29-13-09)	RANCHERS EXP. DEV	MCKINLEY	29	13 N	9.0 W	22
UNDERGRO	100 - 1,000	100	FLAT TOP 1 2 3	HYDE ESTATE	MCKINLEY	30	13 N	9.0 W	22
UNDERGRO	1,000 - 100,000	100	FLAT TOP VIL HYD	HYDE ESTATE	MCKINLEY	30	13 N	9.0 W	22
UNDERGRO	100 - 1,000	50	FOUTZ 1	FOUTZ MNG. CO.	MCKINLEY	4	15 N	16.0 W	22
UNDERGRO	100 - 1,000	100	FOUTZ 2	FOUTZ MNG. CO.	MCKINLEY			0	
UNDERGRO	1,000 - 100,000	50	FOUTZ 3	FOUTZ MNG. CO.	MCKINLEY	31	14 N	16.0 W	22
UNDERGRO	100 - 1,000	0	FRANCIS	CITRUS CNTY LAND	MCKINLEY	8	14 N	11.0 W	22
UNDERGRO	>100,000	100	HOGAN (14-13-9)	UNITED N. MISTKE	MCKINLEY	14	13 N	9.0 W	22
UNDERGRO	1,000 - 100,000	0	HOGBACK 3-5	HYDE ESTATE	MCKINLEY	12	15 N	12.0 W	22
UNDERGRO	1,000 - 100,000	300	ISABELLA (5-7-13-	M. P. GRACE	MCKINLEY	7	13 N	9.0 W	22
UNDERGRO	<100	100	LOST MINE (35-14-	BERRYHILL, ELKINS	MCKINLEY	35	14 N	11.0 W	22
UNDERGRO	1,000 - 100,000	250	MAC 2 (17-18-15-1	INDIAN ALLOTTEE	MCKINLEY	18	15 N	13.0 W	22
UNDERGRO	1,000 - 100,000	200	MALPAIS MINE	SMITH DEVELOPMENT	MCKINLEY	20	13 N	9.0 W	22
UNDERGRO	>100,000	250	MARQUEZ (23-13-0	SANTA FE RAILRO	MCKINLEY	24	13 N	9.0 W	22
UNDERGRO	>100,000	350	MARY MOUNT	UNITED N. MISTKE	MCKINLEY	11	14 N	10.0 W	22
UNDERGRO	>100,000	200	MESA TOP (20-13-	HYDE ESTATE	MCKINLEY	20	13 N	9.0 W	22
UNDERGRO	>100,000	0	NAVAJO RES	KERR-MCGEE CORP	MCKINLEY			0	
SURFACE-	100 - 1,300	50	RED CAP (T-GROUHP	NAVAJO DEVELOPMT.	MCKINLEY			0	
UNDERGRO	100 - 1,000	50	RED POINT 16-13-	STATE OF NEW MEX	MCKINLEY	16	13 N	10.0 W	22
UNDERGRO	100 - 1,000	50	RENTOP 1+2 (18-14	RED TOP URAN. MNG	MCKINLEY	18	14 N	11.0 W	22
SURFACE	<100	0	SANTA FE CHRIST	PEM URANIUM CO	MCKINLEY			0	
UNDERGRO	1,000 - 100,000	200	SEC 16 134 9W	STATE OF NEW MEX	MCKINLEY	16	13 N	9.0 W	22
UNDERGRO	1,000 - 100,000	200	SEC 17 13 9	SANTA FE PACIFIC	MCKINLEY	17	13 N	9.0 W	22
LOWGRADE	1,000 - 100,000	0	SEC 17 144 9W LG	KERR-MCGEE CORP.	MCKINLEY			0	
UNDERGRO	1,000 - 100,000	150	SEC 18 134 9W	CITRUS CNTY LAND	MCKINLEY	18	13 N	9.0 W	22
LOWGRADE	1,000 - 100,000	0	SEC 20 144 9W LG	KERR MCGEE	MCKINLEY			0	
LOWGRADE	1,000 - 100,000	0	SEC 22 144 10W L	KERR MCGEE	MCKINLEY			0	
UNDERGRO	>100,000	450	SEC 22, 14-10	KERR MCGEE	MCKINLEY	22	14 N	10.0 W	22
UNDERGRO	1,000 - 100,000	50	SEC 23 134 10W	SANTA FE PACIFIC	MCKINLEY	23	10 N	13.0 W	22
LOWGRADE	>100,000	0	SEC 24 144 10W L	KERR MCGEE	MCKINLEY			0	
UNDERGRO	100 - 1,000	50	SEC 24 RECENTI	TUCKER, A.W.	MCKINLEY	24	15 N	17.0 W	22
LOWGRADE	1,000 - 100,000	0	SEC 29 144 9W LG	KERR MCGEE	MCKINLEY			0	
LOWGRADE	>100,000	0	SEC 30 144 9W LG	KERR MCGEE	MCKINLEY			0	
UNDERGRO	1,000 - 100,000	600	SEC 31 144 9W	KERR-MCGEE CORP.	MCKINLEY	31	14 N	9.0 W	22
UNDERGRO	>100,000	700	SEC 33, 14-9 ONLY	KERR MCGEE	MCKINLEY	29	14 N	9.0 W	22
LOWGRADE	>100,000	0	SEC 33-14-09 LG	KERR-MCGEE	MCKINLEY			0	
UNDERGRO	1,000 - 100,000	50	SEC 36 134 10W	STATE OF NEW MEX	MCKINLEY	36	13 N	10.0 W	22
UNDERGRO	>100,000	400	SEC. 10-144-10W	UNITED NUC HOMSTKE	MCKINLEY	10	14 N	10.0 W	22
UNDERGRO	1,000 - 100,000	150	SEC. 18, 134-104 3	VANDEVER, N.	MCKINLEY	18	13 N	10.0 W	22
UNDERGRO	1,000 - 100,000	50	SEC. 24-134-11W 4	VANDEVER,	MCKINLEY	24	13 N	11.0 W	22
SURFACE	>100,000	100	SEC. 25-134-10W 3	CITRUS CNTY LAND	MCKINLEY	25	13 N	10.0 W	22
UNDERGRO	1,000 - 100,000	50	SEC. 26-134-10W 4	ESEDEPO, N.	MCKINLEY	26	13 N	10.0 W	22

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 37

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** NE. MEXICO (CONT'D) *****									
SEC.3-15-16LIC41	STATE OF NEW MEX	MCKINLEY	3	15 N	16.0 W	22	UNDERGRO	100 - 1,000	50
SEC.30-13-9 WH 8	BALLEY + FIFF	MCKINLEY	30	13 N	9.0 W	22	UNDERGRO	1,000 - 100,000	200
SEC.31,13N-09N	SANTA FE PACIFIC	MCKINLEY	31	13 N	9.0 W	22	UNDERGRO	1,000 - 100,000	50
SEC.32,13N-9W	STATE OF N.M.	MCKINLEY	32	13 N	9.0 W	22	UNDERGRO	1,000 - 100,000	250
SEC.32-15N-11N	HYDRO NUCLEAR	MCKINLEY	32	15 N	11.0 W	22	UNDERGRO	1,000 - 100,000	300
SEC.32-15N-11N 8	HYDRO NUCLEAR	MCKINLEY	32	15 N	11.0 W	22	UNDERGRO	1,000 - 100,000	300
SEC.33-15-11 N-M	RANCHERS EXP.DEV	MCKINLEY	33	15 N	11.0 W	22	UNDERGRO	1,000 - 100,000	0
SEC.5,13N-10N	SANTA FE PACIFIC	MCKINLEY			0		UNDERGRO	<100	0
SECTION 23-16N-1	GRACE NUCLEAR	MCKINLEY	23	16 N	17.0 W	22	UNKNOWN	<100	700
SILVER BIT	ELKINS, LAWRENCE	MCKINLEY	10	14 N	17.0 W	22	UNDERGRO	100 - 1,000	100
SILVERSPRING(3114)	CITRUS CNTY LAND	MCKINLEY	31	14 N	10.0 W	22	UNDERGRO	1,000 - 100,000	0
U MINE (4-15-16)	CHRISTENSEN EST	MCKINLEY	4	15 N	16.0 W	22	UNDERGRO	1,000 - 100,000	100
VALLEJO (SEC. 34	FARRIS MINES	MCKINLEY	34	13 N	9.0 W	22	UNDERGRO	1,000 - 100,000	200
WESTWATER 1	STATE OF NEW MEX	MCKINLEY			0		UNDERGRO	1,000 - 100,000	150
WHITECAPS	SMITH DEVELOPMENT	MCKINLEY	30	13 N	9.0 W	22	UNDERGRO	1,000 - 100,000	200
LEDOUX RANCH	GALISTEO MINING	MORA	22	7 N	16.0 E	22	SURFACE	<100	0
GOOD LUCK	DEYNNIS, L.T.	QUAY	6	7 N	32.0 E	22	SURFACE	<100	100
SEC.12,11N-33E	HIGHLAND DEVEL.C	QUAY			0		SURFACE	<100	0
TWP.11N+12N-33E	LITTLE RATTLES M	QUAY			0		UNDERGRO	<100	50
BOX CANYON	BOX CANYON MNG.	RIO ARriba			0		UNDERGRO	100 - 1,000	250
COYOTE HILLS	13N LAND+INVEST.	RIO ARriba			0		SURFACE	<100	50
J.O.L.	ARRIBA URANIUM C	RIO ARriba	24	26 N	7.0 F	22	SURFACE	<100	50
LUCKY STRIKE	ARROYA DE AGUA M	RIO ARriba			0		UNDERGRO	<100	300
PINEAPPLE + PALO	WELLS, S.M.	RIO ARriba	30	26 N	9.0 E	22	UNDERGRO	<100	0
RED HEAD 2	BOLIVAR URANIUM	RIO ARriba			0		UNDERGRO	<100	50
TUSAS EAST SLOPE	COLONIAL URANIUM	RIO ARriba	24	28 N	7.0 E	22	UNDERGRO	<100	50
WHITEFLO 1	WHITEFLO MNG.CO.	RIO ARriba	30	23 N	1.0 F	22	SURFACE	<100	50
BUTLER BROTHERS	BUTLER PROS.	SANDOVAL	23	19 N	1.0 W	22	UNDERGRO	<100	50
COLTINS	ZIA IND. RES.	SANDOVAL	24	17 N	1.0 W	22	UNDERGRO	100 - 1,000	700
CORRAL 3	SLA TEX VENTURE	SANDOVAL	25	23 N	1.0 F	22	UNDERGRO	<100	1000
14	MEADOWS MINING C	SAN JUAN			0		UNDERGRO	<100	200
ALONCO MINE	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	<100	50
B 8	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	<100	100
B B 8	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	100 - 1,000	150
RECLABITO	NAVAJO TRIBE	SAN JUAN			0		UNKNOWN	100 - 1,000	0
BEGAY 1	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
BEGAY 2	NAVAJO TRIBE	SAN JUAN	23	29 N	21.0 W	22	UNDERGRO	1,000 - 100,000	50
CANYON VIEW	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	100 - 1,000	150
CARL YAZZIE 1	NAVAJO TRIBE	SAN JUAN			0		SURFACE	<100	150
CASTLE TSOBIE	NAVAJO TRIBE	SAN JUAN			0		SURFACE	<100	100
DENEH NEZ 1	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	100 - 1,000	150
EAST NEW MEXICO	NAVAJO TRIBE	SAN JUAN			0		SURFACE	<100	200
EAST RESERVATION	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
ENOS JOHNSON 1-4	RAY WILLIAMS	SAN JUAN	5	23 N	20.0 W	22	UNDERGRO	1,000 - 100,000	300
H B ROY 2	NAVAJO TRIBE	SAN JUAN			0		SURFACE	<100	50
HOG BACK 2	DAVIDSON, WILLIE	SAN JUAN			0		SURFACE	<100	0
MORACE BEN 1	NAVAJO TRIBE	SAN JUAN			0		SURFACE	<100	250
JACK BOYD CLAIMS	BOYD, JACK	SAN JUAN			0		SURFACE	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 38

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MINING METHOD	RANGE	MERID.	SEC.	TOWNSHIP	COUNTY	CONTROLLER NAME	MINE NAME
***** NEW MEXICO (CONT'D) *****										
UNDERGRO	100 - 1,000	150	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	JOE BEN 1
UNDERGRO	<100	200	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	JOHN JOE 1
UNDERGRO	<100	0	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	JOHN JOHN MP 334
SURFACE	<100	100	SURFACE	0				SAN JUAN	NAVAJO TRIBE	JUNCTION
UNDERGRO	<100	350	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	KEY TONE
UNDERGRO	100 - 1,000	100	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	KING 2
UNDERGRO	100 - 1,000	100	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	KING TUTT 1
UNDERGRO	<100	200	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	KING TUTT POINT
UNDERGRO	<100	50	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	KINGS 6
UNDERGRO	1,000 - 100,000	100	UNDERGRO	31.0 E	14	29	39 N	SAN JUAN	NAVAJO TRIBE	LOWER CANYON
UNDERGRO	1,000 - 100,000	50	UNDERGRO	21.0 W	22	23	28 N	SAN JUAN	NAVAJO TRIBE	NELSON POINT
UNDERGRO	100 - 1,000	250	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	PLOT 1 RED WASH
UNDERGRO	100 - 1,000	200	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	PLOT 2 KING TUTT
UNDERGRO	1,000 - 100,000	100	UNDERGRO	21.0 W	22	11	29 N	SAN JUAN	NAVAJO TRIBE	PLOT 7 OAK SPRIN
UNDERGRO	100 - 1,000	0	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	PLOT 8 COTTONWOOD
UNDERGRO	100 - 1,000	0	UNDERGRO	21.0 W	22	35	30 N	SAN JUAN	NAVAJO TRIBE	PLOT 9 LONE STAR
UNDERGRO	<100	200	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	REED HENDERSON 1
UNDERGRO	100 - 1,000	250	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	ROCKY FLATS MINE
UNDERGRO	<100	100	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	SALT CANYON
UNDERGRO	1,000 - 100,000	100	UNDERGRO	21.0 W	22	23	29 N	SAN JUAN	NAVAJO TRIBE	SHADYSIDE
UNDERGRO	100 - 1,000	100	UNDERGRO	21.0 W	22	23	29 N	SAN JUAN	NAVAJO TRIBE	SHADYSIDE 2
MISC.-PB	100 - 1,000	50	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	SHIPROCK RESIDUE
UNDERGRO	<100	0	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	SUNNYSIDE
UNDERGRO	1,000 - 100,000	100	UNDERGRO	20.0 W	22	18	24 N	SAN JUAN	NAVAJO TRIBE	TENT 1
UNDERGRO	<100	0	UNDERGRO	0				SAN JUAN	NAVAJO TRIBE	UPPER SALT WASH
UNDERGRO	<100	0	UNDERGRO	24.0 E	22	31	17 N	SAN MIGUEL	FURMAN+FOSTER	BLAH 2
UNDERGRO	<100	50	UNDERGRO	14.0 E	22	5	16 N	SAN MIGUEL	SPARKS STONE MNG	SPARKS-STONE 1
UNDERGRO	<100	0	UNDERGRO	23.0 E	22	14	17 N	SAN MIGUEL	SAN CARLOS URAN	WINDY NINE MINE
UNDERGRO	1,000 - 100,000	200	UNDERGRO	7.0 E	22	9	15 N	SANTA FE	LOVE STAR MNG+DV	LA BAJADA
UNDERGRO	<100	0	UNDERGRO	9.0 E	22	29	20 N	SANTA FE	ROYAL, J.C.	SAN JOSE CLAIMS
UNDERGRO	<100	0	UNDERGRO	0				SIERRA	THE EMPIRE PROJ.	EMPIRE CLAIMS
UNDERGRO	<100	50	UNDERGRO	0				SIERRA	ALAMO MINING CO.	LAST CHANCE
SURFACE	<100	50	SURFACE	0				SIERRA	PRICE BUELL	MITCHELL PRICE 1
SURFACE	<100	50	SURFACE	0				SIERRA	SAMANIEGO, BERNIE	PARAN
UNDERGRO	<100	50	UNDERGRO	0				SIERRA	TERRY, EARL H.	PITCHBLEND STRIK
SURFACE	<100	50	SURFACE	0				SIERRA	AUGER, ROBERT	ST MIN LEASE 61
UNDERGRO	100 - 1,000	100	UNDERGRO	0				SOCORRO	CAMPBELL FARMING	AUGUA TORRES
UNDERGRO	100 - 1,000	50	UNDERGRO	6.0 W	22	13	1 N	SOCORRO	LUHMUS, R.H.	HOOKS RANCH
SURFACE	1,000 - 100,000	0	SURFACE	0				SOCORRO	CAPITOL STABOARD	JETER MINE
SURFACE	<100	50	SURFACE	0				SOCORRO	SANTA FE PACIFIC	LITTLE DAVIE
UNDERGRO	1,000 - 100,000	50	UNDERGRO	0				SOCORRO	LUHMUS, R.H.	LUCKY DON SEC 35
UNDERGRO	<100	50	UNDERGRO	0				SOCORRO	CAMPBELL FARMING	MARY BALL 1
SURFACE	<100	50	SURFACE	0				SOCORRO	GRIEGO, JOE	SAN LORENZO 1 NW
UNDERGRO	<100	0	UNDERGRO	15.0 E	22	26	28 N	TAOS	BLACK COPPER MIN	BLACK COPPER 2 L
UNDERGRO	1,000 - 100,000	50	UNDERGRO	9.0 W	22	4	12 N	VALENCIA	GRANTS STATE BNK	BLACK HAWK + BUN
UNDERGRO	1,000 - 100,000	50	UNDERGRO	9.0 W	22	20	11 N	VALENCIA	YUCCA URANIUM	CEDAR CLAIMS 1 7
UNDERGRO	100 - 1,000	50	UNDERGRO	3.0 W	22	22	10 N	VALENCIA	CANONCITO IND,RS	CHAVEZ (22-10-3)
UNDERGRO	1,000 - 100,000	150	UNDERGRO	9.0 W	22	4	12 N	VALENCIA	UNKNOWN CONTRLR	CHRISTMAS DAY PH

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 39

MIKE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** NEW MEXICO (CONT'D) *****									
CRACKPOT(57-8-8-	DEVILLIERS NUC.	VALENCIA			0		UNDERGRO	1,000 - 100,000	450
FALCON 1 + 2	YUCCA URANIUM	VALENCIA	21	43 N	19.0 W	22	SURFACE	<100	50
LA JARA	CIBOLA MNG. CO.	VALENCIA	15	12 N	9.0 W	22	UNDERGRO	1,000 - 100,000	50
LAST CHANCE	BROADBUSH, J. FOLEY	VALENCIA			0		SURFACE	1,000 - 100,000	100
LOVE PINE 3	TYCONDA MINERALS	VALENCIA	8	11 N	9.0 W	22	SURFACE	100 - 1,000	150
PAISANO (16-8-6)	GOOD NEWS MNG. LT	VALENCIA			0		SURFACE	<100	50
RED BLUFF GAY FA	MOISES+HIPANEL	VALENCIA	4	12 N	9.0 W	22	UNDERGRO	1,000 - 100,000	50
SAN MATEO(30-13-	UNITED NUCLEAR	VALENCIA	30	13 N	8.0 W	22	UNDERGRO	>100,000	1050
SANDY (SEC.27-9-	DEVILLIERS NUC.	VALENCIA	27	9 N	5.0 W	22	SURFACE	100 - 1,000	50
SEC 33-12N-9N	HOMESTAKE MING	VALENCIA	33	12 N	9.0 W	22	UNDERGRO	>100,000	150
SEC 9 12N 9W	ELKINS, MARK	VALENCIA	9	12 N	9.0 W	22	UNDERGRO	1,000 - 100,000	50
TAFFY (SEC.11-12	NATIONAL GROWTH	VALENCIA	14	12 N	9.0 W	22	UNDERGRO	100 - 1,000	0
TOM 13(SEC.4-11-	ANACONUA CO.	VALENCIA			0		SURFACE	<100	50
UDC 5 (SEC.4-12-	COFFEY, WILLIAM	VALENCIA			0		UNDERGRO	100 - 1,000	50
WOODROW (36-11-0	ANACONUA CO.	VALENCIA			0		UNDERGRO	1,000 - 100,000	150
***** NORTH DAKOTA *****									
BARANKO LEASE	MCCANN + GETTING	BILLINGS	23	42 N	99.0 W	05	SURFACE	1,000 - 100,000	0
CHURCH 1	AMERICAN LUTH CH	BILLINGS	5	36 N	00.0 W	05	SURFACE	1,000 - 100,000	50
GEARY LEASE	GARDNER+WALSH+S	BILLINGS	4	37 N	00.0 W	05	SURFACE	<100	0
JOHNSON MINE+SFC	US REACQUIRED	BILLINGS	3	40 N	00.0 W	05	SURFACE	1,000 - 100,000	50
KLYM	KLYM, N.O.	BILLINGS	26	42 N	99.0 W	07	SURFACE	1,000 - 100,000	50
ONYUS (NP-1)	NORTHERN PACIFIC	BILLINGS	15	41 N	00.0 W	05	SURFACE	1,000 - 100,000	50
SEC.2,37N-100W,5	GEOLPSOURCES, INC	BILLINGS	2	37 N	00.0 W	05	SURFACE	1,000 - 100,000	50
SMITH 1 LFASE	HAYDEN, GARY	BILLINGS	6	37 N	00.0 W	05	SURFACE	100 - 1,000	50
SPIKE 1	DE SART + KYLE	BILLINGS	10	37 N	01.0 W	05	SURFACE	<100	0
HURICK 1	HURICK, JOE	BLMPE	5	36 N	00.0 W	05	SURFACE	1,000 - 100,000	50
FRANK LEASE	MIKE FRANK	STARK	7	40 N	99.0 W	05	SURFACE	1,000 - 100,000	50
LUPTAK 1	LUPTAK, ANNA	STARK	6	40 N	99.0 W	05	SURFACE	1,000 - 100,000	50
LUPTAK STATE	LUPTAK+ST.N.DAK.	STARK	6	40 N	99.0 W	05	SURFACE	1,000 - 100,000	50
***** OKLAHOMA *****									
CEMENT	LISTER MILLS	CADDO			0		SURFACE	<100	150
SCHOOL LAND	LISTER MILLS	CADDO			0		SURFACE	<100	0
THORPE 1	PRATT, ORVILLE L	CUSTER			0		SURFACE	<100	0
***** OREGON *****									
PINE SPRINGS	UNKNOWN CONTROL	CROOK	13	18 S	16.0 E		UNDERGRO	<100	50
LUCKY LASS	WESTERN NUCLEAR	LAKE	25	37 S	18.0 E	33	SURFACE	1,000 - 100,000	100
WHITE KING	WESTERN NUCLEAR	LAKE	30	37 S	19.0 F	33	SURFACE	>100,000	50
***** SOUTH DAKOTA *****									
APLAND 1	UNKNOWN CONTROL	BUTTE			0		SURFACE	100 - 1,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 40

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MINING METHOD	SEC.	TOWNSHIP	RANGE	MERID.	COUNTY	CONTROLLER NAME	MINE NAME
***** SOUTH DAKOTA (CONT'D) *****										
SURFACE	<100	0	SURFACE	25	8 N	1.0 E	07	BUTTE	ENRICO BONATO	BONATO RANCH
UNDERGRO	1,000 - 100,000	50	UNDERGRO	23	8 N	1.0 E	07	BUTTE	ALEX KLING	KLING LEASE
SURFACE	>100,000	100	SURFACE	34	6 S	1.0 E	07	CUSTER	TENN. VALLEY AUTH	BUD-LUCKY RND
SURFACE	<100	50	SURFACE	6	2 S	7.0 E	07	CUSTER	WESTERN GIANT OI	CAYLOR LEASE
SURFACE	100 - 1,000	0	SURFACE			0		CUSTER	DARROW + COOK	DARK 1
SURFACE	<100	50	SURFACE			0		CUSTER	GENTLEMEN MINING	DEL MUERTO
SURFACE	1,000 - 100,000	100	SURFACE	36	6 S	1.0 E	07	CUSTER	TENN. VALLEY AUTH	FREEZEOUT GROUP
SURFACE	<100	50	SURFACE			0		CUSTER	ROBINSON, DELBERT	JIMBOR 1
SURFACE	<100	50	SURFACE			0		CUSTER	KERCHERVAL, R.S.	LOST CANYON 1
SURFACE	<100	50	SURFACE			0		CUSTER	KENNEDY, MIKE C.	M.C.1
UNDERGRO	1,000 - 100,000	100	UNDERGRO	33	7 S	3.0 E	07	CUSTER	TENN. VALLEY AUTH	TOO LATE
SURFACE	<100	50	SURFACE			0		CUSTER	CRAYEN UPAT. CO.	URANIUM 1
UNDERGRO	100 - 1,000	50	UNDERGRO	31	8 S	4.0 E	07	FALL RIVER	LORFENZ BROTHERS	ACCIDENTAL GROUP
UNDERGRO	<100	0	UNDERGRO			0		FALL RIVER	LORD, B. H. & PARY J.	AMY 1
SURFACE	<100	0	SURFACE			0		FALL RIVER	ALLIED EXPL. CO.	APPLE PIE
SURFACE	100 - 1,000	50	SURFACE			0		FALL RIVER	TENN. VALLEY AUTH	B. & H. GROUP
SURFACE	1,000 - 100,000	0	SURFACE	21	7 S	2.0 E	07	FALL RIVER	TENN. VALLEY AUTH	BARKER + HOWELL
UNDERGRO	<100	50	UNDERGRO			0		FALL RIVER	BAXTER, F.P.	BAXTER 1
SURFACE	<100	50	SURFACE	24	8 S	3.0 F	07	FALL RIVER	TENN. VALLEY AUTH	BEADLE GROUP
SURFACE	<100	50	SURFACE			0		FALL RIVER	DAKOTA URAN+DPLG	BERIT 2
SURFACE	<100	50	SURFACE			0		FALL RIVER	CHORD, ROY E.	BITS GROUP
SURFACE	100 - 1,000	0	SURFACE	24	7 S	2.0 E	07	FALL RIVER	CHORD, ROY F.	BLUE NOTE
UNDERGRO	1,000 - 100,000	200	UNDERGRO	3	8 S	3.0 E	07	FALL RIVER	TENN. VALLEY AUTH	BUDA-DEXTER
UNDERGRO	100 - 1,000	0	UNDERGRO			0		FALL RIVER	TENN. VALLEY AUTH	CHILSON CANYON
UNDERGRO	1,000 - 100,000	100	UNDERGRO	19	7 S	3.0 F	07	FALL RIVER	CHORD, ROY E.	CLARABELLE GROUP
SURFACE	1,000 - 100,000	0	SURFACE			0		FALL RIVER	CHORD, ROY E.	COAL CANYON
SURFACE	100 - 1,000	0	SURFACE	26	7 S	2.0 F	07	FALL RIVER	CHORD, ROY E.	COAL CANYON 1
SURFACE	<100	0	SURFACE			0		FALL RIVER	HUFF J.V.	COAL CANYON 14
SURFACE	1,000 - 100,000	50	SURFACE	12	8 S	3.0 F	07	FALL RIVER	TENN. VALLEY AUTH	CRAM LEASE
SURFACE	<100	50	SURFACE	4	8 S	3.0 F	07	FALL RIVER	TENN. VALLEY AUTH	CRANDALL GROUP
SURFACE	100 - 1,000	50	SURFACE			0		FALL RIVER	TENN. VALLEY AUTH	CYCAD
SURFACE	100 - 1,000	50	SURFACE	25	7 S	6.0 E	07	FALL RIVER	CHORD, ROY E.	DAKOTA FLATS
SURFACE	100 - 1,000	50	SURFACE	31	8 S	4.0 E	07	FALL RIVER	LORFENZ BROTHERS	DAMSITE
SURFACE	<100	200	SURFACE			0		FALL RIVER	DISNEY MINING CO	DARNELL LEASE
SURFACE	>100,000	100	SURFACE	2	7 S	1.0 E	07	FALL RIVER	TENN. VALLEY AUTH	DARROW 1-5 AND 4
SURFACE	1,000 - 100,000	100	SURFACE	25	7 S	2.0 E	07	FALL RIVER	CHORD, ROY E.	DIANE A
SURFACE	1,000 - 100,000	50	SURFACE			0		FALL RIVER	BARKER, JAMES	DRIFTWOOD CANYON
SURFACE	<100	50	SURFACE			0		FALL RIVER	CHILDERS, H.F.	EAGLE AEPIC 1
SURFACE	100 - 1,000	200	SURFACE	24	7 S	2.0 E	07	FALL RIVER	CHORD, ROY E.	EDGE MONT 6
UNDERGRO	1,000 - 100,000	50	UNDERGRO	19	7 S	3.0 E	07	FALL RIVER	CHORD, ROY E.	GERTRUDE + FLORA
UNDERGRO	100 - 1,000	50	UNDERGRO	26	7 S	2.0 F	07	FALL RIVER	CHORD, ROY E.	GET ME WICH 1
UNDERGRO	>100,000	150	UNDERGRO	11	8 S	3.0 E	07	FALL RIVER	TENN. VALLEY AUTH	GOULD LEASES
UNDERGRO	1,000 - 100,000	50	UNDERGRO			0		FALL RIVER	ALBRIGHT, F+J	GREEN ACRES GROU
SURFACE	1,000 - 100,000	100	SURFACE	19	7 S	3.0 E	07	FALL RIVER	CHORD, ROY F.	GREEN SLIPPER 1
SURFACE	1,000 - 100,000	150	SURFACE	27	8 S	3.0 E	07	FALL RIVER	TENN. VALLEY AUTH	GULL LEASES 1-3
UNDERGRO	100 - 1,000	50	UNDERGRO	19	7 S	3.0 E	06	FALL RIVER	CHORD, ROY E.	HELEN
UNDERGRO	100 - 1,000	50	UNDERGRO			0		FALL RIVER	HANSON+HAUPTMAN	HELLS CANYON GRO
SURFACE	1,000 - 100,000	50	SURFACE	20	7 S	3.0 E	07	FALL RIVER	FAY + HEY	HEY + FAY 5 + 6

INACTIVE URANIUM MINES IN THE UNITED STATES
-SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 41

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** SOUTH DAKOTA (CONT'D) *****									
HI POCKETS 4	CHORD, ROY E.	FALL RIVER	20	7 S	3.0 E	07	SURFACE	100 - 1,000	50
HOLDUP 15-KADOS	CHORD, ROY E.	FALL RIVER	33	7 S	3.0 F	07	UNDERGRO	1,000 - 100,000	50
HOLDUP 2	CHORD, ROY E.	FALL RIVER	33	7 S	3.0 E	07	UNDERGRO	100 - 1,000	50
HOLDUP 22	CHORD, ROY E.	FALL RIVER			0		UNDERGRO	<100	50
HOT POINT GRP	CHORD, ROY E.	FALL RIVER	28	7 S	3.0 E	07	UNDERGRO	1,000 - 100,000	100
IMOGENE	CHORD, ROY E.	FALL RIVER	19	7 S	3.0 E	07	SURFACE	<100	0
JACK PINE 2	BONER, ROY	FALL RIVER			0		SURFACE	<100	50
JOANN	CHORD, ROY E.	FALL RIVER			0		SURFACE	<100	50
KELLOGG MINE	TENN, VALLEY AUTH	FALL RIVER	19	8 S	4.0 E	07	UNDERGRO	1,000 - 100,000	100
KING	CHORD, ROY E.	FALL RIVER			0		SURFACE	1,000 - 100,000	50
LAKOTA LODE 11	LAKOTA PROSPECT	FALL RIVER			0		SURFACE	<100	0
LAZYMAN'S LOCE	CHORD, ROY E.	FALL RIVER	20	7 S	3.0 E	07	SURFACE	<100	50
LION 1	TENN, VALLEY AUTH	FALL RIVER			0		SURFACE	1,000 - 100,000	50
LION-MC KNIGHT	TENN, VALLEY AUTH	FALL RIVER	12	8 S	3.0 F	07	SURFACE	1,000 - 100,000	50
LITTLE ANN	TRADE DOLLAR MNG	FALL RIVER			0		SURFACE	<100	0
LODE GROUP	HILL EVERETT	FALL RIVER			0		SURFACE	<100	0
LORA 1	LORD, R.H. & PARY J.	FALL RIVER			0		SURFACE	<100	50
LORD LEASE	BAXTER, F.M.	FALL RIVER			0		SURFACE	<100	50
LUCKY 1	CHORD, ROY E.	FALL RIVER	26	7 S	2.0 E	07	UNDERGRO	100 - 1,000	50
LUCKY STRIKE	EDGEMONT MINING	FALL RIVER			0		SURFACE	<100	0
LUCKY STRIKE	BETTENHAUSEN, RAY	FALL RIVER			0		SURFACE	100 - 1,000	0
LUCKY TOSS	STEVENS, EARL E.	FALL RIVER			0		SURFACE	<100	50
MARY LEASE	TENN, VALLEY AUTH	FALL RIVER	11	8 S	3.0 E	07	SURFACE	1,000 - 100,000	50
MARYJAC 1 9	TENN, VALLEY AUTH	FALL RIVER	10	8 S	3.0 F	07	UNDERGRO	100 - 1,000	50
MATIAS PEAK	TENN, VALLEY AUTH	FALL RIVER	34	7 S	3.0 E	07	SURFACE	1,000 - 100,000	50
MC KNIGHT 1	TENN, VALLEY AUTH	FALL RIVER			0		SURFACE	1,000 - 100,000	50
NIGGER GUICH	TENN, VALLEY AUTH	FALL RIVER	25	8 S	4.0 F	07	SURFACE	<100	50
OPHELIA	CHORD, ROY E.	FALL RIVER	19	7 S	3.0 F	07	SURFACE	100 - 1,000	50
PABST 3	TENN, VALLEY AUTH	FALL RIVER	12	8 S	3.0 E	07	SURFACE	1,000 - 100,000	50
PAT 2	TENN, VALLEY AUTH	FALL RIVER			0		SURFACE	<100	50
PATSY	CHORD, ROY E.	FALL RIVER			0		UNDERGRO	100 - 1,000	0
PAY DAY 3	TENN, VALLEY AUTH	FALL RIVER			0		SURFACE	<100	50
PFE WEE	CHORD, ROY E.	FALL RIVER			0		UNDERGRO	<100	0
PENNY-ITT 1	CHORD, ROY E.	FALL RIVER			0		UNDERGRO	100 - 1,000	50
PICTOGRAPH LODE	TENN, VALLEY AUTH	FALL RIVER	30	7 S	3.0 E	07	SURFACE	1,000 - 100,000	50
RABY LEASE	BETTENHAUSEN, MC	FALL RIVER			0		SURFACE	<100	50
RIDGE RUNNER	CHORD, ROY E.	FALL RIVER	26	7 S	3.0 E	07	SURFACE	1,000 - 100,000	100
RIPSNORTER 1	TENN, VALLEY AUTH	FALL RIVER	27	7 S	3.0 E	07	SURFACE	1,000 - 100,000	0
ROAD HOG	CHORD, ROY E.	FALL RIVER			0		SURFACE	1,000 - 100,000	0
ROWENA	CHORD, ROY E.	FALL RIVER	25	7 S	2.0 F	07	SURFACE	1,000 - 100,000	50
SEC. 11, 24 SEC	TENN, VALLEY AUTH	FALL RIVER			0		SURFACE	100 - 1,000	50
SEC. 36, 35- 3E	BETTENHAUSEN, RAY	FALL RIVER			0		SURFACE	100 - 1,000	100
SHANROCK	CHORD, ROY E.	FALL RIVER	19	7 S	3.0 F	07	UNDERGRO	1,000 - 100,000	100
SKINNEY SOB	SMITH, ROBERT B.	FALL RIVER			0		SURFACE	<100	50
SKIPPER 4	ALLIED EXPL. CO.	FALL RIVER			0		SURFACE	<100	0
SOTO + VERDE	SEAR LODGE MNG.	FALL RIVER	25	7 S	2.0 F	07	UNDERGRO	1,000 - 100,000	100
SOUTH VIEW GROUP	NICOLAYSEN + DAP.	FALL RIVER	25	7 S	2.0 E	07	SURFACE	100 - 1,000	50
SOUTHVIEW	WEIR + HENDERSON	FALL RIVER			0		SURFACE	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 42

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** SOUTH DAKOTA (CONT'D) *****									
STARLIGHT 2	M.I.L. UPAN, INC.	FALL RIVER			0		SURFACE	<100	0
TESS	EYRICH, HAROLD R	FALL RIVER			0		SURFACE	<100	50
TIN SHACK 6 -	BIEBER + MC LEOD	FALL RIVER			0		SURFACE	<100	50
TRAIL FRACTION	CHORD, ROY E.	FALL RIVER			0		UNDERGRO	1,000 - 100,000	0
TRAIL WIND	CHORD, ROY E.	FALL RIVER			0		SURFACE	100 - 1,000	50
VIRGINIA C	SUSQUEHANNA WEST	FALL RIVER	26	7 S	2,0 E	07	SURFACE	1,000 - 100,000	100
WASHBOARD	LORENZ BROTHERS	FALL RIVER	30	8 S	4,0 E	07	UNDERGRO	1,000 - 100,000	150
WESTERN EDGE	CHORD, ROY E.	FALL RIVER			0		SURFACE	<100	50
YELLOW CAT 1	CHORD, ROY E.	FALL RIVER	19	7 S	3,0 E	07	UNDERGRO	1,000 - 100,000	50
BILLY DOLF 1-4	HENDERSON+BLINNET	HARDING	36	22 N	5,0 E	07	SURFACE	1,000 - 100,000	50
BLUE JAY 8	HAIVALA BROS.	HARDING	34	21 N	5,0 E	07	SURFACE	<100	100
BOBCAT GROUP	SCHULL, LESTER	HARDING	30	18 N	8,0 E	07	SURFACE	<100	50
BONE-CHOPPY	PATTERSON, A.J.	HARDING	22	21 N	5,0 E	07	SURFACE	1,000 - 100,000	100
CALAMITY JANE 2	KRAUSE, MELVIN A.	HARDING	14	19 N	7,0 E	07	SURFACE	<100	0
CARBONATE GROUP	SUNDANCE PETRO	HARDING	29	21 N	5,0 E	07	SURFACE	<100	50
CRAZY 1 + 2	HANSON LLEWELYN	HARDING	18	16 N	9,0 E	07	SURFACE	1,000 - 100,000	50
DAISY MAY 5	MTN STATES MNG.	HARDING	24	17 N	1,0 E	07	SURFACE	<100	0
ELFANOR 1	BROWN, E.G.	HARDING	5	20 N	5,0 E	07	SURFACE	<100	0
FLAT TOP GROUP	CREVIER+LUDLOW+S	HARDING	27	22 N	6,0 E	07	SURFACE	1,000 - 100,000	100
HILLTOP GROUP	HANSON LLEWELYN	HARDING	12	16 N	9,0 E	07	SURFACE	1,000 - 100,000	50
JEFFERY LYNA	MILLER, M.D.	HARDING	23	21 N	5,0 E	07	SURFACE	<100	150
JIM GROUP	LEHAR+PATTERSON	HARDING	22	21 N	5,0 E	07	SURFACE	<100	50
LAST CHANCE GROU	KALINAS BROS.	HARDING	20	22 N	5,0 E	07	SURFACE	100 - 1,000	50
LINDAHL	LINDAHL	HARDING	27	22 N	6,0 E	07	SURFACE	1,000 - 100,000	50
LOVESOME PETE	HAIVALA, W.S.	HARDING	26	21 N	4,0 E	07	SURFACE	100 - 1,000	50
LUCKY STRIKE 2	LUCKY SIX MNG.	HARDING			0		SURFACE	<100	0
MC CURDY LEASF	MCCURDY+STENSETH	HARDING	21	22 N	6,0 E	07	SURFACE	<100	50
MOONSHINE	MOONSHINE URAN.	HARDING	29	16 N	9,0 E	07	SURFACE	<100	100
PICKPOCKET	MUNKRES, WM. L.	HARDING	35	22 N	5,0 E	07	SURFACE	1,000 - 100,000	50
QUAD 1 + 2	QUAD MNG CO	HARDING	13	16 N	9,0 E	07	SURFACE	<100	50
QUINN 6	QUINN, J.F. + R.N.	HARDING			0		SURFACE	<100	0
RELF GROUP	YOSS OIL CO.	HARDING	26	22 N	5,0 E	07	SURFACE	1,000 - 100,000	50
RILEY GROUP	OLSON + OAKLAND	HARDING	21	22 N	5,0 E	07	SURFACE	1,000 - 100,000	50
SNAKE EYE	STERNAD + HILL	HARDING	36	22 N	5,0 E	07	SURFACE	1,000 - 100,000	50
SNOOKS	PETERS, M.E.	HARDING	29	22 N	5,0 E	07	SURFACE	<100	50
SUSAN BECKY	RIMROCK MINING	HARDING	23	22 N	5,0 E	07	SURFACE	1,000 - 100,000	50
TRIO LODGE	TRIO MINING CO.	HARDING	18	16 N	9,0 E	07	SURFACE	100 - 1,000	50
BAVARIA	ROCKFORD THREE S	LAWRENCE			0		SURFACE	<100	0
STANLEY	ROCKFORD THREE S	LAWRENCE			0		SURFACE	<100	100
JELD 1	DENFVAN L.R.	PENNINGTON			0		SURFACE	<100	50
KAYLOR LEASE	BALDWIN, F.V.	PENNINGTON			0		SURFACE	<100	50
KOOL 10	ARNOLD, THOMAS F.	PENNINGTON			0		SURFACE	<100	50
RUBE 1	PICTOGRAPH MNG.	PENNINGTON			0		UNDERGRO	100 - 1,000	0
TED 1	MILLER, THEODORE	PENNINGTON			0		SURFACE	<100	50

***** TEXAS *****

SAUL BROS LEASE	BRISCOE CO, URAN	BRISCOE			0		SURFACE	<100	0
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INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 43

MINE NAME	CONTROLLING NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** TEXAS (CONT'D) *****									
TULE RANCH	LOVE MINING CO.	BRISCOL			0		SURFACE	<100	0
WILD HORSE MINE	JOHNSON, JAMES	PURNEY			0		SURFACE	<100	50
EUBANKS RANCH	RADIATION EXPL C	CROSBY			0		SURFACE	100 - 1,000	50
CROWN	RADIATION EXPL C	GARZA			0		SURFACE	<100	50
GARZA COUNTY EXP	ADAMS, JOHN	GARZA			0		SURFACE	<100	0
HILLSIDE	GARZA MINING CO.	GARZA			0		SURFACE	<100	50
MC ARTHUR N40SFC	MC ARTHUR STOCK	GARZA			0		SURFACE	<100	50
TWIN RATTLER	RADIATION EXPL C	GARZA			0		SURFACE	100 - 1,000	50
YELLOW CUB-SEC.4	RADIATION EXPL C	GARZA			0		SURFACE	100 - 1,000	50
DUNERSTADT	CITIES SERVICE	GONZALES			0		IN-SITU	100 - 1,000	0
SMITH E.F. 14	CONOCO-PIONEER	GONZALES			0		SURFACE	>100,000	100
BOSO-HACKNEY 2	SUSQUEHANNA WEST	KARNES			0		SURFACE	1,000 - 100,000	50
BRYSON LEASE	SUSQUEHANNA WEST	KARNES			0		SURFACE	1,000 - 100,000	100
FT. WORTH NATURAL	CONOCO-PIONEER	KARNES			0		SURFACE	1,000 - 100,000	100
HAASE F. (UL1491)	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	100
HACKNEY	SUSQUEHANNA WEST	KARNES			0		SURFACE	1,000 - 100,000	50
KELLNER LEASE 7	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	150
LAUREN LEASE 12	CONOCO-PIONEER	KARNES			0		SURFACE	1,000 - 100,000	100
LYSSY NIESTORY	SUSQUEHANNA WEST	KARNES			0		SURFACE	1,000 - 100,000	200
HANKA, J. (203)	CONOCO-PIONEER	KARNES			0		SURFACE	1,000 - 100,000	150
MC CRADY LEASE	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	200
MOCZYGENBA LSE12	CONOCO-PIONEER	KARNES			0		SURFACE	1,000 - 100,000	200
NIESCHWITZ-725	CONOCO-PIONEER	KARNES			0		SURFACE	1,000 - 100,000	100
NUHN LEASE	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	50
PAWELEK TRACT	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	100
PFEIL-WEIG. 303-1	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	150
SEAPCY-PA-FLEK	CONOCO-PIONEER	KARNES			0		UNKNOWN	1,000 - 100,000	0
SICKENIUS LEASE	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	50
SOUTH BRYSON LEA	SUSQUEHANNA WEST	KARNES			0		SURFACE	1,000 - 100,000	100
STOEL-NIES-BOD. 8	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	150
WEDDINGTON-SUS	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	150
WEDDINGTON-TEHNE	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	400
WRIGHT LEASE 7	CONOCO-PIONEER	KARNES			0		SURFACE	>100,000	200
WURZBACH	SUSQUEHANNA WEST	KARNES			0		SURFACE	1,000 - 100,000	50
CLAY WEST SLUPRY	DALCO-U S STEEL	LIVE OAK			0		IN-SITU	<100	0
E-S-B HEAP LEACH	CONOCO-PIONEER	LIVE OAK			0		HL-DRES	1,000 - 100,000	0
ESSE-SPOON-SCHP2	CONOCO-PIONEER	LIVE OAK			0		SURFACE	>100,000	200
KOPPLIN, K. 363	WYOMING MINERALS	LIVE OAK			0		SURFACE	>100,000	150
NABEL NEW	CONOCO-PIONEER	LIVE OAK			0		SURFACE	1,000 - 100,000	100
MC LEAN 1 + 2	EXXON CO USA	LIVE OAK			0		SURFACE	>100,000	50
RYAN	CONOCO-PIONEER	LIVE OAK			0		SURFACE	1,000 - 100,000	50
***** UTAH *****									
BEEHIVE 5	ATKINSON EXPL. CO	BEAVER			0		UNDERGRO	<100	50
C D P	CARTER-DALTON-PO	BEAVER			0		UNDERGRO	<100	50
DAISY	STAATS + LOWDER	BEAVER			0		UNDERGRO	100 - 1,000	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 44

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
GEYSER BASIN	GEYSER BASIN UPA	BEAVER			0		SURFACE	<100	0
IRON QUEEN	C + H MNG.	BEAVER			0		UNDERGRO	<100	100
LITTLE SISTERS	GRAEFF, RUSSELL	BEAVER			0		SURFACE	<100	50
MERCURY SEC 14	GARRICK, ELDRED M	BEAVER			0		SURFACE	<100	0
MYSTERY SNIFFER	MYSTERY SNIFFER	BEAVER	28	27 S	6,0	24	UNDERGRO	1,000 - 100,000	200
PRODUCER	ATKINSON EXPL.CO	BEAVER			0		UNDERGRO	100 - 1,000	50
WILLARD GROUP	DREYES, VERNON	BOX ELDER			0		SURFACE	<100	150
BLUE SKY 1	GRAHAM, FRED H.	DUCHESNE			0		SURFACE	<100	0
KEG	FEED, G.R.	DUCHESNE			0		UNDERGRO	<100	0
OH HENRY 1	ACADEMY URANIUM	DUCHESNE			0		SURFACE	<100	0
RAINBOW RIDGE GR	PARRISH-DALGLEI	DUCHESNE			0		SURFACE	<100	0
12A	WRIGHT, L.B.	EMERY			0		UNDERGRO	<100	0
4-CORNERS AREA D	ATLAS MINERALS	EMERY			0		DUMPS	1,000 - 100,000	0
A.G.1	ALBRECHT BROS,UP	EMERY			0		UNDERGRO	100 - 1,000	100
A.E.C. INCLINE	CONSOLIDATED URA	EMERY			0		UNDERGRO	100 - 1,000	0
A.E.C.13	CONSOLIDATED URA	EMERY			0		UNDERGRO	<100	250
A.E.C.6+8	UNION CARBIDE CP	EMERY			0		UNDERGRO	>100,000	100
A.E.C.7	CONSOLIDATED URA	EMERY			0		UNDERGRO	<100	150
ACEITE 2	CONSOLIDATED MOM	EMERY			0		UNDERGRO	1,000 - 100,000	50
ADDIT 2	CONSOLIDATED URA	EMERY			0		UNDERGRO	1,000 - 100,000	0
APEX 5	WOODWARD, D.W.	EMERY			0		UNDERGRO	<100	0
BAKER INCLINE	UNION CARBIDE CP	EMERY			0		UNDERGRO	1,000 - 100,000	100
BEAR CLAW	PACHIDE + CHIDES	EMERY			0		UNDERGRO	<100	0
BIG CHIEF 8	SPACE,FEED	EMERY			0		SURFACE	<100	0
BIG FLAT 1	HUNT, RED	EMERY			0		UNDERGRO	<100	0
BIG ROCK	URANIUS INC.	EMERY			0		UNDERGRO	<100	0
BIRTHDAY	WILCOX, BERT L.	EMERY			0		UNDERGRO	100 - 1,000	0
BLACK ROCK 1	FOOTE L.S.	EMERY			0		SURFACE	<100	50
BLANCO	STONE, E.F.	EMERY			0		SURFACE	<100	0
BLOCK C	PLATEAU URANIUM	EMERY			0		UNDERGRO	<100	0
BLOCK G	CONSOLIDATED URA	EMERY			0		UNDERGRO	<100	0
BLUE BELL 2	FARNSWORTH,EMERS	EMERY			0		SURFACE	<100	0
BLUE BUTTE	DAVIS + NIELSON	EMERY			0		SURFACE	<100	0
BLUE BUTTE 1	ADAMS,R.D.+L.A.	EMERY			0		SURFACE	<100	0
BLUE GOOSE	ASIMUS, C.C.	EMERY			0		UNDERGRO	100 - 1,000	0
BLUE JAY 1	JENSEN + KOURIS	EMERY			0		SURFACE	<100	50
BLUEBIRD	HUNT, KAY	EMERY			0		UNDERGRO	100 - 1,000	0
BLUEBIRD	HAYES H.+ M.	EMERY			0		UNDERGRO	100 - 1,000	0
BRIDGE 2-12	HANSEN, HOMER	EMERY			0		UNDERGRO	<100	0
BUCKHORN	KAINSTEAUX,HARVE	EMERY			0		SURFACE	<100	0
BUCKSKIN 2	FAUCETT + JONES	EMERY			0		SURFACE	<100	0
BULL 1	VALLEY MINING CO	EMERY			0		UNDERGRO	<100	0
BUZZARDS ROOST	HAMILTON J.C.	EMERY			0		SURFACE	<100	0
CAMP BIRD GROUP	UNION CARBIDE CP	EMERY			0		UNDERGRO	1,000 - 100,000	100
CANARY GROUP	LITTLE WILD HORSE	EMERY			0		UNDERGRO	100 - 1,000	0
CAT BIRD	WILLIAMS MINERAL	EMERY			0		UNDERGRO	100 - 1,000	0
CECILIAITE 1	ACERSON,ALFRED O	EMERY			0		SURFACE	<100	0
CEDAR MTN SEC 36	R HENDERSON	EMERY			0		SURFACE	100 - 1,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 45

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH	(CONT'D) *****							
CEDAR RIDGE 2	CEDAR RIDGE URAN	EMERY	0	SURFACE	<100	0		
CHERIE 1,2+3	KRUGER, RICHARD	EMERY	0	SURFACE	<100	100		
CISTERN 1 + 2	ENERGY FUELS NUC	EMERY	0	UNDERGRO	1,000 - 100,000	100		
CLIFF DWELLER 1	CHEETLE, E.V.	EMERY	0	UNDERGRO	<100	0		
CONRAD	CONRAD URAN, INC	EMERY	0	UNDERGRO	1,000 - 100,000	100		
CONSOLIDATED	ELLIOTT, JAMES O	EMERY	0	UNDERGRO	1,000 - 100,000	0		
CONTOUR 1	HOWARD, RUCLE	EMERY	0	UNDERGRO	1,000 - 100,000	0		
COPPER HEAD	BEHUNIN, JOANNE	EMERY	0	SURFACE	<100	0		
COTTONWOOD 1	NOYFS, FRED	EMERY	0	SURFACE	<100	50		
COUGAR 2	RAINBOW URANIUM	EMERY	0	SURFACE	<100	150		
CROSSROAD	CULLUM, CARROL	EMERY	0	UNDERGRO	1,000 - 100,000	50		
CROW	HAMILTON J.C.	EMERY	0	SURFACE	<100	0		
D, + R, CLAIMS	CONTIN, URAN, WYO.	EMERY	0	UNDERGRO	<100	0		
DALTON BROS 2	DALTON BROS.	EMERY	0	SURFACE	<100	0		
DARLENE 1-20	ACERSON, ALFRED O	EMERY	0	UNDERGRO	<100	50		
DELTA DUMP	PRESSLEY, G. M.	EMERY	0	DUMPS	1,000 - 100,000	0		
DESEPT MOON	PETRO NUCLEAR	EMERY	0	UNDERGRO	100 - 1,000	50		
DESOLATION	GA, GLEN A.	EMERY	0	SURFACE	<100	0		
DIXIE JOYCE	LINDQUIST, JOE	EMERY	0	SURFACE	<100	0		
DOLLY	ELLIOTT, JAMES O	EMERY	0	UNDERGRO	<100	0		
DONITE GROUP	ACERSON, ALFRED O	EMERY	0	UNDERGRO	100 - 1,000	200		
DRILL 2	INLAND RESOURCE	EMERY	0	SURFACE	<100	0		
EAGLE-BATTLESHIP	OUTWEST URAN+OIL	EMERY	0	UNDERGRO	100 - 1,000	150		
EAGLES NEST	HEATH, WILFORD	EMERY	0	SURFACE	<100	0		
ELMER E CASPER	CASPER, CAPLTON P	EMERY	0	SURFACE	<100	50		
FANTASTIC	JENKES + COVEY	EMERY	0	UNDERGRO	<100	0		
FISHER	HAMILTON J.C.	EMERY	0	SURFACE	<100	0		
FLINTYPE 2	STEEL, L.F.	EMERY	0	UNDERGRO	<100	100		
GREAT BASIN	U.S. URANIUM CORP	EMERY	0	UNDERGRO	<100	100		
GREEN DRAGON	ALBRECHT PROS, UR	EMERY	0	UNDERGRO	100 - 1,000	150		
GREEN VEIN	BRYON, ELDON	EMERY	0	SURFACE	<100	0		
GREEN VEIN 2	JENSEN, WALTER K.	EMERY	0	UNDERGRO	100 - 1,000	0		
GREEN VEIN 5	ENERGY FUELS NUC	EMERY	0	UNDERGRO	100 - 1,000	100		
GREEN VEIN 4	CAPITOL REEF UPA	EMERY	0	UNDERGRO	100 - 1,000	100		
HARD CLIMBING 10	PETITTI, JOHN J.	EMERY	0	UNDERGRO	<100	0		
HARDLUCK	DALE DILLON	EMERY	0	SURFACE	100 - 1,000	0		
HERTZ 1	DONALD HANZI	EMERY	0	UNDERGRO	1,000 - 100,000	100		
HIDDEN POINT	CHIDESTER, JOE	EMERY	0	SURFACE	<100	0		
INCL 6 DUMPS	PETRO NUCLEAR	EMERY	0	DUMPS	1,000 - 100,000	200		
INCLINE 10	PETRO NUCLEAR	EMERY	0	UNDERGRO	100 - 1,000	50		
INCLINE 11	ATLAS MINERALS	EMERY	0	UNDERGRO	100 - 1,000	150		
INCLINE 12	PETRO NUCLEAR	EMERY	0	UNDERGRO	1,000 - 100,000	50		
INCLINE 13	PETRO NUCLEAR	EMERY	0	UNDERGRO	100 - 1,000	100		
INCLINE 14 + 15	PETRO NUCLEAR	EMERY	0	UNDERGRO	1,000 - 100,000	50		
INCLINE 16	PETRO NUCLEAR	EMERY	0	UNDERGRO	100 - 1,000	100		
INCLINE 17	PETRO NUCLEAR	EMERY	0	UNDERGRO	1,000 - 100,000	50		
INCLINE 18	FOUR CORNERS OIL	EMERY	0	UNDERGRO	100 - 1,000	0		
INCLINE 19	FOUR CORNERS OIL	EMERY	0	UNDERGRO	100 - 1,000	100		

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 46

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
INCLINE 2-WED. RE	PETRO NUCLEAR	EMERY			0		UNDERGRO	100 - 1,000	100
INCLINE 20	FOUR CORNERS OIL	EMERY			0		UNDERGRO	<100	100
INCLINE 21	FOUR CORNERS OIL	EMERY			0		UNDERGRO	1,000 - 100,000	100
INCLINE 22	FOUR CORNERS OIL	EMERY			0		UNDERGRO	100 - 1,000	100
INCLINE 3	PETRO NUCLEAR	EMERY			0		UNDERGRO	1,000 - 100,000	250
INCLINE 4-5	PETRO NUCLEAR	EMERY			0		UNDERGRO	1,000 - 100,000	200
INCLINE 7-MINE 7	PETRO NUCLEAR	EMERY			0		UNDERGRO	100 - 1,000	100
INCLINE 8-DINOSA	ATLAS MINERALS	EMERY			0		UNDERGRO	1,000 - 100,000	200
INCLINE 9 NORTH	ATLAS MINERALS	EMERY			0		UNDERGRO	1,000 - 100,000	150
INCLINE 9 (RED 1-	ATLAS MINERALS	EMERY			0		UNDERGRO	1,000 - 100,000	150
JACK POT	CONSOLIDATED URA	EMERY			0		UNDERGRO	<100	0
JOHNNIE BOY 2	JOHNSON & JOHNSON	EMERY			0		SURFACE	<100	0
JOSHUA 1	JOSHUA MINING CO	EMERY			0		UNDERGRO	<100	0
KOH-BIRSEYE	U NEVA URAN. CORP	EMERY			0		UNDERGRO	<100	50
LAST CHANCE	LAST CHANCE MNG	EMERY			0		SURFACE	<100	0
LAST CHANCE 9	WRIGHT, WAYNE S.	EMERY			0		SURFACE	<100	0
LE DUC LEASE	UNION CARBIDE CP	EMERY			0		UNDERGRO	1,000 - 100,000	150
LITTLE EPMA 2	CHUTE CANYON UR.	EMERY			0		UNDERGRO	100 - 1,000	100
LITTLE JOE	CURTIS, RUSSELL	EMERY			0		UNDERGRO	<100	0
LITTLE LIL-JERRY	BLACK BULL MNG.	EMERY			0		UNDERGRO	1,000 - 100,000	300
LITTLE MIKE	GRANLICH MINERAL	EMERY			0		UNDERGRO	100 - 1,000	0
LITTLE SUSAN	ABS MINING CO.	EMERY			0		UNDERGRO	100 - 1,000	150
LITTLE WILDCAT	JACKSON, GEORGE B	EMERY			0		SURFACE	<100	0
LONE TREE GROUP	EMERY ENTERPRISE	EMERY			0		UNDERGRO	<100	0
LOOKOUT 1	HAMILTON, J.C.	EMERY			0		SURFACE	<100	0
LORETTA	WRIGHT, L.P.	EMERY			0		UNDERGRO	<100	0
LUCKY DOG 1	FOOTE L.S.	EMERY			0		SURFACE	<100	50
LUCKY SQUIRREL	GRANLICH MINERAL	EMERY			0		UNDERGRO	<100	0
LUCKY STRIKE	NIXON URANIUM CO	EMERY			0		UNDERGRO	100 - 1,000	50
LUCKY STRIKE 1,2	ASIMUS, C.C.	EMERY			0		UNDERGRO	100 - 1,000	0
LUCKY STRIKE GRO	FEHRON URAN MNG	EMERY			0		UNDERGRO	1,000 - 100,000	200
LUCKY STRIKE GRO	JACKSON, GEORGE B	EMERY	6	24 S	9,0 E	24	UNDERGRO	1,000 - 100,000	150
MAGPIE	JENKES M.K.	EMERY			0		SURFACE	<100	0
MAJESTIC 2	RAE JEAN MINING	EMERY			0		UNDERGRO	<100	150
MANILA GROUP	BEKUYIN, JOANNE	EMERY			0		SURFACE	<100	150
MARY JO 2	PETERSON, JESS	EMERY			0		SURFACE	<100	0
MAYFLOWER	U.S. URANIUM CORP	EMERY			0		UNDERGRO	<100	100
MAYFLOWER 1	CALDERWOOD, J.M.	EMERY			0		UNDERGRO	100 - 1,000	0
MAYFLOWER 32	U.S. URANIUM CORP	EMERY			0		UNDERGRO	<100	150
MESA 1	BENTLEY, JIM	EMERY			0		UNDERGRO	1,000 - 100,000	0
MISCELLANEOUS	FOUR CORNERS OIL	EMERY			0		UNDERGRO	<100	0
MORLENE 1	ATILSON, ORSON	EMERY			0		SURFACE	<100	0
MOUNTAIN KING	SKIDMORE, T.H.	EMERY			0		UNDERGRO	100 - 1,000	0
MOUNTAIN KING 1	CONSOLIDATED URA	EMERY			0		UNDERGRO	<100	0
MOUNTAIN KING 2	SKIDMORE, T.H.	EMERY			0		UNDERGRO	100 - 1,000	0
MOUNTAIN KING 3	SKIDMORE, T.H.	EMERY			0		UNDERGRO	1,000 - 100,000	0
MOUNTAIN KING GR	CONSOLIDATED URA	EMERY			0		UNDERGRO	1,000 - 100,000	0
MUCHO	ATLAS MINERALS	EMERY			0		UNDERGRO	100 - 1,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 47

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
MUDDY 7	CIMMARON MNG.	EMERY			0		UNDERGRO	100 - 1,000	100
MUDDY 8	HUNTER L.T.	EMERY			0		UNDERGRO	100 - 1,000	0
MUDDY RIVER	HUNTER L.T.	EMERY			0		SURFACE	<100	0
NO. 2	FRANDSEN, LAVERN	EMERY			0		SURFACE	<100	0
NORMA JOE	JOHNSON MELVIN C	EMERY			0		UNDERGRO	<100	0
NORTH MESA 1	UNION CARBIDE CP	EMERY			0		UNDERGRO	1,000 - 100,000	150
NORTH MESA 12	CONSOLIDATED URA	EMERY			0		UNDERGRO	<100	0
NORTH MESA 17	CONSOLIDATED URA	EMERY			0		UNDERGRO	<100	0
NORTH MESA 6	CONSOLIDATED URA	EMERY			0		SURFACE	<100	0
NORTH MESA 9	UNION CARBIDE CP	EMERY			0		UNDERGRO	1,000 - 100,000	50
NORTH MOUNTAIN 2	WILKERSON, THOMAS	EMERY			0		SURFACE	<100	300
NORTH SLOPE GROU	JACOBSON, LEO D.	EMERY			0		UNDERGRO	100 - 1,000	0
NORTH STAR	HAYS, MONTY	EMERY			0		SURFACE	<100	0
PACIFIC VENTURES	BAY WEST INC.	EMERY			0		UNDERGRO	1,000 - 100,000	0
PAY DAY	AGAARD, ELMER R. A	EMERY			0		UNDERGRO	100 - 1,000	100
PEARCE + KELLEY	FOUR CORNERS OIL	EMERY			0		UNDERGRO	<100	0
POSSIBILITY 1	HEATH, WILFORD	EMERY			0		SURFACE	<100	0
RABBIT 1	GALLAGHER+PIECRUCC	EMERY			0		UNDERGRO	100 - 1,000	0
RAINY DAY	GREEN FORK MNG.	EMERY			0		UNDERGRO	100 - 1,000	250
RED 1,2+5	PETRO NUCLEAR	EMERY			0		UNDERGRO	1,000 - 100,000	250
RED BONE 10	VITRO MINERALS	EMERY			0		UNDERGRO	100 - 1,000	100
RED BONE 3	WELCH MINING CO	EMERY			0		UNDERGRO	<100	100
RED BONE 4	PETRO NUCLEAR	EMERY			0		UNDERGRO	1,000 - 100,000	150
RED BUTTE 1-4	ADAMS URANIUM CO	EMERY			0		SURFACE	<100	100
RED CLIFF 3	WINDER, WALLACE	EMERY			0		SURFACE	<100	0
RED FACT	KNIGHT, MYRON	EMERY			0		SURFACE	<100	50
RED FORT	HEATH, WILFORD	EMERY			0		SURFACE	<100	0
RED SEEPS	MOORE URANIUM CO.	EMERY			0		UNDERGRO	<100	0
REX 1	UNION CARBIDE CP	EMERY			0		UNDERGRO	1,000 - 100,000	0
RYAN 11	MAYARD + RYAN	EMERY			0		SURFACE	<100	0
SAHARA 16	EMERY FIELDS NUC	EMERY			0		UNDERGRO	1,000 - 100,000	250
SC. SEC. 36-18S-10	RAM MINING CO	EMERY			0		SURFACE	<100	100
SCH SEC 2	ACEPSON, ALFRED D	EMERY			0		UNDERGRO	100 - 1,000	100
SCH. SEC. 2+16	ASIMUS, C.C.	EMERY			0		UNDERGRO	<100	0
SOUTH TEMPLE	SKIDMORE, T.H.	EMERY			0		UNDERGRO	<100	0
SOUTHERN CROSS	WINN, JOHN	EMERY			0		UNDERGRO	<100	0
SQUIRREL	GRACKLICK MINERAL	EMERY			0		SURFACE	<100	0
STILSON 5	STILSON, DOYLE	EMERY			0		SURFACE	<100	0
TEMPLE 1	CONSOLIDATED URA	EMERY			0		SURFACE	<100	0
TOTEM POLE 1	BABBLE, GORDON	EMERY			0		SURFACE	<100	0
TOUGH LUCK 1	CONSOLIDATED URA	EMERY			0		UNDERGRO	<100	0
TWIN PEAKS	WINN, JOHN	EMERY			0		SURFACE	<100	0
UNION GULF 1	THOMAS, HARVEY	EMERY			0		UNDERGRO	100 - 1,000	100
UNKNOWN	NUCLEAR MINES IN	EMERY			0		UNDERGRO	<100	0
VAN BEG 2	KNIGHT, H.R.	EMERY			0		UNDERGRO	100 - 1,000	0
VANADIUM KING 4+	ROBINSON, ELWIN	EMERY			0		UNDERGRO	1,000 - 100,000	0
VARIOUS	CONSOLIDATED URA	EMERY			0		UNDERGRO	1,000 - 100,000	100
VENTURE	GREEN HORNET MNG	EMERY			0		SURFACE	<100	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 48

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
VIRGINIA VALLEY	LAW, GLEN A.	EMERY			0		SURFACE	100 - 1,000	100
VITRO DUMP	VITRO CHEMICAL	EMERY			0		DUMPS	100 - 1,000	0
WEDDING RFL	H.F.W. MINERALS	EMERY			0		UNDERGRO	1,000 - 100,000	150
WHITE STAR 1-10	B + B MINING CO.	EMERY			0		UNDERGRO	100 - 1,000	50
WICKIUP	ARAGON, A.R.	EMERY			0		UNDERGRO	<100	200
WILD HORSE 22,27	CISTERN MINING	EMERY			0		UNDERGRO	100 - 1,000	100
WILDCAT	JENSEN + JACKSON	EMERY			0		SURFACE	<100	0
WILLOW SPRINGS	WINTERS + JONES	EMERY			0		SURFACE	<100	0
WINDY	BLACK DRAGON URA	EMERY			0		SURFACE	<100	0
AGATE GROUP	UNITED ENERGY	GARFIELD			0		UNDERGRO	1,000 - 100,000	100
ALLEN 3	KAISER URAN.CO.	GARFIELD			0		UNDERGRO	<100	150
B + M CLAIM 1	UNKNOWN	GARFIELD			0		SURFACE	<100	50
BEAR CANYON	HEIMECKE PROS.	GARFIELD			0		UNDERGRO	<100	100
BETTY JACK	FEDERAL RESOURCE	GARFIELD			0		SURFACE	<100	100
BIG TREE	UNIVERSAL URAN.C	GARFIELD			0		SURFACE	<100	300
BLACK CAT 2	20TH CENTURY POW	GARFIELD			0		SURFACE	<100	100
BLACK MOUNTAIN	MC ELROY, JOHN	GARFIELD			0		SURFACE	<100	100
BLACK WIDOW	HOLMES, J.MARK	GARFIELD			0		UNDERGRO	100 - 1,000	150
BLITZ	ENERGY FUELS NUC	GARFIELD			0		UNDERGRO	1,000 - 100,000	50
BLUE BIRD	ACHE URANIUM MNG	GARFIELD			0		UNDERGRO	<100	50
BLUE BIRD 4	RAINBOW URANIUM	GARFIELD			0		SURFACE	<100	150
BLUE GOOSE	BLUE GOOSE MNG	GARFIELD			0		SURFACE	<100	100
BROWN TOP	DAVISON, EMERY L	GARFIELD			0		UNDERGRO	<100	150
BUFF	CHRISTENSEN, ARMI	GARFIELD			0		UNDERGRO	<100	200
BULL 4	WRIGHT, B.K.	GARFIELD			0		UNDERGRO	<100	150
BUST 2	HUNT, RFO	GARFIELD			0		UNDERGRO	<100	50
BUTTLAR WASH 1	SMITH + STEWART	GARFIELD			0		SURFACE	<100	100
CARRON 1	EKKER, HAROLD C.	GARFIELD			0		SURFACE	<100	100
CEDAR POINT 2	SILVER BELL INDU	GARFIELD			0		UNDERGRO	100 - 1,000	100
CIRCLE CLIFFS	COOPER + BROWN	GARFIELD			0		SURFACE	<100	100
COFFEE ROYAL 1 +	BROWN, A.P., ASSOC	GARFIELD			0		SURFACE	<100	100
CONGRESS 14	BULLDOG MINING C	GARFIELD			0		UNDERGRO	<100	250
CONGRESS 22	GUTHRIE + ASSOC.	GARFIELD			0		UNDERGRO	<100	250
CONGRESS 24	GUTHRIE + ASSOC.	GARFIELD			0		UNDERGRO	<100	300
CONGRESS 25	GUTHRIE + ASSOC.	GARFIELD			0		UNDERGRO	<100	300
CONGRESS 47	GENERAL UTILITIES	GARFIELD			0		SURFACE	<100	100
CONGRESS 47+48+5	EKKER, HAROLD C.	GARFIELD			0		SURFACE	<100	150
CONGRESS 53	INDUSTRIES+MINES	GARFIELD			0		UNDERGRO	<100	200
CROW-LUCKY STATE	SMITH, C.C.	GARFIELD			0		UNDERGRO	<100	200
DEAN 4	GENERAL UTILITIES	GARFIELD			0		SURFACE	<100	100
DEEP CANYON 1	EKKER, HAROLD C.	GARFIELD			0		SURFACE	100 - 1,000	150
DEEP CANYON 2	EKKER, HAROLD C.	GARFIELD			0		SURFACE	<100	150
DENNEA LOU	HERCULES URANIUM	GARFIELD			0		SURFACE	<100	50
DIRTY SHANE 7	DIRTY SHANE MNG	GARFIELD			0		SURFACE	<100	200
DOMS 1	EKKER, RITTER	GARFIELD			0		SURFACE	<100	50
DREAM	BROWN + SONS CO	GARFIELD			0		UNDERGRO	<100	100
DREAM CLAIM 1	DREAM MINING CO	GARFIELD			0		SURFACE	<100	100
DUKE MINE	FRANDSEN BROS.	GARFIELD			0		SURFACE	<100	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 49

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
EAGLE GROUP	EKKER, NOPACE	GARFIELD			0		UNDERGRO	100 - 1,000	250
EAST COVE 9	MONTY POT URAN C	GARFIELD			0		SURFACE	<100	100
EDNA GROUP	EKKER, HAROLD C.	GARFIELD			0		SURFACE	<100	150
ELLEN 1	HUNT, PEO	GARFIELD			0		UNDERGRO	100 - 1,000	100
ELLSWORTH 1	PROSPECTORS, INC.	GARFIELD			0		SURFACE	<100	100
ELORA 1	EKKER, RITTER	GARFIELD			0		SURFACE	100 - 1,000	50
ELSWORTH 1	EKKER, HAROLD C.	GARFIELD			0		SURFACE	<100	50
ERMA MAE 1 + 2	FAIRBANKS, NEIL	GARFIELD			0		SURFACE	<100	50
EUNICE 1	WOOD, HOWARD P.	GARFIELD			0		SURFACE	<100	50
FIVE STAR GROUP	DUNFIELD, MABLE P.	GARFIELD			0		SURFACE	<100	100
FREMONT ASSOC 1	FREMONT ASSOC. D.	GARFIELD			0		SURFACE	<100	200
GARFIELD	STUD HORSE BUTTE	GARFIELD			0		UNDERGRO	<100	200
GENERAL	EKKER, HAROLD C.	GARFIELD			0		SURFACE	<100	50
H., H. 3-5+15	BAKER, JOE	GARFIELD			0		UNDERGRO	100 - 1,000	50
HANSEN 4	HANSEN, J. VERN	GARFIELD			0		SURFACE	<100	100
HARD ROCK	HOLMES, J. MARK	GARFIELD			0		SURFACE	<100	150
HARD SCRABBLE 1	HUNT, KAY	GARFIELD			0		SURFACE	<100	150
HENRY 12	BROWN, J. F.	GARFIELD			0		SURFACE	<100	100
HENRY MOUNTAINS	HUNT, KAY	GARFIELD			0		UNDERGRO	1,000 - 100,000	50
HOPE	HOPE MINING CO	GARFIELD			0		UNDERGRO	100 - 1,000	200
HOPE-CIRCLE CLIF	STUD HORSE BUTTE	GARFIELD			0		UNDERGRO	100 - 1,000	50
HOT SHOT+MT. SHT.	HOWARD S. K.	GARFIELD			0		UNDERGRO	100 - 1,000	100
JAKE	BUCHANAN, SANDY L.	GARFIELD			0		SURFACE	<100	50
JIM DANDY	HUNT, REO	GARFIELD			0		UNDERGRO	100 - 1,000	50
JUNE BELL GROUP	FOOTE MINFRALS	GARFIELD			0		UNDERGRO	100 - 1,000	50
KING GROUP	HUNT, KAY	GARFIELD			0		SURFACE	1,000 - 100,000	50
LAST CHANCE 1	UTAH ENERGY CO.	GARFIELD			0		UNDERGRO	<100	150
LITTIE SUE	ALLEN PAUL K	GARFIELD			0		SURFACE	<100	50
LODE	C-H MINING	GARFIELD			0		SURFACE	<100	50
LONE F.	MORTENSEN, NEAL J.	GARFIELD			0		UNDERGRO	<100	50
LONE-EAGLE 1-9	SHOOTERING CREEK	GARFIELD			0		UNDERGRO	100 - 1,000	100
LOUISE 1	JUSTENSEN, LEE	GARFIELD			0		SURFACE	<100	100
LUCY DAY	WARDEN+TWITCHELL	GARFIELD			0		UNDERGRO	<100	100
LUCKY STRIKE 1-7	JENSEN + JACKSON	GARFIELD			0		UNDERGRO	1,000 - 100,000	100
LUCKY STRIKE 10	KLIPPEL, BEN F.	GARFIELD			0		UNDERGRO	1,000 - 100,000	300
LUCKY STRIKE 8 +	HYDRO JET SERVIC	GARFIELD			0		UNDERGRO	1,000 - 100,000	150
LUCKY STRIKE 1 9+	EKKER, HAROLD C.	GARFIELD			0		SURFACE	<100	50
MAJESTIC + MODOC	P+P ASSOCIATES	GARFIELD			0		UNDERGRO	100 - 1,000	150
MAJESTIC 1	BASTIAN, G. A.	GARFIELD			0		SURFACE	<100	50
MAUD 9	NOVELLE, E. D.	GARFIELD			0		UNDERGRO	<100	100
MERRI MAC GROUP	WOOD, HOWARD P.	GARFIELD			0		UNDERGRO	100 - 1,000	50
MIDNIGHT	TANNER + TROUT	GARFIELD			0		SURFACE	<100	100
MIDNIGHT 1	WRIGHT, W. F.	GARFIELD			0		SURFACE	<100	50
MISFIT 1	SELF, ERROL	GARFIELD			0		SURFACE	<100	50
MODOC 2	RIFER EKKER	GARFIELD			0		UNDERGRO	100 - 1,000	100
NANCY 2	B. + A. MINING CO.	GARFIELD			0		SURFACE	<100	50
NANCY JANE	BREI MINING CO	GARFIELD			0		SURFACE	<100	50
NAVAJO 1	HUNT, REO	GARFIELD			0		UNDERGRO	100 - 1,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 30

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
OIL BEEP 8	EKKER, HAROLD C.	GARFIELD			0		SURFACE	<100	50
PENNEL 11	HUNT, RED	GARFIELD			0		UNDERGRO	100 = 1,000	150
PHYLLIS	EKKER, RITTER	GARFIELD			0		UNDERGRO	<100	200
QUEEN OF SHEBA	JENSEN, WESLEY	GARFIELD			0		SURFACE	<100	0
RAINY DAY	ACHE URANIUM MNG	GARFIELD			0		UNDERGRO	1,000 = 100,000	350
RATS NEST	PROSPECTORS, INC.	GARFIELD			0		UNDERGRO	<100	100
RATTLESNAKE	DAVIS, RAY	GARFIELD			0		UNDERGRO	100 = 1,000	0
RED CLIFF	TANNER, RUTHERFORD	GARFIELD			0		SURFACE	<100	50
RENE BOB	SEVEY, JAMES	GARFIELD			0		SURFACE	<100	50
ROARING ATON	JOHNSON + ROSS	GARFIELD			0		UNDERGRO	<100	100
ROBBERS PODST	JENSEN MC MINING	GARFIELD			0		UNDERGRO	<100	0
ROSE AN' 1	APMSTRONG, LYLE C	GARFIELD			0		SURFACE	<100	100
ROSLAND 2	EKKER + SANDERS	GARFIELD			0		SURFACE	<100	50
ROY DEL GROUP	FEDERAL RESOURCE	GARFIELD			0		SURFACE	<100	50
RUBILEE 2	ARCO URANIUM CO	GARFIELD			0		UNDERGRO	<100	100
S.Y. 36	GUTHRIE + ASSOC.	GARFIELD			0		UNDERGRO	<100	100
SALINA 2	MATHIS, R.G.	GARFIELD			0		SURFACE	<100	0
SEC. 36-32S-11E	SHLUCCO, INC.	GARFIELD			0		UNDERGRO	100 = 1,000	0
SEC. 8-32S-11E(T-8)	P+P ASSOC	GARFIELD			0		UNDERGRO	100 = 1,000	0
SHOOTPRING 1 + 2	U NEVA URAN. CORP	GARFIELD			0		UNDERGRO	<100	50
SILVER FALLS	KINDS, FRED	GARFIELD			0		SURFACE	<100	100
SILVER LEAF	KINDS, FRED	GARFIELD			0		SURFACE	<100	100
SILVER ROCK GROUP	SELF, EMROL	GARFIELD			0		SURFACE	<100	50
SPRING CANYON	EKKER, HORACE	GARFIELD			0		SURFACE	<100	50
STRAIGHT CREEK	ENERGY FUELS NUC	GARFIELD			0		UNDERGRO	100 = 1,000	50
SUNRISE 1-6	S. + A. MINING CO.	GARFIELD			0		SURFACE	<100	50
SYLVIA 2 + 14	CREEK FARM KING J	GARFIELD			0		UNDERGRO	<100	150
TENDERFOOT	HULEN, BRADLEY	GARFIELD			0		UNDERGRO	<100	50
TRACKYTE 12	MILLARD GARDNER	GARFIELD			0		UNDERGRO	<100	50
TRACKYTE 17, 18+1	DENNIS EKKER	GARFIELD			0		UNDERGRO	1,000 = 100,000	100
TRACKYTE 9 10	CHESTNUT + HARTLEY	GARFIELD			0		UNDERGRO	100 = 1,000	100
TRAIL CANYON 1	EKKER, HORACE	GARFIELD			0		SURFACE	<100	100
TRU ANN GROUP	PROSPECTORS, INC.	GARFIELD			0		UNDERGRO	100 = 1,000	50
TUNNISON	ACHE URANIUM MNG	GARFIELD			0		UNDERGRO	<100	150
VA LEAR	HUNT, KAY	GARFIELD			0		SURFACE	<100	50
WALTER GROUP	DAY, ORSON	GARFIELD			0		UNDERGRO	100 = 1,000	150
WAR PAINT 1	WALLACE, LYMAN	GARFIELD			0		SURFACE	<100	50
WEST BANK	ROGERS, R.S.	GARFIELD			0		SURFACE	<100	50
WHITE BULL GROUP	TYRRE, CHAS. H. EST	GARFIELD			0		UNDERGRO	100 = 1,000	150
WILLIE 1	URANIUM SERVICE	GARFIELD			0		SURFACE	<100	50
WOODRUFF GROUP	ENERGY FUELS NUC	GARFIELD			0		UNDERGRO	100 = 1,000	50
YELLOW CAT 1-3	NOVILL, F.D.	GARFIELD			0		UNDERGRO	<100	50
YELLOW JACKET	BARNEY + HOWARD	GARFIELD			0		UNDERGRO	100 = 1,000	150
YELLOW PAINT	ATLAS-AMAX	GARFIELD			0		UNDERGRO	100 = 1,000	100
3 RS 5	BALSLEY, TOM	GRAND			0		UNDERGRO	100 = 1,000	0
A GROUP	URANIUM PROD CO	GRAND	12	26 S	17.0 E	24	UNDERGRO	1,000 = 100,000	50
AEC GROUP	BUTTERLAND + SUTH	GRAND			0		SURFACE	<100	0
AUQUA	HARRIS MINING CO	GRAND		21 S	17.0 E	24	SURFACE	<100	150

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 51

MIKE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
AVIS-HORN	UNION CARRIDE CP	GRAND	1	25 S	25.0 E	24	UNDERGRO	1,000 - 100,000	100
BARE SPOT	SWARTZ, HAROLD H	GRAND	5	23 S	21.0 E	24	SURFACE	<100	50
BEAR POINT	LILE, GLEN	GRAND	29	25 S	25.0 E	24	SURFACE	100 - 1,000	0
BERTHA + FALCON	CLINE + CO.	GRAND	34	23 S	20.0 F	24	UNDERGRO	100 - 1,000	100
BIG BUCK 17	LAHMEPT+BURGESS	GRAND		26 S	17.0 F	24	SURFACE	<100	150
BIG LOUIE 1	ANCAN	GRAND	21	24 S	26.0 E	24	UNDERGRO	1,000 - 100,000	0
BLACK APE 1 + 2	TERRA MARINE MNG	GRAND	7	23 S	22.0 E	24	UNDERGRO	1,000 - 100,000	50
BLUE BOY	HARRICK, FARL	GRAND			0		SURFACE	<100	0
BLUE CHIEF	NICHOLS, CARL A.	GRAND			0		SURFACE	<100	0
BLUE CHIEF	CONGER W.R.	GRAND			0		SURFACE	<100	50
BORRY 1	CIRCLE DOT INC.	GRAND			0		UNDERGRO	<100	5
BRUMLEY BASIN 1	STRATEGIC MRLS	GRAND			0		UNDERGRO	<100	0
BUCKHORN 1	COLVIN +BRESNAHA	GRAND	3	23 S	25.0 E	24	SURFACE	<100	250
BULL CANYON GROU	ADAIR, IVOR	GRAND	4	26 S	20.0 E	24	UNDERGRO	100 - 1,000	0
C D WHEELER CLAL	BLACK ROCK URAN.	GRAND			0		UNDERGRO	<100	0
CAPTAIN JACK	UNION CARRIDE CP	GRAND	26	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	200
CAPTAIN JINKS	UNION CARRIDE CP	GRAND			0		UNDERGRO	<100	0
CEDAR POINT 2	FOSTER, LFONA	GRAND	33	24 S	26.0 E	24	UNDERGRO	1,000 - 100,000	200
CIE DOG	COMANCHE URAN.CO	GRAND	29	22 S	20.0 E	24	UNDERGRO	<100	0
CLEANUP MATERIAL	LANCE HARDMAN	GRAND			0		MISC.-PB	<100	0
CONSTELLATION 3	SECREST, ART O.	GRAND	12	23 S	21.0 E	24	SURFACE	<100	0
COPPER 2 B	K.+ J. MINING CO.	GRAND	11	25 S	20.0 E	24	SURFACE	<100	0
CORRAL CUP	THORNBERG MNG.	GRAND			0		DUMPS	1,000 - 100,000	0
COTTONWOOD 6	URANIUM PROO CO	GRAND	25	23 S	17.5 F	24	UNDERGRO	1,000 - 100,000	650
COUNTER 1	ABERNATHY, JESS	GRAND			0		SURFACE	<100	0
DAHICE	UNKNOWN CONTROLR	GRAND	26	25 S	17.0 E	24	UNDERGRO	100 - 1,000	200
DILLERS 1 + 3	GOLD CLOUD URAN	GRAND			0		UNDERGRO	<100	100
DINE	MC NEIL + HUGHES	GRAND	24	24 S	20.0 F	24	SURFACE	<100	0
DINE	STOCKS, JOHN	GRAND	24	24 S	20.0 E	24	UNDERGRO	100 - 1,000	0
DISCOVERY	MFRAB CO.	GRAND			0		UNDERGRO	<100	150
DRAGON FLY 4	GREAT WESTERN UR	GRAND		26 S	18.0 E	24	UNDERGRO	<100	50
EDNA 3	COLUMBIA URAN.CO	GRAND	15	25 S	17.5 E	24	UNDERGRO	100 - 1,000	0
EFFIE F.	UNION CARRIDE CP	GRAND	34	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	200
END OF TRAIL	BLACK CAT EXPL.C	GRAND	10	5 S	91.0 W	24	SURFACE	<100	0
F.W. 3	UNION CARRIDE CP	GRAND	35	24 S	25.0 F	24	UNDERGRO	1,000 - 100,000	250
FALCON 1	HASLEY, JACK	GRAND	27	23 S	20.0 E	24	SURFACE	<100	100
FLAT TOP	WILSON, R.P.	GRAND	25	22 S	22.0 E	24	UNDERGRO	100 - 1,000	0
FOOLS LUCK	SWARTZ, HAROLD H	GRAND	10	23 S	21.0 E	24	SURFACE	<100	0
FOSTER AREA	UNION CARRIDE CP	GRAND	34	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	200
FREEDA 2	MONARCH URANIUM	GRAND	9	27 S	23.0 F	24	UNDERGRO	<100	0
FRFEPORF	UNION CARRIDE CP	GRAND	1	25 S	25.0 E	24	UNDERGRO	<100	0
GAP	WOODS, DUKE B.	GRAND			0		SURFACE	<100	0
GEIGER 1	EMERSON EL +ASSO	GRAND			0		SURFACE	<100	0
GLADSTONE	UTAH ALLOY ORFS	GRAND			0		UNDERGRO	<100	0
GLORY HOLE	URADO MINING	GRAND	24	23 S	24.0 E	24	SURFACE	<100	0
GREAT DANE 1	ABRAHAMSON, J.F.	GRAND			0		UNDERGRO	100 - 1,000	0
GREEN LIZARD 1	SMITH + LIKES	GRAND			0		SURFACE	<100	0
GREEN TREE	MCGEEHEE, L.	GRAND			0		UNDERGRO	<100	150

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 33

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
GREY FAULT	BALCONES EXPL. DR	GRAND	21	26 S	20.0 E	24	UNDERGRO	100 - 1,000	200
HEBER 1	ZUEFELT C. & P.	GRAND	12	29 S	21.0 E	24	SURFACE	<100	0
HEY JOE	JIM TWITCHELL	GRAND	16	25 S	17.0 E	24	UNDERGRO	1,000 - 100,000	50
HILL TOP 1	DE ROSS, ROSE MAP	GRAND	4	22 S	14.0 E	24	UNDERGRO	100 - 1,000	50
MIROHITO	GRAMLICH MINERAL	GRAND			0		UNDERGRO	<100	0
MOPE	IRVINE, DON J.	GRAND			0		SURFACE	<100	0
MORSETHIEF 7 MIN	UNIVERSAL URAN. C	GRAND		26 S	17.0 E	24	SURFACE	<100	150
INCLINE 1,2,3,4+	ATLAS MINERALS	GRAND	22	2 S	14.0 E	24	UNDERGRO	1,000 - 100,000	0
INGERSOLL 13	SUNSET MINES INC	GRAND			0		SURFACE	<100	0
JIM	V. & L. DEVELOPMEN	GRAND	10	25 S	17.5 E	24	UNDERGRO	<100	0
JOHN	UNKNOWN CONTROL	GRAND	1	23 S	21.0 E	24	UNDERGRO	1,000 - 100,000	200
JOHNNIE A. 1	DAVIS, LEON L.	GRAND			0		SURFACE	<100	0
JUANITA GROUP	BARTON, C. S.	GRAND	23	23 S	21.0 E	24	SURFACE	<100	0
JUNCTION	MOAR URANIUM CO.	GRAND			0		SURFACE	<100	50
JUNIPER GROUP	PETRO NUCLEAR	GRAND	26	22 S	22.0 E	24	UNDERGRO	100 - 1,000	50
KELLY 1	ABERNATHY, JESS	GRAND			0		UNDERGRO	<100	50
KLONDIKE	KNIGHT, N. B. JR	GRAND	15	25 S	20.0 E	24	UNDERGRO	100 - 1,000	0
LAST INDIAN 1	LEWIS, J. W.	GRAND			0		UNDERGRO	<100	0
LENA 1	MINERAL HILL URA	GRAND			0		UNDERGRO	<100	0
LEWIS A	UNION CARBIDE CP	GRAND	26	24 S	24.0 E	24	UNDERGRO	100 - 1,000	0
LITTLE EYE	TONY PENE	GRAND	6	22 S	22.0 E	24	UNDERGRO	1,000 - 100,000	50
LITTLE LIZZARD	JONES, E. L.	GRAND			0		UNDERGRO	<100	0
LITTLE MEDICINE	WEAVER, CAPL	GRAND			0		UNDERGRO	<100	0
LONE PINE 2	FOSTER & WAREHAM	GRAND	28	24 S	26.0 E	24	SURFACE	<100	0
LOOKOUT 1	CATO, CHARLES	GRAND			0		SURFACE	<100	0
LOUISE GROUPS	BROWN & DARROWS	GRAND			0		UNDERGRO	100 - 1,000	0
LUCKY DAY	BUTHERLAND & SUTH	GRAND	7	23 S	22.0 E	24	SURFACE	<100	0
M GROUP	AIRBORNE PROSPECT	GRAND	32	26 S	20.0 E	24	UNDERGRO	1,000 - 100,000	0
MATCHLESS	DEVELOPS CO.	GRAND		26 S	18.0 E	24	UNDERGRO	100 - 1,000	50
MAYBELL	PENE, ANTHONY JR	GRAND			0		SURFACE	<100	0
MERCURY 1	BEHUNIN, JOANNE	GRAND			0		SURFACE	<100	0
MILE HIGH TANK F	INDUSTRIES & MINES	GRAND	27	26 S	20.0 E	24	UNDERGRO	100 - 1,000	0
MINERAL 10-BLK J	JERRY STOCKS	GRAND			0		UNDERGRO	1,000 - 100,000	50
MINERAL 11	SHUWAY & WESTHO	GRAND	15	26 S	18.0 E	24	SURFACE	<100	0
MINERAL POLAR 8+	ATLAS-AMAX	GRAND	35	24 S	25.0 E	24	UNDERGRO	<100	400
MINERAL-RIG BLUE	INTERMOUNTAIN	GRAND			0		UNDERGRO	100 - 1,000	100
MISTAKE	KNIGHT, N. B.	GRAND	7	26 S	24.0 E	24	SURFACE	<100	0
MORRIS CLAIMS	LEE MORRIS	GRAND			0		UNDERGRO	100 - 1,000	0
NEW ARMSTRONG	UNION CARBIDE CP	GRAND	3	25 S	25.0 E	24	UNDERGRO	100 - 1,000	0
NORTH MINE 2	THORNBURG URAN P	GRAND			0		UNDERGRO	<100	0
NORTH SLOPE 2	JENSEN & KOURIS	GRAND			0		SURFACE	<100	0
OLD MC COY	WRIGHT, L. P.	GRAND			0		SURFACE	<100	0
OLD SPOON	TURNER BROS	GRAND			0		SURFACE	<100	0
OSCAR 1	ABERNATHY, JESS	GRAND			0		SURFACE	<100	0
OXIDE 1	CATO, CHARLES	GRAND			0		SURFACE	<100	0
P. F. G. E.	UNION CARBIDE CP	GRAND	1	24 S	25.0 E	24	UNDERGRO	100 - 1,000	0
PARCO 6	UTAH ALLOY ORES	GRAND	6	23 S	22.0 E	24	UNDERGRO	100 - 1,000	50
PARIS 25	UTAH ALLOY ORES	GRAND	6	23 S	22.0 E	24	SURFACE	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 33

MINES NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
PET.TREE 7-8	UNION CARBIDE CP	GRAND	27	24 S	25.0 F	24	UNDERGRO	1,000 - 100,000	100
PET.TREE-ELVA W.	UNION CARBIDE CP	GRAND	34	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	100
PETERSON PROPERTY	UNKNOWN CONTROL	GRAND			0		UNDERGRO	<100	0
PETRIFIED TREE 2	UNION CARBIDE CP	GRAND	27	24 S	25.0 F	24	UNDERGRO	1,000 - 100,000	150
PETRIFIED TREE 9	UNION CARBIDE CP	GRAND	35	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	250
PINE TREE 1	IRVINE, DON J.	GRAND	20	19 S	19.0 E	24	SURFACE	<100	0
PINTO JACK	UNION CARBIDE CP	GRAND	10	25 S	25.0 E	24	UNDERGRO	<100	0
PITTSBURG	UNION CARBIDE CP	GRAND	25	24 S	25.0 E	24	UNDERGRO	100 - 1,000	0
POLAR KING	UNION CARBIDE CP	GRAND	11	25 S	25.0 E	24	UNDERGRO	1,000 - 100,000	150
POND + SHUBERT	UNKNOWN	GRAND	16	25 S	26.0 E	24	UNDERGRO	1,000 - 100,000	100
PROSPECT 1	HETZEL, LFE	GRAND	18	25 S	25.0 E	24	SURFACE	<100	0
PROSPECT 2	HARBISON, JAMES L	GRAND	18	25 S	26.0 E	24	UNDERGRO	1,000 - 100,000	0
PROSPECT 4	BOOMERANG MINING	GRAND	18	25 S	26.0 F	24	SURFACE	<100	0
PROSPECT 7	WIENAN, S.W.	GRAND	18	25 S	26.0 E	24	SURFACE	<100	0
PROSPECT 9	MC FARLAND, BOB	GRAND	18	25 S	26.0 F	24	UNDERGRO	<100	0
PROSPECTOR	MONTGOMERY, JACK	GRAND			0		SURFACE	<100	0
QUESTA 3	BUSEY, L.E.	GRAND			0		SURFACE	<100	0
RANCH VIEW	YOSEMITE URAN., CO	GRAND	24	23 S	24.0 E	24	UNDERGRO	<100	100
RED HEAD 2	SHURE, IKE W.	GRAND	22	24 S	23.0 E	24	SURFACE	<100	0
RED OXIDE 1 + 2	CATO, CHARLES	GRAND	4	23 S	23.0 F	24	SURFACE	<100	50
RED TOP 3	CALLAHAN, JAY	GRAND			0		UNDERGRO	<100	0
PED VANADIUM GRO	UNION CARBIDE CP	GRAND	26	22 S	23.0 E	24	UNDERGRO	1,000 - 100,000	50
RIBBON RIDGE	KFELER, T.H + DALLAS	GRAND			0		SURFACE	<100	250
RIMROCK	UNION CARBIDE CP	GRAND	26	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	50
RIVER VIEW	EVANS R.C.	GRAND			0		SURFACE	<100	200
ROCKET JP. GROUP	ADAIR, IVOR	GRAND		25 S	18.0 F	24	UNDERGRO	100 - 1,000	50
ROBE	KEOGH, JOHN	GRAND	12	23 S	21.0 E	24	UNDERGRO	<100	0
RUSTY BUCKET	DETROITICS INC.	GRAND			0		SURFACE	<100	0
RUTH 1	MAY DAY URAN CO	GRAND			0		SURFACE	<100	0
S.R. FRACTION	UNION CARBIDE CP	GRAND	26	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	100
SADDLE	ZIAK, PAUL	GRAND			0		SURFACE	<100	0
SAND FLAT	MIDLAND URANIUM	GRAND	24	23 S	24.0 F	24	SURFACE	<100	0
SC. SEC. 36 (PARTIAL)	ATLAS MINERALS	GRAND	36	22 S	21.0 E	24	UNDERGRO	1,000 - 100,000	200
SECK SEC. 2	STATE OF UTAH	GRAND			0		SURFACE	<100	0
SEC. 2, 23S-21E	UNION CARBIDE CP	GRAND	2	23 S	21.0 E	24	UNDERGRO	1,000 - 100,000	200
SEC. 32, 22S-22E	HUFF, LILLIAN	GRAND	32	22 S	22.0 E	24	UNDERGRO	1,000 - 100,000	50
SHINARUMP 1	POWER OIL CO.	GRAND			0		UNDERGRO	1,000 - 100,000	50
SHINARUMP 1, 1A, 3	POWER OIL CO.	GRAND			0		UNDERGRO	1,000 - 100,000	50
SILVER MOON	UTAH ALLOY ORES	GRAND	11	23 S	21.0 E	24	UNDERGRO	<100	50
SLICK ROCK	LILY, GLEN	GRAND	28	23 S	20.0 E	24	SURFACE	<100	0
SNOW FLAKE 1	SUTHERLAND + SUTH	GRAND			0		SURFACE	<100	0
SOCKO 2	ATOMIC POWER UR	GRAND	11	47 N	20.0 W	22	UNDERGRO	100 - 1,000	50
SQUAW PARK GROUP	PENL + RUGGRI	GRAND	4	23 S	23.0 E	24	UNDERGRO	1,000 - 100,000	50
SUE + RUTH	SUNRAY MINING CO	GRAND			0		UNDERGRO	<100	0
SURE FIRE	MURPHY, OTTO	GRAND			0		SURFACE	<100	0
TEATON 2	MANGER + JONES	GRAND			0		SURFACE	<100	0
THOMAS B.	UNION CARBIDE CP	GRAND	34	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	50
THOMPSON C.	UNION CARBIDE CP	GRAND	34	24 S	25.0 E	24	UNDERGRO	1,000 - 100,000	200

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 54

MIKE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
THORN	GARDNER L.L.	GRAND	31	22 S	23.0 E	24	SURFACE	<100	0
THREE JAY 19	JONES, LEO W.	GRAND			0		SURFACE	<100	0
TOBY GROUP	DOYLE H.	GRAND			0		UNDERGRO	<100	250
TURTLE	MERIMETHUR, JAMES	GRAND			0		SURFACE	<100	0
TWIN BUTTE	LEWIS, C.L.	GRAND			0		SURFACE	<100	0
UTAH ST. LSE. 573	STATE OF UTAH	GRAND	32	22 S	25.0 E	24	UNDERGRO	<100	50
VALLEY VIEW	ROWLES + WRIGHT	GRAND	18	26 S	24.0 E	24	UNDERGRO	1,000 - 100,000	50
VIRGIN MARY 1	DAVIS R.L. + BESS	GRAND	13	23 S	21.0 E	24	SURFACE	<100	0
VIVIAN + NEW VIV	UNION CARRIAGE CO	GRAND	3	25 S	25.0 E	24	UNDERGRO	100 - 1,000	0
WASP	ABERNATHY, JESS	GRAND			0		SURFACE	<100	0
WEDD 1-10	JAHNKE, ROLAND	GRAND			0		SURFACE	<100	0
WINDY POINT	SMITH, CHARLES H	GRAND	35	22 S	22.0 E	24	SURFACE	<100	0
YELLOW GRASS POO	ABERNATHY, JESS	GRAND			0		SURFACE	<100	0
YELLOW HORSE	MC COLOUGH, W.J.	GRAND			0		UNDERGRO	100 - 1,000	0
YELLOW JACK	CATO, CHARLES	GRAND			0		UNDERGRO	<100	0
YELLOW ROAD	ABERNATHY, JESS	GRAND			0		SURFACE	<100	0
YELLOW VANADIUM	WRIGHT, WILLIAM	GRAND			0		UNDERGRO	<100	0
YIP YIP	SCHUMACHER J. I.	GRAND			0		UNDERGRO	100 - 1,000	0
DESERT VIEW	ATKINSON, W.H.	IRON			0		SURFACE	<100	50
BELL HILL	MINERALS PEFIN.	JUAB			0		UNDERGRO	100 - 1,000	150
CARNOTITE KING	GOLDEN GLOVES MC	JUAB			0		UNDERGRO	<100	50
EAGLE ROCK	CLARIDGE + WILDEN	JUAB	35	12 S	12.0 W	24	UNDERGRO	<100	50
YELLOW CHIEF	BLACK ROCK MNG.	JUAB	35	12 S	12.0 W	24	UNDERGRO	>100,000	100
LYNN GROUP	SALINA MNG. + SMC	KANE	16	40 S	9.0 W	24	UNDERGRO	100 - 1,000	50
RADIANCE	FARMUSSEN, W. + V.	KANE			0		UNDERGRO	100 - 1,000	50
YOUNG 1	BLACK MTN. URAN. C	KANE			0		SURFACE	<100	0
BUDDY	U.S. SMELT-REFNG	PIUTE	4	27 N	3.0 W	24	UNDERGRO	1,000 - 100,000	500
CLOYS MINE	BERGMILLER, P.	PIUTE	4	27 N	3.0 W	24	UNDERGRO	1,000 - 100,000	150
FAST SLGPF	MAGNOLIA LEAD+OI	PIUTE			0		SURFACE	<100	0
PAPER JOHN	BILLION MONARCH	PIUTE	4	27 S	3.0 W	24	UNDERGRO	1,000 - 100,000	500
FREEDOM GROUP	BERGMILLER, P.	PIUTE	4	27 S	3.0 W	24	UNDERGRO	>100,000	500
J D G	GLENNY P.A.	PIUTE			0		SURFACE	<100	0
LUCKYSTRIKE ANNE	WILHELM ESTATE	PIUTE	4	27 S	3.0 W	24	UNDERGRO	<100	50
POTTS FRACTION	BLACK BEAR CONSO	PIUTE	26	26 S	4.0 W	24	UNDERGRO	1,000 - 100,000	200
PROSPECTOR	SMITH, REX	PIUTE	4	27 S	3.0 W	24	UNDERGRO	1,000 - 100,000	200
VFG CLAIMS	ATHERLEY + SMITH	PIUTE			0		UNDERGRO	100 - 1,000	100
D+RCW RR PAPER Y	MECHAM, V.W.	SALT LAKE			0		MISC. - PM	<100	0
A FAULT THERE WA	HUGHES C.J.	SAN JUAN			0		SURFACE	<100	0
ABE 11	INTERNATNL ENRGY	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
ABE 8	INTERNATNL ENRGY	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
ACE	CLEON SHUNWAY	SAN JUAN	15	37 S	21.0 E	24	UNDERGRO	100 - 1,000	0
AJAX	EDDS, FRANK	SAN JUAN			0		SURFACE	<100	50
ANFTH 1	NAYAJO TRIBE	SAN JUAN			0		UNDERGRO	<100	100
ANK	ATLAS-AMAX	SAN JUAN			0		UNDERGRO	100 - 1,000	1150
ANN	ATLAS MINIPALS	SAN JUAN	11	30 S	24.0 E	24	UNDERGRO	1,000 - 100,000	500
ABH	ATLAS MINIPALS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	50
AVALANCHE 13	URADCO	SAN JUAN			0		UNDERGRO	1,000 - 100,000	350
AVALANCHE 9	URADCO	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 55

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
B + I	WHITE CANYON MNG	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
P VEE	HILL M.F.	SAN JUAN			0		UNDERGRO	<100	100
BACARDI CUTLER	ATLAS MINERALS	SAN JUAN	29 S	28,0 E	24		UNDERGRO	>100,000	350
BANGER HOLF	UNKNOWN CONTROL	SAN JUAN			0		SURFACE	<100	0
WALSEY	UNKNOWN CONTROL	SAN JUAN			0		SURFACE	<100	0
BASIN GROUP	SHUMWAY BROS. MG	SAN JUAN	3	37 S	21,0 E	24	UNDERGRO	1,000 - 100,000	100
BEE	ATOMIC RESOURCES	SAN JUAN	8	31 S	25,0 F	24	UNDERGRO	100 - 1,000	50
BELL MINE	PARLOW + SITTON	SAN JUAN	8	37 S	17,0 E	24	UNDERGRO	100 - 1,000	100
BENCH 1,2,3 + 4	MCDONALD+PATRSON	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
BETTY GROUP	ELK RIDGE MINING	SAN JUAN			0		UNDERGRO	1,000 - 100,000	250
BIG BEN 1	FLECK, SAMUEL	SAN JUAN			0		SURFACE	<100	0
BIG BLUFF	S. + R. WILKES	SAN JUAN	9	37 S	25,0 F	24	SURFACE	<100	0
BIG BOWL	PITTMAN, EMIT	SAN JUAN	31	27 S	23,0 F	24	SURFACE	<100	0
BIG RUCK	BAILEY, WALLACE	SAN JUAN			0		UNDERGRO	1,000 - 100,000	0
BIG BUCK 11	ATLAS MINERALS	SAN JUAN			0		UNDERGRO	100 - 1,000	400
BIG BUCK 7	ATLAS MINERALS	SAN JUAN	11	30 S	24,0 E	24	UNDERGRO	<100	0
BIG BUCK 7,8+8A	ATLAS MINERALS	SAN JUAN	11	30 S	24,0 E	24	UNDERGRO	1,000 - 100,000	100
BIG BUCK 8A(624K)	ATLAS MINERALS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
BIG BUCK 9,9A,10,	ATLAS MINERALS	SAN JUAN	11	30 S	24,0 F	24	UNDERGRO	>100,000	500
BIG CREEK	BAILEY, WALLACE	SAN JUAN			0		SURFACE	<100	0
BIG INDIAN 2	UTAH COLORADO OF	SAN JUAN			0		UNDERGRO	<100	150
BIG MIKE 1	LAMBERT, JAMES R	SAN JUAN			0		SURFACE	<100	0
BIG MONKEY	OPINMAN, WILLIAM	SAN JUAN			0		UNDERGRO	<100	50
BIG SADDLE 1	MISSOURI EXPL.CO	SAN JUAN			0		SURFACE	<100	0
BIG TREE	AKIN APQS.	SAN JUAN			0		SURFACE	<100	0
BILL 5	WIGHT, BILL	SAN JUAN			0		SURFACE	<100	300
BILLY JOE WHEEL	BLACK ROCK URAN.	SAN JUAN			0		SURFACE	<100	0
BIRTHDAY	FOOTE MINERALS	SAN JUAN	10	37 S	21,0 E	24	UNDERGRO	1,000 - 100,000	0
BLACK ACE 2	LYENS, W.F.	SAN JUAN			0		UNDERGRO	<100	0
BLACK BUTT	SWIPROCK, LTD	SAN JUAN	19	31 S	25,0 E	24	UNDERGRO	100 - 1,000	0
BLACK CATS 1-10	MAXWELL, MELVIN L.	SAN JUAN			0		UNDERGRO	<100	200
BLACK HAT	GEO-ENERGY RES	SAN JUAN	28	28 S	26,0 E	24	UNDERGRO	1,000 - 100,000	200
BLACK OXIDE	ABC EXPLORATION	SAN JUAN			0		UNDERGRO	100 - 1,000	200
BLACK WATER	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	1,000 - 100,000	50
BLACKSTONE DUMPS	EXCALIBUR IND.	SAN JUAN			0		DUMPS	1,000 - 100,000	100
BLO SYAKE	TIDWELL, CLAUDE E	SAN JUAN			0		SURFACE	<100	0
BLOCK 42	PLATEAU MNG, CO.	SAN JUAN			0		UNDERGRO	<100	0
BLUE BELL	MALIN, OREN	SAN JUAN			0		UNDERGRO	<100	0
BLUE BIRD	HARRISON, JAMES L	SAN JUAN			0		UNDERGRO	<100	0
BLUE BIRD 1	UNKNOWN CONTROL	SAN JUAN			0		UNDERGRO	100 - 1,000	0
BLUE BONNET	PRESLEY + TANNER	SAN JUAN			0		SURFACE	<100	0
BLUE BUTTE 1	UNKNOWN CONTROL	SAN JUAN			0		SURFACE	<100	0
BLUE CLAY	PICKENS, CHARLES	SAN JUAN			0		UNDERGRO	1,000 - 100,000	0
BLUE GOOSE	BALSLEY + WRIGHT	SAN JUAN	15	27 S	23,0 E	24	UNDERGRO	100 - 1,000	0
BLUE HILL	JOHNSON, DUANE	SAN JUAN	36	27 S	22,0 F	24	SURFACE	<100	0
BLUE JAY	BALSLEY, H.W.	SAN JUAN	28	28 S	23,0 E	24	UNDERGRO	1,000 - 100,000	0
BLUE JAY 3	RUST+BAIRD+PIECE	SAN JUAN	20	37 S	25,0 E	24	UNDERGRO	<100	0
BLUE JAY DUMPS	BIERSCHIED+SWISH	SAN JUAN			0		DUMPS	1,000 - 100,000	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 56

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
BLUE LIZZARD	ENERGY FUELS MNC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	300
BOBCAT	UNKNOWN CONTRLR	SAN JUAN	11	30 S	24.0 E	24	UNDERGRO	1,000 - 100,000	450
BOJO	BOJO URANIUM CO.	SAN JUAN			0		UNDERGRO	100 - 1,000	100
BONANZA 1 + 2	BALSLEY, H.W.	SAN JUAN	27	31 S	24.0 E	24	UNDERGRO	100 - 1,000	0
BONNIE LEE	BOJO URANIUM CO.	SAN JUAN			0		SURFACE	<100	100
BOULDER	PETTERSON, FRANK	SAN JUAN			0		UNDERGRO	<100	0
BOW	INSPIRATION LEAD	SAN JUAN			0		SURFACE	<100	50
BOULEGS	ATLAS MINERALS	SAN JUAN	33	29 S	24.0 E	24	UNDERGRO	>100,000	550
BOX CANYON	FOOTE MINERALS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	50
BRADFORD 1-5	DEVILS CANYON UR	SAN JUAN			0		UNKNOWN	100 - 1,000	0
BREEZY GROUP	STOCKS+LAMBERT	SAN JUAN			0		UNDERGRO	100 - 1,000	250
BRIGHT STAR	LEWIS, C.L.	SAN JUAN	14	36 S	24.0 E	24	UNDERGRO	<100	0
BROKEN SHOE	DANVERS, DON	SAN JUAN	33	29 S	24.0 E	24	SURFACE	<100	0
BROWNIE 1	URANIUM CENTER C	SAN JUAN			0		SURFACE	<100	200
BUB	SHUMWAY BOB	SAN JUAN			0		SURFACE	<100	0
BUCK 3	BIGHORN GOLD CO	SAN JUAN			0		UNDERGRO	100 - 1,000	250
BUCK SKIN 1 + 2	ALBERT BALLARD	SAN JUAN			0		UNDERGRO	100 - 1,000	0
BUCKEYE NO 1	WEGNER W.J.	SAN JUAN			0		SURFACE	<100	50
BUCKHORN	UTOMIC EXP. + MNC	SAN JUAN	23	37 S	24.0 E	24	SURFACE	<100	150
BUCKHORN 1-8	BUCKHORN MINING	SAN JUAN			0		SURFACE	<100	250
BUCKSKIN 1-2	JIN WINBOURN	SAN JUAN			0		UNKNOWN	<100	0
BUCKSKIN ROBBY	UNKNOWN CONTRLR	SAN JUAN			0		UNDERGRO	<100	0
BUCKSKIN STAIN	SHIPROCK, ITO	SAN JUAN			0		UNDERGRO	<100	100
BUD 1	BALDWIN, ROBERT D	SAN JUAN	29	51 N	18.0 W	22	UNDERGRO	<100	0
BUSTER	NICHOLS+SUTYAK	SAN JUAN	29	27 S	23.0 E	24	UNDERGRO	<100	0
BUTLER CANYON	UNKNOWN CONTRLR	SAN JUAN			0		SURFACE	<100	0
BUTTE 1	SHUMWAY SETH	SAN JUAN			0		UNDERGRO	100 - 1,000	0
C GROUP	AIRBORNE PRSPCTP	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
C.C.C.	DFLAY SHUMWAY	SAN JUAN			0		UNDERGRO	<100	0
CALICO	RANCHERS EXPL+D	SAN JUAN			0		SURFACE	<100	100
CAMEL (CIGARETTE)	BABBELS UPAN, ENG	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
CANARY	BADGER UPAN, CORP	SAN JUAN	18	36 S	26.0 E	24	SURFACE	<100	0
CANARY	ABERNATHY, JESS	SAN JUAN	19	27 S	24.0 E	24	UNDERGRO	<100	50
CANDY	REDD+BASILIERE ME	SAN JUAN			0		SURFACE	<100	50
CANYON 1	BARBER MINING CO	SAN JUAN			0		UNDERGRO	100 - 1,000	150
CANYON 2	COL TEX URANIUM	SAN JUAN			0		UNDERGRO	<100	100
CARL, LOOK, PAYDAY	NIELSON, FRANCIS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
CAROL B	DURAY URANIUM	SAN JUAN	3	28 S	23.0 E	24	UNDERGRO	100 - 1,000	0
CARTER 2	MORGAN, CHARLES G	SAN JUAN			0		UNDERGRO	<100	0
CASMIN 3	SHUMWAY + WESTAD	SAN JUAN			0		SURFACE	<100	0
CEDAR	ZIAK, PAUL	SAN JUAN			0		SURFACE	<100	0
CEDAR 37	YUBA DEVELOP. CO.	SAN JUAN			0		UNDERGRO	100 - 1,000	250
CEDAR BIRD	BRADFORD, SYLVEST	SAN JUAN	9	37 S	21.0 E	24	UNDERGRO	<100	150
CEDAR INCLINE	YUBA DEVELOP. CO.	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
CHAMP 1	CHAMP MINING CO	SAN JUAN			0		SURFACE	<100	0
CHARLES KEITH	NAVJO TRIBE	SAN JUAN			0		UNDERGRO	<100	150
CHARLEY 3	WHITE CANYON MNC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	250
CHESS RIDGE	SHUMWAY PETE	SAN JUAN	7	27 S	24.0 E	24	UNDERGRO	<100	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 57

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH	(CONT'D) *****								
CHIPMUNK	BROPHY + YACKEL	SAN JUAN	15	27 S	23.0 E	24	SURFACE	<100	0
CLAIM 22	DALPEZ+MORTISON	SAN JUAN			0		SURFACE	<100	100
CLAIM 4	UNKNOWN CONTROLLER	SAN JUAN			0		UNDERGRO	100 - 1,000	50
CLAIM 8	STANDARD METALS	SAN JUAN			0		UNDERGRO	<100	0
CLIFF HOUSE	UNION CARBIDE CP	SAN JUAN	14	36 S	25.0 E	24	UNDERGRO	100 - 1,000	50
CLIFFAX 2	ATLAS MINERALS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	50
COAL CREEK	BITTON + DULANEY	SAN JUAN			0		UNDERGRO	<100	0
COLUMBIA 1	ATLAS MINERALS	SAN JUAN	33	29 S	24.0 E	24	UNDERGRO	1,000 - 100,000	700
CONCENTRATE	C.O.G. MINERALS	SAN JUAN			0		LOW GRADE	1,000 - 100,000	0
CONGLOMERATE 2	INTERMOUNTAIN UR	SAN JUAN		31 S	21.0 E	24	SURFACE	<100	0
COPPER TOP 5	WRIGHT, L.A.	SAN JUAN			0		SURFACE	<100	300
COSTANZA	PIONEER DRUG CO.	SAN JUAN	35	30 S	25.0 E	24	UNDERGRO	1,000 - 100,000	100
COTTONWOOD	BALSLEY, H.W.	SAN JUAN			0		UNDERGRO	100 - 1,000	0
COTTONWOOD	UNKNOWN CONTROLLER	SAN JUAN			0		UNDERGRO	1,000 - 100,000	0
COTTONWOOD 1	DENTON F.J.	SAN JUAN			0		UNDERGRO	100 - 1,000	100
COTTONWOOD 2	BLACK, CALVIN	SAN JUAN	3	37 S	21.0 E	24	UNDERGRO	1,000 - 100,000	0
COTTONWOOD 2+3	SHUMWAY+HOLLIDAY	SAN JUAN	4	37 S	21.0 E	24	UNDERGRO	1,000 - 100,000	100
COTTONWOOD 3	BLACK, C.+JONES K	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
COTTONWOOD 4	HAWKEY BELLUM	SAN JUAN	4	37 S	21.0 E	24	UNDERGRO	1,000 - 100,000	100
COTTONWOOD TAILI	BLANDING MINES C	SAN JUAN			0		MISC.+PB	1,000 - 100,000	50
COUGAR	INDUSTRIAL URAN.	SAN JUAN			0		UNDERGRO	<100	100
COVE GROUP	C.O.G. MINERALS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	250
COYOTE 1	BURDETT SHUMWAY	SAN JUAN	35	35 S	24.0 F	24	UNDERGRO	<100	50
CUTLER	CLIFF MINING CO.	SAN JUAN			0		UNDERGRO	100 - 1,000	50
CYS MONUMENT	ENERGY FUELS NUC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
DAISY 2	JIM C. BUTT	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
DARK HORSE	DAVIS, HOWER	SAN JUAN			0		SURFACE	<100	0
DELAWARE CHIFF 1	GIBBALTAP URAN.C	SAN JUAN			0		UNDERGRO	<100	250
DEVIL 3	CROSS, ODDIE V.	SAN JUAN			0		UNDERGRO	<100	250
DEVIL CANYON 1	HURST,QUINTON R.	SAN JUAN			0		UNDERGRO	<100	0
DIABLO	CRA+FOORD+RIGHT	SAN JUAN			0		SURFACE	<100	250
DIME	SHUMWAY+ DADE	SAN JUAN	7	31 S	24.0 E	24	UNDERGRO	1,000 - 100,000	50
DIP GROUP	MINERAL HILL UR	SAN JUAN			0		UNDERGRO	100 - 1,000	50
DISMONT SLIMES	ATOMIC RESOURCES	SAN JUAN			0		UNDERGRO	<100	0
DISSIPATION	HOMESTEAD MNG CO	SAN JUAN	28	29 S	24.0 E	24	UNDERGRO	1,000 - 100,000	500
DIVIDE+YELLOW JC	RED ROCK DEV C	SAN JUAN	35	30 S	25.0 F	24	UNDERGRO	1,000 - 100,000	150
DIXIE 2	SURPLUS URANIUM	SAN JUAN	19	37 S	25.0 E	24	SURFACE	<100	150
DOLLAR	C.O.S.URANIUM CO	SAN JUAN	26	36 S	24.0 E	24	UNDERGRO	<100	250
DOROTHY 2	BARRY, ROBERT	SAN JUAN			0		SURFACE	<100	100
DOROTHY WAY	ATLAS MINERALS	SAN JUAN	13	30 S	24.0 E	24	UNDERGRO	>100,000	400
DRY HOLE(NE 35)	COTTONWOOD MNG.	SAN JUAN	4	37 S	21.0 F	24	UNDERGRO	1,000 - 100,000	100
DUGWAY	FOOTE MINERALS	SAN JUAN			0		UNDERGRO	100 - 1,000	50
DUSTY	ALONGO, E.J.	SAN JUAN			0		UNDERGRO	100 - 1,000	50
E.TAYLOR CANYON	DELHI URAN + OIL	SAN JUAN			0		UNDERGRO	100 - 1,000	50
EAGLE NEST	MOAB URANIUM CO.	SAN JUAN			0		SURFACE	<100	100
EARLY DAWN	STOCKS, H.P.	SAN JUAN			0		UNDERGRO	<100	0
EARLY HORN	STOCKS, H.P.	SAN JUAN			0		SURFACE	100 - 1,000	0
EAST BANK	SHUMWAY BROS.+NG	SAN JUAN	3	37 S	21.0 E	24	UNDERGRO	1,000 - 100,000	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 58

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
EAST RIM	BLEAK, FLOYD	SAN JUAN			0		UNDERGRO	100 - 1,000	50
EAST WOODEN SHOUL	SETH SHUMWAY	SAN JUAN			0		UNDERGRO	100 - 1,000	150
ELLA	WASHBURN, LARK	SAN JUAN	28	28 S	26,0 E	24	UNDERGRO	1,000 - 100,000	300
ELMORE 1	PRESLEY + TANNER	SAN JUAN			0		UNDERGRO	<100	0
EVA	UNION CARRIAGE	SAN JUAN	13	47 N	20,0 W	22	UNDERGRO	1,000 - 100,000	0
EXPECTATION	ATLAS MINERALS	SAN JUAN	3	30 S	24,0 E	24	UNDERGRO	>100,000	750
F.D.P. 9	NEILSON, D.J.	SAN JUAN			0		UNDERGRO	<100	250
F.I. GROUP	HILLSBOPO URAN	SAN JUAN			0		UNDERGRO	<100	100
FAIR DAY	JONES + SIMPSON	SAN JUAN			0		SURFACE	<100	0
FAIR	MERIT URAN, CORP.	SAN JUAN			0		UNDERGRO	<100	100
FAULT 1	ARBOT, DAVID	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
FIREFLY	GEO-ENERGY RES	SAN JUAN	30	28 S	26,0 E	24	UNDERGRO	1,000 - 100,000	300
FIRST+LAST CHANC	WRIGHT, L.R.	SAN JUAN			0		UNDERGRO	<100	200
FORK VIEW	RANDOLPH + DILLON	SAN JUAN			0		UNDERGRO	<100	0
FOUND	FOOTE MINERALS	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
FOUND	WHITE CANYON MNG	SAN JUAN			0		UNDERGRO	1,000 - 100,000	50
FOXY DAW	WINBOURN, JAMES	SAN JUAN			0		UNDERGRO	<100	0
FRACTION	OURAY URANIUM	SAN JUAN	2	28 S	23,0 E	24	UNDERGRO	100 - 1,000	100
FRACTION CLAIM	SHUMWAY + DAVIS	SAN JUAN			0		SURFACE	<100	100
FRANCE	ENGEL, MAX W.	SAN JUAN			0		UNDERGRO	<100	100
FRIDAY	BROWN, ALBERT J.	SAN JUAN			0		UNDERGRO	<100	0
FRUNDELL 1	UNKNOWN CONTROL	SAN JUAN	9	37 S	24,0 E	24	UNDERGRO	<100	0
FRY-E, COMMAND	TANNER + TANGREE	SAN JUAN			0		SURFACE	<100	100
G.D. 1	DEAN + FERRANDO	SAN JUAN			0		SURFACE	<100	50
GARY	SHUPE BROS	SAN JUAN			0		SURFACE	<100	0
GAYLENE 2	AMALGAMATED URAN	SAN JUAN			0		SURFACE	<100	50
GILMAN + GILMAN	SHUMWAY + DADE	SAN JUAN	6	31 S	25,0 F	24	UNDERGRO	1,000 - 100,000	50
GILPAN	FOOTE MINERALS	SAN JUAN			0		UNDERGRO	<100	50
GIVELWAY	JIM C. BUTT	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
GIZNO GROUP	BLAKE, DONALD V.	SAN JUAN			0		UNDERGRO	1,000 - 100,000	450
GLADE 11	NUGENT, CLAUDE E	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
GOLD CROWN	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	<100	150
GOOD HOPE	DAY, ABE	SAN JUAN	7	31 S	25,0 E	24	UNDERGRO	<100	0
GRAND VIEW	BLUE CHIEF URAN	SAN JUAN			0		SURFACE	<100	100
GRASS ROOT	HAZER, ORSON	SAN JUAN	27	28 S	23,0 E	24	SURFACE	<100	0
GREEN + GREEN	ELK MTR, URAN, COR	SAN JUAN			0		SURFACE	<100	250
GREEN FLY 1	LAUGHTER, GERALD	SAN JUAN	13	31 S	24,0 E	24	UNDERGRO	<100	0
GREEN ROCK	WEGNER + VEGELIK + M	SAN JUAN			0		SURFACE	<100	0
GRFY DAWN	WASHBURN, LARK	SAN JUAN	30	28 S	26,0 E	24	UNDERGRO	1,000 - 100,000	0
H. BLACKWATER 2+3	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	100 - 1,000	50
HALF + HALF	JOHNSON, DAVE	SAN JUAN			0		SURFACE	<100	0
HANGOVER	C. SHUMWAY	SAN JUAN	15	37 S	21,0 E	24	UNDERGRO	100 - 1,000	0
HAPPY JACK	BRUNKE, W.W.	SAN JUAN	13	32 S	23,0 E	24	UNDERGRO	100 - 1,000	0
HAPPY SURPRISE 1	HAPPY SURPRISE M	SAN JUAN			0		UNDERGRO	<100	150
HARD ROCK	GODDWIN, RAY	SAN JUAN			0		SURFACE	<100	0
HART DRAW	CLIFF MINING CO.	SAN JUAN			0		UNDERGRO	<100	100
HATCHET 3	TURNER + TIDWELL	SAN JUAN			0		UNDERGRO	<100	100
HATTIE	MURPHY, VICTOR	SAN JUAN			0		UNDERGRO	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 59

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
HENRY L. SAMPSON	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	<100	250
HERSEY-RADIUM #1	ATLAS MINERALS	SAN JUAN			0		UNDERGRO	>100,000	500
HESPERUS	MAYFIELD, JERRY	SAN JUAN			0		UNDERGRO	100 - 1,000	50
HIDDEN LOG	TOWN MINING CO	SAN JUAN			0		UNDERGRO	100 - 1,000	0
HIDDEN VALLEY 5	M. + M. DEVELOPMENTS	SAN JUAN			0		SURFACE	<100	100
HIDEDUT MINE	UTAH MIN RESOU	SAN JUAN			0		UNDERGRO	>100,000	200
HIGH HOPES	UNKNOWN CONTROL	SAN JUAN			0		UNDERGRO	<100	0
HIGHBALL	INDUSTRIAL URAN.	SAN JUAN			0		UNDERGRO	100 - 1,000	200
HIGHWAY	DAY, ABE	SAN JUAN			0		SURFACE	<100	0
HILLTOP	STADLER, ROBERT I.	SAN JUAN			0		SURFACE	<100	0
HONEYMOON	DOUGLAS + SEARS	SAN JUAN			0		UNDERGRO	100 - 1,000	0
HOOT	SHUMWAY BROS. MC	SAN JUAN	15	37 S	21.0 E	24	UNDERGRO	100 - 1,000	100
HOOT OWL	CROSS, ODDIE V.	SAN JUAN	3	37 S	24.0 E	24	UNDERGRO	<100	200
HOPE 1	CAMPBELL, W.B.	SAN JUAN			0		UNDERGRO	100 - 1,000	0
HOPE-SAGE, ETC.	BUTT, H.D. ET AL	SAN JUAN			0		UNDERGRO	<100	150
HOPPERS	PEARSON, T.E.	SAN JUAN			0		SURFACE	<100	0
HORSEFLY	UNKNOWN CONTROL	SAN JUAN			0		SURFACE	<100	100
HORSESHOE 1	KNIGHT, M.R.	SAN JUAN	7	32 S	21.0 E	24	UNDERGRO	1,000 - 100,000	0
HOUSE	UNION CARRIAGE CO	SAN JUAN			0		UNDERGRO	100 - 1,000	0
I DON T KNO	BRUCE + CUNNINGHAM	SAN JUAN			0		SURFACE	<100	0
IDA	GRANLICH EXPLOR.	SAN JUAN	25	28 S	25.0 E	24	UNDERGRO	<100	150
JOE-NIXON SHAFT	ATLAS MINERALS	SAN JUAN	34	29 S	24.0 E	24	UNDERGRO	>100,000	500
INDEPENDENT	TUSING + DEAN	SAN JUAN			0		SURFACE	<100	150
INDIAN CREEK	PARADOX MINING CO	SAN JUAN			0		UNDERGRO	<100	50
INDIAN CREEK GRP	PARADOX MINING CO	SAN JUAN	10	33 S	22.0 E	24	UNDERGRO	1,000 - 100,000	0
J.R.P. 16	ALADDIN EXPL. INC	SAN JUAN			0		SURFACE	<100	150
JACK	GLENNY CUTLER	SAN JUAN			0		UNDERGRO	<100	50
JACKPOT	LONG STAR MNG. CO	SAN JUAN			0		UNDERGRO	<100	50
JACKPOT	BENTLEY + GARCIA	SAN JUAN			0		UNDERGRO	100 - 1,000	0
JACKS	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	100 - 1,000	300
JACORS CHAIR	ATOMIC EXTRACT, N	SAN JUAN			0		UNDERGRO	<100	100
JIM DANDY	ROGERS, DARRYL	SAN JUAN			0		SURFACE	<100	0
JOAN	WATKINS, LYMAN M.	SAN JUAN			0		SURFACE	<100	0
JOAN	WASHBURN, LARK	SAN JUAN	28	28 S	26.0 E	24	UNDERGRO	1,000 - 100,000	0
JOE MISHOP	REDD, ELLIOTT	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
JOHN CLAIMS	SHUMWAY M + B	SAN JUAN			0		UNDERGRO	100 - 1,000	150
JOHN D 2	CLOUSE + FOSTER	SAN JUAN			0		SURFACE	<100	100
JOHNIE	SHUMWAY BROS. MC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	250
JOHNNIE MINE	VIGIL BROS	SAN JUAN	11	32 S	23.0 E	24	UNDERGRO	<100	0
JOKER	COTTONWOOD MNG.	SAN JUAN			0		UNDERGRO	100 - 1,000	0
JOKER 1	MINERALS WEST	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
JOHAC 1, 2 + 3	CENTURY 21 MNG	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
JUDY KAY	CRAIG, EARL M.	SAN JUAN			0		SURFACE	<100	0
JUNCTION GROUP	APPLEBEE, GLEN	SAN JUAN	3	29 S	23.0 E	24	UNDERGRO	1,000 - 100,000	0
KIEV	AKIN, M.E.	SAN JUAN	10	37 S	21.0 E	24	SURFACE	<100	0
KING 3	SESAME MINING CO	SAN JUAN			0		UNDERGRO	<100	0
KING JAMES VIRGE	URADCO	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
KITTY	SHUMWAY KENNETH	SAN JUAN			0		SURFACE	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 60

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
KNOB	REDD, PRESTON	SAN JUAN			0		UNDERGRO	100 - 1,000	150
LA LOMA 2	BAILEY, R.D.	SAN JUAN			0		UNDERGRO	100 - 1,000	100
LAST CHANCE 1	ATLAS URANIUM CO	SAN JUAN	17	28 S	23.0 E	24	UNDERGRO	100 - 1,000	150
LAST CHANCE 1	ALCO URANIUM	SAN JUAN	31	35 S	25.0 E	24	UNDERGRO	1,000 - 100,000	150
LAST CHANCE 3	SHUPE, WADE	SAN JUAN			0		UNDERGRO	<100	150
LAVENDER 5	FROST, ALFRED	SAN JUAN			0		SURFACE	<100	0
LEASE 3226	STATE OF UTAH	SAN JUAN			0		UNDERGRO	<100	100
LEASE 3919	STATE OF UTAH	SAN JUAN			0		SURFACE	<100	0
LEDGE	ELLISON, W.	SAN JUAN	10	37 S	21.0 F	24	UNDERGRO	1,000 - 100,000	0
LENA 16	TURNER BROS	SAN JUAN			0		UNDERGRO	100 - 1,000	0
LENA 20	TURNER BROS	SAN JUAN			0		UNDERGRO	<100	0
LENA 4	KELLEY, JANE S.	SAN JUAN			0		UNDERGRO	<100	0
LEO J.	J. + J. URANIUM	SAN JUAN			0		UNDERGRO	<100	150
LINERTY	ERNEST, HAROLD	SAN JUAN	29	31 S	26.0 E	24	UNDERGRO	100 - 1,000	100
LINDA MUJAR	ATLAS MINERALS	SAN JUAN	11	30 S	24.0 E	24	UNDERGRO	1,000 - 100,000	50
LITTLE DAWN	STOCKS, DONALD	SAN JUAN			0		UNDERGRO	<100	0
LITTLE DEVIL	BOWLES + HEFLIN	SAN JUAN	3	28 S	23.0 E	24	UNDERGRO	<100	0
LITTLE DIAM 3	LONE BUTTE MNG.	SAN JUAN			0		UNDERGRO	100 - 1,000	50
LITTLE DOROTHY	UNKNOWN CONTROLP	SAN JUAN			0		SURFACE	<100	0
LITTLE FAIR	NICHOLS, CARL L.	SAN JUAN	15	27 S	23.0 E	24	UNDERGRO	<100	0
LITTLE INDIAN	SITTON F A	SAN JUAN			0		SURFACE	<100	0
LITTLE JACK	QUIGLEY, FRANK	SAN JUAN			0		UNDERGRO	<100	100
LITTLE PETER	WASHAURN, LARK	SAN JUAN			0		UNDERGRO	100 - 1,000	150
LIZARD	CLEVELAND, J.G.	SAN JUAN	16	28 S	23.0 E	24	SURFACE	<100	0
LOGAN	UNKNOWN CONTROLP	SAN JUAN	7	31 S	25.0 E	24	SURFACE	<100	0
LONE BUTTE	MOONAN RALPH	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
LONE PINE	DYER, DELBERT	SAN JUAN	6	32 S	24.0 E	24	SURFACE	<100	0
LONE STAR	BAILEY + FILSON	SAN JUAN	7	31 S	26.0 E	24	UNDERGRO	100 - 1,000	0
LONE WOLF	HARD, W.D.	SAN JUAN			0		SURFACE	<100	0
LONSONE 5 + 6	STOCKS, CLAYTON	SAN JUAN			0		UNDERGRO	100 - 1,000	150
LONG HOLE	RUST, GLEN	SAN JUAN			0		UNDERGRO	<100	0
LOST BOY + LOST RD	BARBER MINING CO	SAN JUAN			0		UNDERGRO	1,000 - 100,000	250
LOST INDIAN 1	DOHILL MNG. CO.	SAN JUAN			0		SURFACE	<100	0
LOUISA	ATLAS MINERALS	SAN JUAN	14	30 S	24.0 E	24	UNDERGRO	>100,000	550
LOYA RAY	COLORADO DRUG. +	SAN JUAN	6	32 S	24.0 F	24	UNDERGRO	100 - 1,000	50
LUCKY BOY 1	CENTURY 21 MNG.	SAN JUAN	34	35 S	24.0 E	24	UNDERGRO	100 - 1,000	300
LUCKY DAY 2	ARMSTRONG GERMAN	SAN JUAN			0		SURFACE	<100	0
LUCKY LADY	CHITONWOOD MNG.	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
LUCKY STRIKE	SHUMWAY RAY + PE	SAN JUAN			0		UNDERGRO	1,000 - 100,000	0
LUCKY STRIKE	GOFORTH, GAY	SAN JUAN			0		UNDERGRO	100 - 1,000	100
LUCKY STRIKE 1 +	CABOT CORP.	SAN JUAN			0		UNDERGRO	<100	0
LUCKY STRIKE 2	CASTER, LESTER	SAN JUAN			0		SURFACE	<100	150
LULU 1	URANIUM CORPORAT	SAN JUAN			0		UNDERGRO	<100	100
MABLE DEE	LISBON URAN. COR	SAN JUAN			0		UNDERGRO	100 - 1,000	450
MACHINE REJECTS	MARCY SHENANDOAH	SAN JUAN			0		MISC. - PB	<100	0
MAMIE	ATLAS MINERALS	SAN JUAN	27	29 S	24.0 E	24	UNDERGRO	>100,000	400
MARCY GROUP	ENERGY FUELS NUC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
MARTON	UNKNOWN	SAN JUAN			0		UNDERGRO	100 - 1,000	150

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 61

MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****		
MARY JANE 7	SMITH, RICHARD F.	SAN JUAN
MAXINE	MARSHON, JAMES L.	SAN JUAN
MAXINE 2	TRANS WESTERN UR	SAN JUAN
MAXINE 3	TRANS WESTERN UR	SAN JUAN
MAYBE MINE	NEILSON, MILTON	SAN JUAN
MESA	GOOD EARTH MNG.	SAN JUAN
MEXICAN HAT STOC	NAVAJO TRIBE	SAN JUAN
MI CORAZON	ATLAS MINERALS	SAN JUAN
MIDVALE	SHUMWAY + DADE	SAN JUAN
MILL	R + J MINING	SAN JUAN
MINERAL 10	SHUMWAY + WESTHO	SAN JUAN
MIRADOR-HORNY TO	ELAM J.D.	SAN JUAN
MITTEN 1	NAVAJO TRIBE	SAN JUAN
MOE 1	BLAKE, DONALD V.	SAN JUAN
MOKI 1	MOKI MNG CO	SAN JUAN
MOLLIE, UT. ST. 56	STATE OF UTAH	SAN JUAN
MONTEZUMA	BLACK THUNDER OI	SAN JUAN
MONTEZUMA 1	BLACK THUNDER OI	SAN JUAN
MONUMENT 3	BUTT, H.D.	SAN JUAN
MOONLIGHT 1,2+3	SECURITY URA+OIL	SAN JUAN
NAP	TERRELL + BLACK	SAN JUAN
NEW AL POGERS	M.T. + G. MNG. CO.	SAN JUAN
NIPPLES	TORRES, DAVID	SAN JUAN
NOSE SUCK-RIG LE	FOOTE MINERALS	SAN JUAN
NORTH MESA	NICHOL-ADAIR+BP	SAN JUAN
NORTH POINT 6	WHITE CANYON MNG	SAN JUAN
NORTH PT.-GONEA-	YUBA DEVELOP.CO.	SAN JUAN
OKIE	A+B MINING CO	SAN JUAN
OLD POWDER	HELMAN, KATT+DA	SAN JUAN
PAGODA	EAST BASIN OIL +	SAN JUAN
PANDIA	HURST, BOR	SAN JUAN
PASCO, JEN, JACKIE	ATLAS MINERALS	SAN JUAN
PAY DAY	QUIGLEY, FRANK	SAN JUAN
PAY OFF	WESTWOOD + SMITH	SAN JUAN
PAYDAY-CLOUDY DA	ENERGY FUELS MUC	SAN JUAN
PEACH A 1	RICHEY, LEE	SAN JUAN
PEARL	ATLAS MINERALS	SAN JUAN
PEAVINE QUEEN	KIMMERLE, HOWARD	SAN JUAN
PEGGY	SHUPE, WADE	SAN JUAN
PETE GPOUP	CANYONLANDS URAV	SAN JUAN
PETERING	SHUMWAY PETP	SAN JUAN
PHILLIP DEE 1	NAVAJO TRIBE	SAN JUAN
PICKALO PETE	BLEAK URANIUM CO	SAN JUAN
PIGMY	STERN + ZASTROW	SAN JUAN
PINCH	UNKNOWN CONTROLR	SAN JUAN
PINE TREE	PETE SHUMWAY	SAN JUAN
PIUTE	CONSOLIDATED URA	SAN JUAN
POINT	FOOTE MINERALS	SAN JUAN

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 62

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
POINT	YUBA DEVELOP.CO.	SAN JUAN		36 S	16,0 E	24	SURFACE	<100	50
POPEYE	JARAMILLO, THOS.	SAN JUAN	4	32 S	23,0 E	24	SURFACE	<100	0
PORCUPINE	HARBISON, JAMES L	SAN JUAN	7	28 S	23,0 E	24	UNDERGRO	<100	0
PORCUPINE 1	CHATU URAN.MNG.	SAN JUAN			0		SURFACE	<100	200
PRICE-LEUC	SLAGTER EXPLORAT	SAN JUAN			0		UNDERGRO	<100	100
PRINCE ALBERT	SILVER EAGLE MNG	SAN JUAN	16	28 S	23,0 E	24	UNDERGRO	100 - 1,000	0
PRODUCTION	SKYLAND DEVEL.CO	SAN JUAN			0		UNDERGRO	100 - 1,000	0
PROFIT GROUP	ATLAS-AMAX	SAN JUAN	30	31 S	25,0 E	24	UNDERGRO	1,000 - 100,000	150
PURE LUCK	EDGAR EXPLORATIO	SAN JUAN	35	36 S	25,0 E	24	UNDERGRO	100 - 1,000	150
R.H.H.	DECURSE, JACK	SAN JUAN			0		SURFACE	<100	0
R.O.METALS	APEX EXPLORATION	SAN JUAN			0		UNDERGRO	<100	100
RADON HOT ROCK C	ATLAS MINERALS	SAN JUAN	29	29 S	24,0 E	24	UNDERGRO	>100,000	700
RAINBOW	SMITH, W.I.	SAN JUAN			0		UNDERGRO	100 - 1,000	0
RAM 1	CIMOTA EXPLORATI	SAN JUAN			0		UNDERGRO	100 - 1,000	0
RAM GROUP	LYLE PROS	SAN JUAN			0		SURFACE	<100	0
RAM JET	WESTERN MINERALS	SAN JUAN			0		UNDERGRO	<100	500
RATTLESNAKE 4 U	ATLAS MINERALS	SAN JUAN			0		UNDERGRO	100 - 1,000	150
RATTLESNAKE PIT	WOODMONT INC	SAN JUAN	12	29 S	23,0 E	24	UNDERGRO	>100,000	200
RAVEN 1	RADIO GEOPHYSICA	SAN JUAN			0		SURFACE	<100	50
REFEL	R J WOODMAN	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
RECLAIMED ORE	BLACK, CECIL	SAN JUAN			0		MISC.-PB	<100	0
RED BOX + RED MO	ERNST + DILLON	SAN JUAN			0		UNDERGRO	<100	250
RED CANYON 1	PED CANYON MINER.	SAN JUAN			0		UNDERGRO	<100	100
PED DEVIL 1 + 2	OURHAM, JASPER H.	SAN JUAN			0		SURFACE	<100	0
PED HOT 1-36	ALTURA MINING CO	SAN JUAN			0		UNDERGRO	100 - 1,000	50
RED HOTEL	UNKNOWN CONTROLR	SAN JUAN			0		SURFACE	<100	0
RED PATCH	SHUMWAY, DADY	SAN JUAN	9	31 S	25,0 E	24	UNDERGRO	<100	50
RED ROCK	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	<100	150
PED ROCK GROUP	DENTON F.J.	SAN JUAN			0		UNDERGRO	1,000 - 100,000	300
REDLANDS GROUP	MUPPHY, VICTOR	SAN JUAN			0		SURFACE	<100	0
RENEGADE	PIETERSON, FRANK	SAN JUAN	14	27 S	23,0 E	24	UNDERGRO	100 - 1,000	0
RENEGADE 1	MINERAL HILL UPA	SAN JUAN	14	27 S	23,0 E	24	UNDERGRO	<100	0
REPRISE	ATLAS MINERALS	SAN JUAN	3	30 S	24,0 E	24	UNDERGRO	1,000 - 100,000	900
REX GROUP (DEAR F	GRANT SHUMWAY	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
REYNOLDS 1	REYNOLDS, WOPACE	SAN JUAN			0		SURFACE	<100	100
RICHARDSON	HOMESTAKE MNG CO	SAN JUAN	28	29 S	24,0 E	24	UNDERGRO	>100,000	500
RIDGE 1	STIMPSON, WOODROW	SAN JUAN	15	37 S	21,0 E	24	UNDERGRO	1,000 - 100,000	0
RIM ROCK	KASSEL URAN, ENTE	SAN JUAN			0		UNDERGRO	<100	200
ROAN CREEK	CONSOLIDATED URA	SAN JUAN			0		UNDERGRO	<100	0
ROBIN	TURNER, FRED	SAN JUAN	36	27 S	22,0 E	24	SURFACE	<100	0
ROCK 1 + 2	BURBRIIDGE, DEL	SAN JUAN	34	35 S	24,0 E	24	UNDERGRO	1,000 - 100,000	0
ROCK DOOR 1	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	<100	50
ROTTEN LOG	UNKNOWN CONTROLR	SAN JUAN			0		SURFACE	<100	0
ROYAL FLUSH	SHUMWAY PROS.MG	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
PURY	SHUPE, WADE	SAN JUAN			0		SURFACE	<100	100
RUSTY 4	TURNER URAN CORP	SAN JUAN			0		SURFACE	<100	0
RUSTY CAN 5	WRIGHT, L.B.	SAN JUAN	26	36 S	25,0 E	24	UNDERGRO	<100	250
RUSTY-LITTLE MAE	GENERAL ELECTRIC	SAN JUAN			0		UNDERGRO	1,000 - 100,000	250

INACTIVE URANIUM MINES IN THE UNITED STATES
-SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 63

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
SADDLE	WHITE CANYON MNG	SAN JUAN			0		UNDERGRO	100 - 1,000	150
SADDLE-SCENIC	YUBA DEVELOP.CO.	SAN JUAN			0		UNDERGRO	100 - 1,000	100
SALT CEDAR	BARBER MINING CO	SAN JUAN	34	29 S	20.0 E	24	UNDERGRO	100 - 1,000	0
SALT CREEK 2	JONES BROS.MNG.	SAN JUAN			0		SURFACE	<100	100
SAN JUAN 4	LINDA GAY MNG.	SAN JUAN			0		SURFACE	<100	100
SAN JUAN SHAFT	RANCHERS EXPL+D	SAN JUAN	34	29 S	24.0 F	24	UNDERGRO	1,000 - 100,000	250
SANDY	KNAPP URAN.DEVEL	SAN JUAN	33	29 S	21.0 E	24	UNDERGRO	<100	50
SC,SEC.16,275-23	STATE OF UTAH	SAN JUAN	16	27 S	23.0 E	24	UNDERGRO	<100	0
SCENIC	JIM BUTT	SAN JUAN			0		UNDERGRO	1,000 - 100,000	200
SCH SEC 2	STATE OF UTAH	SAN JUAN			0		SURFACE	100 - 1,000	50
SCH SEC. 36	STATE OF UTAH	SAN JUAN			0		SURFACE	<100	0
SCH SEC 36 A	STATE OF UTAH	SAN JUAN			0		UNDERGRO	<100	0
SCH. SEC. 16	LAMBERT +PALSLEY	SAN JUAN			0		SURFACE	<100	0
SCH. SEC. 32	STATE OF UTAH	SAN JUAN			0		SURFACE	<100	0
SCH. SEC. 32	STATE OF UTAH	SAN JUAN	32	37 S	23.0 E	24	SURFACE	100 - 1,000	0
SCH,SEC.16-285-2	BALDLEY, TOM	SAN JUAN	16	28 S	23.0 E	24	UNDERGRO	100 - 1,000	100
SCH,SEC.16-395-2	JIM BUTT	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
SCH,SEC.36,305-2	TRIDENT MIN CO	SAN JUAN	36	30 S	25.0 E	24	UNDERGRO	1,000 - 100,000	150
SCH,SEC.36,435-1	STATE OF UTAH	SAN JUAN	36	43 S	14.0 E	24	UNDERGRO	1,000 - 100,000	100
SCHMITZ FOLLY	BADGER URAN,CORP	SAN JUAN			0		SURFACE	<100	50
SCHOOL SECTION	AMERICAN URAN EN	SAN JUAN			0		UNDERGRO	<100	150
SEBASTPOOL	FERGUSON AV.+ CT	SAN JUAN			0		UNDERGRO	100 - 1,000	0
SEC.36,245-24E U	UNION CARBIDE	SAN JUAN	36	28 S	24.0 E	24	UNKNOWN	1,000 - 100,000	450
SEC.36,315-24E	HOPKINS, ED	SAN JUAN	36	21 S	24.0 F	24	UNDERGRO	1,000 - 100,000	150
SELDON	DELAH SHUMWAY	SAN JUAN			0		UNDERGRO	100 - 1,000	0
SERVICE	LYEN,POLLOCK+KUM	SAN JUAN			0		SURFACE	<100	0
SERVICE BERRY	LYLE FRANCIS	SAN JUAN	34	30 S	25.0 E	24	UNDERGRO	100 - 1,000	0
SHALE	SHUMWAY BROS.MG	SAN JUAN	3	37 S	21.0 E	24	UNDERGRO	1,000 - 100,000	100
SHALE BLOCK 1 +	FOOTE MINERALS	SAN JUAN			0		UNDERGRO	100 - 1,000	100
SHINE	SHUMWAY BROS.MG	SAN JUAN			0		UNDERGRO	100 - 1,000	0
SHIKLEY 1	CRAIG MINING CO,	SAN JUAN			0		UNDERGRO	<100	0
SIMPATICA 3	SHUMWAY BROS.MG	SAN JUAN			0		UNDERGRO	1,000 - 100,000	0
SIR SNAPPER	NOLAND, FRANK	SAN JUAN			0		UNDERGRO	<100	250
SKEETER	STOCKS, JOHN	SAN JUAN			0		UNDERGRO	<100	0
SKIP	BISHOP + DAVIS	SAN JUAN			0		SURFACE	<100	0
SKUNKOVICH 1	ATLAS MINERALS	SAN JUAN	11	30 S	24.0 E	24	UNDERGRO	1,000 - 100,000	550
SKUNKOVICH 2	ATLAS MINERALS	SAN JUAN	14	30 S	24.0 E	24	UNDERGRO	1,000 - 100,000	650
SKYLINE	NAVAJO TRIBE	SAN JUAN	27	43 S	15.0 E	24	UNDERGRO	1,000 - 100,000	200
SLIDE	SHUMWAY H + L	SAN JUAN			0		UNDERGRO	<100	0
SLIDE-BIG THREE	CHESS RIDGE MNG	SAN JUAN	7	25 S	24.0 F	24	UNDERGRO	<100	100
SLUM	MONTEZUMA URAN,	SAN JUAN	28	35 S	24.0 E	24	UNDERGRO	<100	150
SMALL FRY 5 + 7	RANCHERS EXPL+D	SAN JUAN	35	29 S	24.0 E	24	UNDERGRO	1,000 - 100,000	200
SOLOMAN EXHA	TATMAN+GILLESPIE	SAN JUAN			0		SURFACE	<100	500
SOUTH ALICE	HOMESTAKE MNG CO	SAN JUAN	33	29 S	24.0 E	24	UNDERGRO	1,000 - 100,000	700
SOUTH ALMAR GROU	ATLAS MINERALS	SAN JUAN	28	29 S	24.0 E	24	UNDERGRO	>100,000	750
SOUTH NOTCH	BERNSTEIN+CARTER	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
SOUTH SLOPE	BENTLEY, JIM	SAN JUAN	13	31 S	24.0 E	24	SURFACE	<100	0
SOUTH WIND 1	DAKOTA MINING CO	SAN JUAN			0		UNDERGRO	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 64

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
BPOOK-DEE	CBC MINING CO.	SAN JUAN			0		UNDERGRO	1,000 = 100,000	300
SPRING CREEK	FOOTE MINERALS	SAN JUAN	3	37 S	21,0 E	24	UNDERGRO	1,000 = 100,000	50
SPRING WATER GRO	SHUMWAY BROS. MC	SAN JUAN	3	37 S	21,0 E	24	UNDERGRO	1,000 = 100,000	150
SSL 2470	STATE OF UTAH	SAN JUAN			0		SURFACE	<100	0
ST. LSE, 5691 UTA	STATE OF UTAH	SAN JUAN			0		UNDERGRO	<100	150
STARLIGHT	SHUPE, IKE W.	SAN JUAN			0		SURFACE	<100	0
STATE LINE	NEILSON, MILTON	SAN JUAN	26	31 S	24,0 E	24	UNDERGRO	<100	0
STINKO	ATLAS MINERALS	SAN JUAN	3	30 S	24,0 E	24	UNDERGRO	>100,000	650
STONE LOG 1	BASHAW, FRED L.	SAN JUAN			0		UNDERGRO	<100	50
SUNDAY 4	NATURAL POWERS C	SAN JUAN			0		SURFACE	<100	50
SUNDOON	STEWART, WESLEY	SAN JUAN	14	27 S	23,0 E	24	UNDERGRO	<100	0
SUNDOON	NAVAJO TRIBE	SAN JUAN			0		SURFACE	<100	200
SUNNY DAY	BLAKE, DONALD V.	SAN JUAN	6	24 S	24,0 E	24	SURFACE	<100	0
SUNNYSIDE	BALSLEY, H.W.	SAN JUAN	33	28 S	23,0 E	24	UNDERGRO	100 = 1,000	0
SUNNYSIDE	WOODMONT INC.	SAN JUAN	11	31 S	24,0 E	24	UNDERGRO	1,000 = 100,000	300
SUNNISE	ATLAS MINERALS	SAN JUAN			0		UNDERGRO	1,000 = 100,000	50
SUNRISE	STEWART, WESLEY	SAN JUAN	14	27 S	23,0 E	24	SURFACE	<100	0
SUNSET	NICK J. MIPPHY	SAN JUAN	30	31 S	25,0 E	24	UNDERGRO	1,000 = 100,000	50
SUNSHINE	MAXWELL, MELVIN I.	SAN JUAN	26	35 S	24,0 E	24	UNDERGRO	1,000 = 100,000	0
SURPRISE	SAYFO, CHARLIE	SAN JUAN			0		SURFACE	<100	0
TAYLOR PEID 1+2	NAVAJO TRIBE	SAN JUAN	15	43 S	14,0 E	24	UNDERGRO	1,000 = 100,000	200
TE QUIERO	LARRY SHUMWAY	SAN JUAN			0		UNDERGRO	1,000 = 100,000	0
TERGIRA	TERGIRA INC	SAN JUAN			0		SURFACE	<100	50
TEXTAR	MOONAN R J	SAN JUAN			0		UNDERGRO	<100	250
TOP	JOHNSON, DAVE	SAN JUAN	31	27 S	23,0 E	24	SURFACE	<100	0
TREASURE TROVE	ASIMUS, C.C.	SAN JUAN			0		SURFACE	<100	0
TREE	BLFAK, CHAPLES	SAN JUAN			0		UNDERGRO	100 = 1,000	100
UNIT 7	LEWIS, W.E	SAN JUAN			0		SURFACE	<100	150
UNKNOWN	WOKI URAN, SYNDIC	SAN JUAN			0		UNDERGRO	<100	50
URACOP 1+3	DENTON F.J.	SAN JUAN			0		UNDERGRO	1,000 = 100,000	150
URANIUM KING	SUTHERLAND M C	SAN JUAN	6	37 S	16,0 E	24	UNDERGRO	1,000 = 100,000	300
UTAH ST. LSE 2291	STATE OF UTAH	SAN JUAN			0		SURFACE	<100	100
UTAH ST. LSE, 233	SHUMWAY, CADE	SAN JUAN			0		SURFACE	<100	150
UTAH ST. LSE, 510	STATE OF UTAH	SAN JUAN			0		UNDERGRO	<100	0
UTAH ST. LSE, 1760	ANDERSON BRUS, DR	SAN JUAN			0		UNDERGRO	1,000 = 100,000	150
UTAH ST. LSE, 2333	COSTANZA, JOSEPH	SAN JUAN			0		UNDERGRO	100 = 1,000	150
VAL VISTA 1	CONFEDERATED MET	SAN JUAN			0		UNDERGRO	<100	150
VALLEY POINT	THOMPSON, NOEL	SAN JUAN			0		SURFACE	<100	0
VALLEY VIEW	ATOMIC RESOURCES	SAN JUAN	11	31 S	24,0 E	24	UNDERGRO	<100	0
VANADIUM QUEEN	BRUNKE, H.W.	SAN JUAN			0		UNDERGRO	100 = 1,000	0
VARIOUS	W.C.T. ENGINEERING	SAN JUAN			0		UNDERGRO	1,000 = 100,000	0
VARIOUS	URANIUM PROCESSE	SAN JUAN			0		SURFACE	<100	0
VEDURE	PARAMOUNT URAN, C	SAN JUAN	34	35 S	25,0 E	24	UNDERGRO	100 = 1,000	250
VELDERS 1	VELDERS, JOHN	SAN JUAN			0		UNDERGRO	<100	0
VELVET	FOOTE MINERALS	SAN JUAN	19	31 S	25,0 E	24	UNDERGRO	1,000 = 100,000	0
VICTOR	MURPHY, OTTO	SAN JUAN			0		SURFACE	<100	0
VIEW	FOOTE MINERALS	SAN JUAN			0		UNDERGRO	<100	50
VIEW NORTH 1	SOUTHERN CROSS U	SAN JUAN			0		UNDERGRO	<100	200

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 65

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UTAH (CONT'D) *****									
VISION	PLATEAU MNG.CO.	SAN JUAN			0		UNDERGRO	100 - 1,000	100
M.J.1	CASTOR & STEWART	SAN JUAN			0		UNDERGRO	100 - 1,000	200
M.N.	YUBA DEVELOP.CO.	SAN JUAN	21	3 S	1,7 E	24	UNDERGRO	1,000 - 100,000	150
WARAN MINE	RALPH J MOONAN	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
WATERLOO	MALEY MINING CO.	SAN JUAN	16	37 S	21,0 E	24	SURFACE	<100	0
WATERLOO-STEPLAD	DUNCAN WALTER JR	SAN JUAN		5	0 F	24	UNDERGRO	<100	250
WEST DAY DAY	DANIEL R.F.	SAN JUAN	36	24 S	19,0 E	24	UNDERGRO	100 - 1,000	250
WHEFLER	OSULLIVAN-THIRKE	SAN JUAN			0		SURFACE	<100	250
WHIRLWIND	NAVAJO TRIBE	SAN JUAN			0		UNDERGRO	1,000 - 100,000	150
WHITE CANYON STR	FOOTE MINERALS	SAN JUAN			0		UNDERGRO	<100	0
WHITE HORSE GROU	BARRY, ROBERT A.	SAN JUAN	11	36 S	25,0 E	24	UNDERGRO	<100	0
WILLIE	UNKNOWN CONTROL	SAN JUAN			0		SURFACE	<100	0
WIND BLOWN	SAN MIGUEL MINES	SAN JUAN			0		SURFACE	<100	0
WINDFALL 27	ATLAS-FOOTE	SAN JUAN			0		UNDERGRO	1,000 - 100,000	100
WOLTER	LYMAN, BOB	SAN JUAN			0		SURFACE	<100	0
WOODROW WILSON	NEILSON, MILTON	SAN JUAN	35	30 S	24,0 E	24	UNDERGRO	1,000 - 100,000	0
WYOMING	NEILSON, MILTON	SAN JUAN			0		UNDERGRO	100 - 1,000	0
YELLOW BARY	MERZ & KNOX	SAN JUAN			0		UNDERGRO	<100	0
YELLOW GINL	KNOX, HARRY	SAN JUAN	13	44 N	20,0 W	22	SURFACE	<100	0
YELLOW JOHN	JIM C. BUTT	SAN JUAN			0		UNDERGRO	1,000 - 100,000	50
YELLOW PAUL 1	F. C. WESTERN MG	SAN JUAN			0		UNDERGRO	100 - 1,000	0
ZIG ZAG	HAUGUM, CARL	SAN JUAN			0		UNDERGRO	<100	0
R.B.R.	APPIER, DAVID	SEVIER			0		SURFACE	<100	50
FLAT TIRE 2	MARYSVILLE URAN.C	SEVIER	24	26 S	4,0 W	24	UNDERGRO	<100	100
BOBO	TEXAS PLUNGERS-A	UINTAH			0		UNDERGRO	<100	0
BRENDA GROUP	PERMAN CO.	UINTAH			0		SURFACE	<100	0
BUZZARD 1	HAMILTON ASSOC.	UINTAH			0		SURFACE	<100	0
CROW 1	HAMILTON, STEVE	UINTAH			0		SURFACE	<100	50
DEVILS CAVE	MARTINSEN, M. LYNN	UINTAH			0		UNDERGRO	<100	50
EUREKA-CANARY A	COPPER & SANDS	UINTAH			0		UNDERGRO	<100	0
GREEN ROCK	SMOYTS, DARRELL	UINTAH			0		SURFACE	<100	50
HORSE EAR	MOTT, ORSON	UINTAH			0		SURFACE	<100	0
LION 5	MORGETTS, SUMNER	UINTAH			0		SURFACE	<100	0
LONE PEAK	ERRISA L.M.	UINTAH			0		SURFACE	<100	0
RATTLESNAKE	HARRISON J.W.	UINTAH			0		SURFACE	<100	0
SANDY	ANSGY, DELMER C.	UINTAH			0		SURFACE	<100	0
SATIN GROUP	COLOTAN URANIUM	UINTAH			0		SURFACE	<100	100
STEPHENS	FLUXEY & HUNT	UINTAH			0		SURFACE	<100	0
ANNAS PRIDE	LA FLOPECITA MNG	WASHINGTON			0		SURFACE	<100	0
KOLOB	STATE OF UTAH	WASHINGTON			0		UNDERGRO	100 - 1,000	50
LOCALORE 1	LA VORFIN MNG.	WASHINGTON			0		SURFACE	<100	0
SILVER POINT LDD	LA FLOPECITA MNG	WASHINGTON			0		SURFACE	<100	0
SILVERMAN 2	WESTERN EQUITIES	WASHINGTON	6	41 S	13,0 W	24	UNDERGRO	1,000 - 100,000	150
VANDERBILT	EAGER, WALTER C.	WASHINGTON	17	41 S	13,0 W	24	SURFACE	<100	0
BIG JIM-LITTLE J	SMITH & HUNT	WAYNE			0		UNDERGRO	100 - 1,000	50
BLACK JACK	MC CAULEY, EUGENE	WAYNE			0		SURFACE	<100	50
BLUE RIBBON	BRONK, J.F.	WAYNE			0		SURFACE	<100	50
BLUE STONE	BLUE STONE MNG	WAYNE			0		SURFACE	<100	100

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 66

1	MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
*****	UTAH	(CONT'D)	*****							
	BRIDGER JACK	BRIDGER, JACK INC	WAYNE			0		UNDERGRO	100 - 1,000	0
	CONGRESS 28	INDUSTRIES+MINES	WAYNE			0		UNDERGRO	<100	50
	FAIRVIEW	ROBINSON, EDWIN	WAYNE			0		SURFACE	<100	50
	GRAYBAR	ROCKY MTH URA	WAYNE			0		SURFACE	<100	50
	GREEN HORNET	ALAMCO INC.	WAYNE			0		UNDERGRO	100 - 1,000	50
	GREEN MONSTER	CAPITOL REEF URA	WAYNE			0		UNDERGRO	<100	100
	GREENLIGHT + KIM	ELLETT, HUION S.	WAYNE			0		UNDERGRO	<100	150
	HECLA	HECLA MINING CO.	WAYNE			0		UNDERGRO	<100	0
	HELM GROUP	HATCH, DMAIN + CL	WAYNE			0		UNDERGRO	100 - 1,000	150
	KILLERS 1	PENNEL URANIUM	WAYNE			0		UNDERGRO	<100	50
	ISSIANN 1	ISABEL MNG.CO.	WAYNE			0		UNDERGRO	<100	150
	JANUARY	LYMAN, BOB	WAYNE			0		SURFACE	<100	100
	LA VELL 1	SKINER+SKINER+B	WAYNE			0		UNDERGRO	<100	150
	LAST CHANCE 2	GREAT WESTERN UR	WAYNE			0		SURFACE	<100	50
	LOVELL 1	IMPERIAL URANIUM	WAYNE			0		SURFACE	<100	100
	LUSCOMBE 2	PELICAN URANIUM	WAYNE			0		SURFACE	<100	50
	MAMA + PAPA BEAR	P + P ASSOCIATES	WAYNE			0		SURFACE	100 - 1,000	0
	NORTON	ROCKY MTH, URA.	WAYNE			0		SURFACE	<100	50
	OAK CREEK	LYMAN, BOB	WAYNE			0		UNDERGRO	<100	100
	OAK RIDGE	UNKNOWN CONTROL	WAYNE			0		SURFACE	<100	50
	OLD CRO-	WONDERLAND MNG	WAYNE			0		UNDERGRO	<100	0
	ORAL 1 + 2	JENSEN + KOURIS	WAYNE			0		UNDERGRO	<100	50
	POOR BOY 1	ROPER, R.L.	WAYNE			0		SURFACE	<100	100
	SOUTH FORK 1	CROWN LEVEL CO	WAYNE			0		SURFACE	<100	50
	THUNDERING HERD	TACONY URANIUM	WAYNE			0		SURFACE	<100	50
	TURRET 1	UTAH SOUTHERN UR	WAYNE			0		UNDERGRO	<100	150
	WILD HORSE MESA	US COMMON WEALTH	WAYNE			0		UNDERGRO	<100	50
	YELLOW CANARY	CHRISTENSEN+MAT	WAYNE			0		UNDERGRO	100 - 1,000	50
	RIG DANE	CURTIS E.F.	UNKNOWN			0		SURFACE	<100	0
	BULL DOG 1	PEACOCK+NEILSON	UNKNOWN			0		SURFACE	<100	0
	CAT 1	WALLACE, MELVIN J	UNKNOWN			0		UNDERGRO	<100	0
	CUB 1	TUSHER MINING CO	UNKNOWN			0		UNDERGRO	<100	0
	HOLE 4	PETTY MNG + EXPL	UNKNOWN			0		SURFACE	<100	0
	SUSIE WELL	SADDLER, DON	UNKNOWN			0		SURFACE	<100	0
*****	WASHINGTON	*****								
	H.P.S. CLAIMS	GREEN AUGGET MNG	PEND OREILLF			0		SURFACE	100 - 1,000	50
	LOST CREEK 4	ATOMIC SILVER CP	PEND OREILLF			0		SURFACE	1,000 - 100,000	100
	QUARTZ RIDGE GRO	ATOMIC SILVER CP	PEND OREILLF			0		SURFACE	100 - 1,000	0
	CLINE LEASE	UNKNOWN CONTROL	SPOKANE			0		SURFACE	100 - 1,000	50
	DAHL LEASE	HAREM, OSCAR	SPOKANE	11	28 N	44,0 F	33	SURFACE	1,000 - 100,000	50
	DAHL TRACT A	HAREM, MRS. LETHA	SPOKANE			0		SURFACE	100 - 1,000	100
	HANSON LEASE	UNKNOWN CONTROL	SPOKANE			0		SURFACE	100 - 1,000	50
	HERM MOORE LEAS	HAREM, OSCAR	SPOKANE			0		SURFACE	1,000 - 100,000	100
	HUFFMAN LEASE	HUFFMAN, ELLA	SPOKANE			0		SURFACE	100 - 1,000	100
	INGRAM LEASE	UNKNOWN CONTROL	SPOKANE	18	28 N	45,0 E	33	SURFACE	100 - 1,000	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 67

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** WASHINGTON (CONT'D) *****									
LEHMBECKER LEASE	LEHMBECKER, W.E.	SPOKANE			0		SURFACE	100 - 1,000	50
MOURNING LEASE	DIMITROFF, JACK	SPOKANE			0		SURFACE	100 - 1,000	50
LOWLEY LEASE	WESTERN NUCLEAR	STEVENSON			0		SURFACE	100 - 1,000	50
***** WYOMING *****									
AJAX 42	ASPEN MINING CO.	ALBANY			0		SURFACE	<100	0
ALBANY	WATERS + CUNDALL	ALBANY			0		SURFACE	<100	0
DESERT ROSE LEAS	SIXTON, LINCOLN W	ALBANY	9	13 N	78.0 W	06	SURFACE	<100	50
NIGHT OWL	BATTLE AXE MNG. CO	ALBANY			0		SURFACE	<100	50
HASHAWN 2	BONNEVILLE BASIN	BIG HORN	29	45 N	75.0 W	06	SURFACE	<100	50
BROKEN HEART 8	SUPER CUB MNG. CO	BIG HORN			0		SURFACE	<100	50
HIGH NOON 3	SKYWAYS EXPLORAT	BIG HORN			0		UNDERGRO	100 - 1,000	100
HORSESHOE JOHN	SANTA RITA MNG. CO	BIG HORN			0		UNDERGRO	100 - 1,000	50
JET 8	MODERN MINES DEV	BIG HORN	22	58 N	94.0 W	06	UNDERGRO	1,000 - 100,000	200
LEO 1-6	FINK, WYTHAN	BIG HORN	23	58 N	94.0 W	06	UNDERGRO	100 - 1,000	50
MIKE MICKEY	HANSON	BIG HORN	22	58 N	94.0 W	06	UNDERGRO	1,000 - 100,000	50
BANQUINE 12	HOLLAND, ROBERT L	BIG HORN			0		SURFACE	<100	50
TRI-PACER	TITAN MINING CO.	BIG HORN	28	58 N	94.0 W	06	UNDERGRO	1,000 - 100,000	50
ABLE, BAKER, CHARL	HIGHLAND URANIUM	CAMPBELL	23	43 N	76.0 W	06	SURFACE	<100	50
ANOMALY 261, 4-42-	KERR-MCGEE CORP.	CAMPBELL	4	42 N	76.0 W	06	SURFACE	<100	50
ANOMALY 96, 16-45-	KERR-MCGEE CORP.	CAMPBELL	16	43 N	76.0 W	06	SURFACE	<100	0
AXE 10	AXE URANIUM CO.	CAMPBELL	8	43 N	75.0 W	06	SURFACE	100 - 1,000	50
B GROUP	LOGAN CHUFCHILL	CAMPBELL	17	44 N	75.0 W	06	SURFACE	100 - 1,000	50
BAR NONE 33	UNKNOWN CONTROLR	CAMPBELL	20	43 N	75.0 W	06	SURFACE	<100	50
BIG HORN 3	BIG HILL MINING	CAMPBELL			0		SURFACE	<100	0
BILL + EARL CLAI	GARDNER, F.L.	CAMPBELL	33	44 N	75.0 W	06	SURFACE	<100	50
BLACK STAR-BLUF	ROWEN, HEPLDON H.	CAMPBELL	10	44 N	75.0 W	06	SURFACE	1,000 - 100,000	50
BRUCE CLAIMS	ROSENBERGER, GENE	CAMPBELL	9	43 N	76.0 W	06	SURFACE	<100	100
CAMBELIA 2	WHYNOT PROSPECT?	CAMPBELL	28	44 N	75.0 W	06	SURFACE	<100	50
CAMBLIN WEST	HURD H.R. + ASSOC	CAMPBELL	4	43 N	75.0 W	06	SURFACE	100 - 1,000	150
CHRYSOPS 1, 30-41-	KERR-MCGEE CORP.	CAMPBELL	20	41 N	73.0 W	06	SURFACE	1,000 - 100,000	50
COLD SPOT 4 + 5	GILBERT, MORSE C	CAMPBELL	25	43 N	76.0 W	06	SURFACE	1,000 - 100,000	50
COLORADO CHRISTE	ROWEN, HEPLDON H.	CAMPBELL	27	45 N	75.0 W	06	SURFACE	1,000 - 100,000	50
COLUMBUS GROUP	BALL, GRACE D. MRS	CAMPBELL	5	43 N	76.0 W	06	SURFACE	<100	50
CORA + BETTY	HAMLIN EXPL. MNG.	CAMPBELL	29	43 N	75.0 W	06	SURFACE	<100	50
DOE + CHRISTENSE	WESTERN URANIUM	CAMPBELL	21	45 N	76.0 W	06	SURFACE	100 - 1,000	50
DOVE BUTTE	MELSON + SPAIN	CAMPBELL	35	45 N	76.0 W	06	SURFACE	100 - 1,000	50
GAP GROUP	LITTLE PHOENIX UP	CAMPBELL	5	43 N	75.0 W	06	SURFACE	<100	50
HA, RUTH SCHLAUTM	LITTLE STAR MNG	CAMPBELL			75.0 W	06	SURFACE	<100	50
HOC CLAIMS	AMERICAN NUCL. CP	CAMPBELL	3	46 N	76.0 W	06	SURFACE	100 - 1,000	50
INNES LEASE	WESTERN URANIUM	CAMPBELL	11	45 N	75.0 W	06	SURFACE	1,000 - 100,000	50
IRA 1	BUTLER, RYBURN	CAMPBELL			75.0 W	06	SURFACE	100 - 1,000	50
JAKE CME 1920-41-	KERR-MCGEE CORP.	CAMPBELL	19	41 N	73.0 W	06	SURFACE	1,000 - 100,000	50
JEANETTE 1	ROWEN, HEPLDON H.	CAMPBELL	22	45 N	75.0 W	06	SURFACE	1,000 - 100,000	100
JOE 1	CORBELL, E.O.	CAMPBELL			0 W	06	SURFACE	<100	50
JOYCE	GILBERT, MORSE C	CAMPBELL	33	44 N	75.0 W	06	SURFACE	<100	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 68

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** WYOMING (CONT'D) *****									
LAUBY SCHLAUTMAN	LITTLE STAR MNG	CAMPBELL	27	45 N	75.0 W	06	SURFACE	<100	50
LAUR 5, 19	WHYNOT PROSPECT	CAMPBELL	19	43 N	74.0 W	06	SURFACE	<100	50
LUCKY EIGHT 1	BOND, JACF	CAMPBELL	10	44 N	76.0 W	06	SURFACE	<100	50
MARY 1	BOWEN, H.H.	CAMPBELL	22	45 N	75.0 W	06	SURFACE	100 - 1,000	50
MC CLAIN 8	WILK, MAX	CAMPBELL	27	43 N	76.0 W	06	SURFACE	100 - 1,000	50
MIRACLE MILE	MOORE, GEORGE R.	CAMPBELL	27	45 N	75.0 W	06	SURFACE	100 - 1,000	50
MYSTERY 2	HOT SANDS URAN C	CAMPBELL	28	43 N	76.0 W	06	UNDERGRD	<100	50
NERO 5 + 17	WTH STATES MNG.	CAMPBELL	15	43 N	76.0 W	06	SURFACE	100 - 1,000	50
ORR 2	GILA URAN, CORP.	CAMPBELL	5	45 N	74.0 W	06	SURFACE	100 - 1,000	50
OSAGE LEASE	BOWEN, HERLOOY H.	CAMPBELL	32	46 N	74.0 W	06	SURFACE	100 - 1,000	150
PAT 3	MIDDLE BUTTE MNG	CAMPBELL	9	41 N	75.0 W	06	SURFACE	<100	50
PETE GROUP	WESTERN URANIUM	CAMPBELL	3	43 N	76.0 W	06	SURFACE	1,000 - 100,000	50
PRAEST LEASE	GILBERT, MORSE	CAMPBELL	11	45 N	75.0 W	06	SURFACE	<100	50
QUEEN 2	BOWEN, HERLOOY H.	CAMPBELL	22	45 N	75.0 W	06	SURFACE	<100	100
R.H.O. TENRA	WESTERN URANIUM	CAMPBELL	19	43 N	75.0 W	06	SURFACE	<100	50
RENO LEASE	GEORESOURCES, INC	CAMPBELL	34	42 N	73.0 W	06	SURFACE	100 - 1,000	50
SABLE	GILBERT, MORSE	CAMPBELL	9	42 N	75.0 W	06	SURFACE	<100	50
SCOTT 1	W.S. SCHLAUTMAN	CAMPBELL	11	44 N	75.0 W	06	SURFACE	100 - 1,000	100
SEC. 36, 43N-76W	GILBERT, MORSE	CAMPBELL	36	43 N	76.0 W	06	SURFACE	<100	50
SIMMONS 2	WYO ROC DEVEL	CAMPBELL	28	44 N	75.0 W	06	SURFACE	<100	50
STUPLIN MINE	BOWEN, H.H.	CAMPBELL	30	43 N	75.0 W	06	SURFACE	<100	50
SUE 3	PASEK, J.I.	CAMPBELL	17	44 N	75.0 W	06	SURFACE	100 - 1,000	50
SYL DEL MINE	SYL DEL MINES	CAMPBELL	3	43 N	75.0 W	06	SURFACE	100 - 1,000	50
TRIANGLE	AMERICAN URAN, CO	CAMPBELL	28	41 N	75.0 W	06	UNDERGRD	100 - 1,000	100
TRIX GROUP	URANIUM CORPORAT	CAMPBELL	29	43 N	75.0 W	06	SURFACE	<100	50
VAN BUGGENUM SEC	MURD H.R. + ASSOC	CAMPBELL	34	44 N	75.0 W	06	SURFACE	<100	100
WHITE MULE 1	W.S. SCHLAUTMAN	CAMPBELL	10	44 N	75.0 W	06	SURFACE	<100	50
AJO	UNKNOWN	CARBON			0		SURFACE	<100	0
BALD KNOB	WESTERN STAY, URAN	CARBON	14	25 N	81.0 W	06	SURFACE	<100	50
CEDAR HLS-JACKRA	UNION CARBIDE CP	CARBON	32	13 N	92.0 W	06	SURFACE	1,000 - 100,000	50
DAVE LEAN ORE	KGS JOINT VENTUR	CARBON			0		LO-GRADE	>100,000	200
DAVE SEC 9+10+15	GETTY-SKELLY UJV	CARBON	10	27 N	78.0 W	06	SURFACE	>100,000	200
DEL ORO 1	PFLUG, RICHARD I	CARBON			0		SURFACE	<100	50
HELEN MAY	ARMSTRONG, H.T.	CARBON	17	28 N	92.0 W	06	SURFACE	1,000 - 100,000	50
KETCHUM BUTTE L	UNKNOWN CONTROLR	CARBON			0		SURFACE	100 - 1,000	50
LITTLE LILL-JB D	KERR MCGEE	CARBON			0		HL-DUMPS	<100	0
LITTLE MAN 1	UNKNOWN CONTROLR	CARBON			0		UNDERGRD	<100	50
MALL LEASE	TEXAS UTILITIES	CARBON	26	28 N	78.0 W	06	SURFACE	1,000 - 100,000	250
POISON BASIN	HOMESTEAD MINRLS	CARBON	4	12 N	92.0 W	06	SURFACE	1,000 - 100,000	50
SEC 10 LEAN ORE	KGS JOINT VENTUR	CARBON			0		SURFACE	1,000 - 100,000	250
SEC. 36, 25N-W	BAILEY, ROBERT V.	CARBON			0		SURFACE	<100	50
TETON-LUCKY SIRI	HOMESTEAD MINRLS	CARBON	6	12 N	92.0 W	06	SURFACE	1,000 - 100,000	100
TSG 1	KGS JOINT VENTUR	CARBON	9	27 N	78.0 W	06	SURFACE	>100,000	150
TSG HEAP LEACH	KGS JOINT VENTUR	CARBON			0		HL-DUMPS	1,000 - 100,000	250
URANIUM KING 1	PLATT, RE + NORISS	CARBON			0		SURFACE	<100	0
WALKER-GULLIVAN	KERR MCGEE	CARBON	11	27 N	78.0 W	06	SURFACE	>100,000	200
ANOMALY 07 28-37-	KERR-MCGEE CORP.	CONVERSE	28	37 N	73.0 W	06	SURFACE	1,000 - 100,000	150
BETTY	KERR-MCGEE CORP.	CONVERSE	25	41 N	74.0 W	06	SURFACE	<100	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 69

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** MONTANA (CONT'D) *****									
BETZFF LEASE	SMATHERS + DILTS	CONVERSE			0		SURFACE	<100	50
BOX CREEK	UNITED NUCLEAR	CONVERSE	2	35 N	72.0 W	06	SURFACE	1,000 - 100,000	50
CANON BALL 1	UNKNOWN CONTROL.	CONVERSE	1	35 N	72.0 W	06	SURFACE	100 - 1,000	50
CONVERSE MINE	MOORE-HAMBLIN	CONVERSE	14	40 N	75.0 W	06	SURFACE	1,000 - 100,000	150
CURRIER 1	SHAWNEE URAN-MAC	CONVERSE	7	32 N	69.0 W	06	SURFACE	<100	50
HARDY LSE 27-3B-	KERR-MCGEE CORP.	CONVERSE	15	38 N	73.0 W	06	SURFACE	1,000 - 100,000	100
JACKALOPE 13	JACKALOPE OIL+M.	CONVERSE	17	35 N	72.0 W	06	SURFACE	<100	50
JOE GROUP	CRABTREE, JOHN M	CONVERSE	26	38 N	73.0 W	06	SURFACE	1,000 - 100,000	50
JUDY 1-14	COWAN J.L.	CONVERSE	2	32 N	69.0 W	06	SURFACE	<100	50
LAMB	UNKNOWN CONTROL.	CONVERSE	12	35 N	72.0 W	06	SURFACE	1,000 - 100,000	50
LILA BUZZ MINES	NEOCO	CONVERSE	31	35 N	71.0 W	06	UNDERGRO	<100	50
LUCKY ALN MIKEY	STARVFS, JOEL	CONVERSE			0		SURFACE	100 - 1,000	50
MINE 1 (D-B)	JARKETT ENTERPRI	CONVERSE	3	37 N	73.0 W	06	UNDERGRO	<100	50
MINE 20-B 53-37-	KFRP-MCGEE CORP.	CONVERSE	3	37 N	73.0 W	06	UNDERGRO	<100	100
NORTH POLE 1	BRINSON + CAMBLIN	CONVERSE			0 W	06	SURFACE	<100	50
PEYNOLDS T36N-P7	KERR-MCGEE CORP.	CONVERSE	17	36 N	73.0 W	06	SURFACE	<100	50
SEC. 3,37N-73W	KERR-MCGEE CORP.	CONVERSE	3	37 N	73.0 W	06	SURFACE	1,000 - 100,000	100
SEC. 9,37N-73W	KERR-MCGEE CORP.	CONVERSE	9	37 N	73.0 W	06	SURFACE	1,000 - 100,000	100
SEC.10,37N-73W 0	KERR-MCGEE CORP.	CONVERSE	10	37 N	73.0 W	06	SURFACE	1,000 - 100,000	50
SEC.15,37N-73W	KERR-MCGEE CORP.	CONVERSE	15	37 N	73.0 W	06	SURFACE	100 - 1,000	150
SEC.16,37N-73W	KERR-MCGEE CORP.	CONVERSE	16	37 N	73.0 W	06	SURFACE	1,000 - 100,000	100
SEC.21,37N-73W W	KERR-MCGEE CORP.	CONVERSE	21	37 N	73.0 W	06	SURFACE	1,000 - 100,000	100
SHEEP-SHEP GROUP	ROVEN,HERIDON H.	CONVERSE	12	35 N	72.0 W	06	SURFACE	<100	50
SPOOK	HORNBUCKLE,DICK	CONVERSE	28	38 N	73.0 W	06	SURFACE	>100,000	100
SPRING GROUP	ALLSUP, DARREL B	CONVERSE	30	40 N	74.0 W	06	SURFACE	<100	50
STROCK-ROSCAT-LE	JOHN PETERSON AS	CONVERSE	27	33 N	72.0 W	06	SURFACE	<100	50
TRAIL CREEK GROU	PHINLEY,B.D.ETAL	CONVERSE			0		UNDERGRO	<100	50
TURNERCREST	KERR-MCGEE CORP.	CONVERSE	25	41 N	74.0 W	06	SURFACE	1,000 - 100,000	50
ZEE 1 + 2	KERR-MCGEE CORP.	CONVERSE	27	38 N	73.0 W	06	SURFACE	1,000 - 100,000	100
A.A.H.	UNKNOWN CONTROL	CROOK			0		SURFACE	<100	0
A.A.SISSON	KERR MCGEE	CROOK	7	54 N	60.0 W	06	SURFACE	<100	50
ACKERMAN LEASE	FEDERAL RESOURCE	CROOK	34	56 N	66.0 W	06	SURFACE	1,000 - 100,000	50
BUSFIELD LEASE	FEDERAL RESOURCE	CROOK	26	56 N	66.0 W	06	SURFACE	1,000 - 100,000	50
CASIN CREEK 6	TRI STATE MINING	CROOK			0		SURFACE	<100	50
DENNIS 2 LEASE	MICHAUD MINING C	CROOK	3	55 N	67.0 W	06	SURFACE	1,000 - 100,000	50
GRIFFITH LEASE	GRIFFITH,LYLE	CROOK	26	52 N	68.0 W	06	UNDERGRO	1,000 - 100,000	100
HAUBER MINE UG	HOMESTAKE MNG CO	CROOK	2	55 N	67.0 W	06	UNDERGRO	>100,000	400
HELMER RANCH	CHRIS HELMER	CROOK	19	54 N	60.0 W	06	SURFACE	100 - 1,000	0
HOLMES LEASE	TENN,VALLEY AUTH	CROOK	18	54 N	65.0 W	06	SURFACE	1,000 - 100,000	100
HOMESTAKE 1-4	GEORESOURCE,INC	CROOK	26	52 N	66.0 W	06	SURFACE	1,000 - 100,000	50
K GROUP	TENN,VALLEY AUTH	CROOK	26	56 N	66.0 W	06	SURFACE	1,000 - 100,000	100
LAYMON LEASE	FEDERAL RESOURCE	CROOK	27	52 N	66.0 W	06	UNDERGRO	1,000 - 100,000	200
LEWIS DENNIS LEA	LITTLE MQ MINING	CROOK			0		SURFACE	1,000 - 100,000	50
MEYERS LEASE	TENN,VALLEY AUTH	CROOK	21	52 N	65.0 W	06	UNDERGRO	100 - 1,000	50
NEW HAVEN 20-SEC	HOMESTAKE MNG CO	CROOK	10	53 N	67.0 W	06	UNDERGRO	1,000 - 100,000	50
NORTH SLOPE	UNKNOWN CONTROL	CROOK	7	54 N	60.0 W	06	SURFACE	<100	50
POISON CREEK CLA	BAKER, IRVIN	CROOK	12	54 N	67.0 W	06	SURFACE	<100	50
SADDLE GROUP	LYNARD +KIPLEAN	CROOK	22	53 N	65.0 W	06	SURFACE	<100	50

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 70

WINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** WYOMING (CONT'D) *****									
STATE LEASE	STATE OF WYOMING	CROOK	17	54 N	60.0 W	06	SURFACE	100 - 1,000	50
STORM MINE	HONESTAKE MNG CO	CROOK	6	55 N	66.0 W	06	SURFACE	1,000 - 100,000	50
T.F. WOLF SKILL LS	ROUNDS + SHPNER	CROOK	15	54 N	66.0 W	06	SURFACE	1,000 - 100,000	50
VIRGINIA 18	ROCKY MTH. URAN.	CROOK	32	57 N	64.0 W	06	SURFACE	<100	0
ANDRIA (C PIT)	FEDERAL AMERICAN	FREMONT			0		SURFACE	1,000 - 100,000	0
ANDRIA GROUP	FEDERAL AMERICAN	FREMONT	1	32 N	91.0 W	06	SURFACE	1,000 - 100,000	50
ARROWHEAD (COP. MT)	ROCKY MTH. ENERGY	FREMONT	28	40 N	92.0 W	06	UNDERGRO	>100,000	100
B.H. 1	HAYS, RICHARD	FREMONT	8	33 N	95.0 W	06	SURFACE	<100	50
B.H. HALL-BONANZA	FORD, ROBERT + ASSOC	FREMONT	33	40 N	92.0 W	06	SURFACE	1,000 - 100,000	50
BEATRICE 3	ARMSTRONG, H.T.	FREMONT			0		SURFACE	<100	50
BIG RED 4	BRIDGER MNG CO	FREMONT			0		SURFACE	<100	50
BLACKSTONE 51	WESTERN NUCLEAR	FREMONT	22	33 N	89.0 W	06	SURFACE	<100	50
BLARCO	J.M. HADE	FREMONT	12	32 N	91.0 W	06	SURFACE	1,000 - 100,000	0
BLUE ROCK-RED NO	VIPONT MINING CO	FREMONT	9	33 N	89.0 W	06	SURFACE	1,000 - 100,000	50
BOUNTIFUL 1, 9 + 10	WESTERN NUCLEAR	FREMONT	22	33 N	89.0 W	06	SURFACE	1,000 - 100,000	150
BULLRUSH DUMP	WESTERN NUCLEAR	FREMONT			0		LONGRADE	100 - 1,000	0
BULLRUSH GROUP	WESTERN NUCLEAR	FREMONT	29	33 N	90.0 W	06	SURFACE	>100,000	100
CAL 15	FEDERAL AMERICAN	FREMONT	29	33 N	90.0 W	06	SURFACE	1,000 - 100,000	200
CINCH 15	WILLIAMS, A.E.	FREMONT			0		SURFACE	<100	50
CLEANUP MATERIAL	PETERS, REX	FREMONT			0		MISC.-PB	100 - 1,000	0
DAY 26	WESTERN NUCLEAR	FREMONT	24	32 N	91.0 W	06	SURFACE	>100,000	350
DAY-BERGER LEASE	REEVES, M.J.	FREMONT	3	39 N	92.0 W	06	UNDERGRO	1,000 - 100,000	50
DAY-LOMA	ENERGY FUELS NUC	FREMONT	24	32 N	91.0 W	06	SURFACE	>100,000	100
DISCOVERY	WELLS, JOHN	FREMONT	29	40 N	92.0 W	06	SURFACE	<100	50
DONNA LEE	PETERS, REX	FREMONT	5	33 N	93.0 W	06	SURFACE	<100	50
DUBOIS 1	KAYE MINERALS INC	FREMONT	24	41 N	08.0 W	06	SURFACE	<100	50
EUREKA-IMPERIAL	WYOMING NUCLEAR	FREMONT	29	33 N	90.0 W	06	SURFACE	1,000 - 100,000	100
FANNIE 1-4	STAMM, ED	FREMONT	2	39 N	92.0 W	06	SURFACE	100 - 1,000	50
FEDERAL	FEDERAL AMERICAN	FREMONT			0		SURFACE	1,000 - 100,000	100
FRAZIER-LAMAC GR	WESTERN NUCLEAR	FREMONT	27	33 N	90.0 W	06	SURFACE	>100,000	300
GEORGE + VEP GRO	FEDERAL AMERICAN	FREMONT	22	33 N	90.0 W	6	SURFACE	1,000 - 100,000	150
GEORGE 1 AND 2	UNION CARBIDE CP	FREMONT	30	33 N	90.0 W	06	SURFACE	1,000 - 100,000	100
GEORGE 12 + 14	UNION CARBIDE CP	FREMONT	30	33 N	90.0 W	06	SURFACE	1,000 - 100,000	50
HADA 17	REX PETERS	FREMONT			0		SURFACE	100 - 1,000	0
HAPPY 19	HACK LANG URAN. C	FREMONT			0		SURFACE	<100	50
HAZEL GROUP	WESTERN NUCLEAR	FREMONT	9	28 N	92.0 W	06	SURFACE	100 - 1,000	50
HEAP LEACH (UTAH)	PATHFINDER	FREMONT			0		HL-DUMPS	<100	200
HESITATION	GENEVA URAN CORP	FREMONT	27	40 N	92.0 W	06	SURFACE	<100	50
HOPE-STAR	UNKNOWN CONTROLR	FREMONT	12	32 N	90.0 W	06	SURFACE	1,000 - 100,000	150
HUNTER LEASE	DALE B. LEVI	FREMONT	1	32 N	91.0 W	06	SURFACE	1,000 - 100,000	100
IDOTTS DELIGHT	PATHFINDER	FREMONT	12	32 N	91.0 W	06	SURFACE	<100	300
JACK 2	UNION CARBIDE CP	FREMONT	6	32 N	91.0 W	06	UNDERGRO	1,000 - 100,000	50
JAY GROUP	FEDERAL AMERICAN	FREMONT	26	33 N	90.0 W	06	SURFACE	>100,000	200
JEM 1	HERRSTROM H.O.	FREMONT	15	39 N	92.0 W	06	SURFACE	<100	50
JOHN-GUNNEL	FEDERAL AMERICAN	FREMONT	25	33 N	90.0 W	06	SURFACE	>100,000	100
JOY GROUP	UNION CARBIDE CP	FREMONT	6	32 N	90.0 W	06	SURFACE	>100,000	150
LAST CHANCE	SAN JUAN URAN. EX	FREMONT			0		SURFACE	<100	50
LITTLE STORY 6	LITTLE STORY GRP	FREMONT			0		SURFACE	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 71

WINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** WYOMING (CONT'D) *****									
LOMA HEAP LEACH	WESTERN NUCLEAR	FREMONT			0		HL-DUMPS	<100	250
MIDNIGHT 1	BIG SKY URANIUM	FREMONT			0		SURFACE	<100	0
NELS GROUP	UNKNOWN CONTROL	FREMONT	13	32 N	91.0 W	06	SURFACE	1,000 - 100,000	150
NEVIN GF4	JAY, G.I.	FREMONT	33	40 N	92.0 W	06	SURFACE	100 - 1,000	50
PAN-PAT	HENRY HUDSPETH	FREMONT	34	32 N	99.0 W	06	SURFACE	100 - 1,000	50
PEACH GROUP(UG)	AMERICAN NUCLEAR	FREMONT	2	32 N	90.0 W	06	UNDERGRO	1,000 - 100,000	400
PHIL (P-1)	P.JENKINS-HAAD	FREMONT	12	32 N	91.0 W	06	SURFACE	1,000 - 100,000	50
PIT A-6	UNION CARBIDE CP	FREMONT	14	33 N	89.0 W	06	SURFACE	1,000 - 100,000	150
PIT M-1	UNION CARBIDE CP	FREMONT			0		SURFACE	100 - 1,000	0
QUEENS GROUP	SWANTON, L.F	FREMONT			0		SURFACE	<100	50
RAY 6	WESTERN NUCLEAR	FREMONT	4	32 N	90.0 W	06	SURFACE	<100	50
REX CLAIMS	PATHEINDER	FREMONT	11	32 N	91.0 W	06	SURFACE	1,000 - 100,000	100
RIM GROUP	PATHEINDER	FREMONT	9	33 N	89.0 W	06	SURFACE	>100,000	150
RING + SANDUSKY	WAR BONNET URAN.	FREMONT	10	30 N	97.0 W	06	SURFACE	<100	50
ROX GROUP	ENERGY FUELS NUC	FREMONT	21	33 N	89.0 W	06	UNDERGRO	1,000 - 100,000	350
ROX LEACH	ENERGY FUELS NUC	FREMONT			0		HL-DUMPS	<100	0
SAGEBUSH-TRIST	FEDERAL AMERICA	FREMONT	32	33 N	90.0 W	06	UNDERGRO	>100,000	200
SEC.16,33.-89A	GREAT BASIN PETR	FREMONT	16	33 N	89.0 W	06	SURFACE	>100,000	50
STAN	UNION CARBIDE CP	FREMONT	28	33 N	90.0 W	06	SURFACE	1,000 - 100,000	100
STOCKPILE CLEAVU	WESTERN NUCLEAR	FREMONT			0		MISC.-PB	100 - 1,000	0
SUN-SNOWHALL	WESTERN NUCLEAR	FREMONT	28	28 N	92.0 W	6	SURFACE	1,000 - 100,000	250
TORCHY	WOOTEN, CHARLENF	FREMONT			0		UNDERGRO	<100	0
TWO C + BOZO	JADEN, JIN	FREMONT	17	30 N	96.0 W	06	SURFACE	<100	0
WASHAKIE	ALLIED NUCLEAR	FREMONT	10	32 N	94.0 W	06	SURFACE	1,000 - 100,000	50
WASHOUT 1	WALKER, RAYMOND	FREMONT			0		SURFACE	<100	0
YELLOW RUCK-BEND	VIPONT MINING CO	FREMONT	4	33 N	89.0 W	06	SURFACE	100 - 1,000	50
YELLOW-STONE 1	SWANTON, L.F	FREMONT	31	33 N	89.0 W	06	SURFACE	<100	50
ANTELOPE GROUP	GEORESOURCES,INC	JOHNSON	10	43 N	77.0 W	06	SURFACE	1,000 - 100,000	50
HANNA GROUP	HANLIN EXPL.MNG.	JOHNSON	14	42 N	77.0 W	06	SURFACE	1,000 - 100,000	50
JERRIE MARIE GRO	LITTLE GIANT URA	JOHNSON			0		SURFACE	<100	0
KELL ROY 3 + 4	YOUNG, ROY O.	JOHNSON		42 N	78.0 W	06	SURFACE	<100	50
LUCKY SEVEN 2	TREBELCOCK + KWI	JOHNSON	20	44 N	77.0 W	06	SURFACE	<100	50
MASEK 1	ATOE MINING CO.	JOHNSON	3	44 N	83.0 W	06	SURFACE	<100	50
MISSEY GROUP	AMERICAN NUCLEA	JOHNSON	3	42 N	78.0 W	06	SURFACE	100 - 1,000	50
MOF 14(CANDALY 8	UNKNOWN CONTROL	JOHNSON	19	45 N	76.0 W	06	SURFACE	100 - 1,000	50
N,STAR ISF(COYMO	SPEERBERG,JOHN J.	JOHNSON	11	43 N	80.0 W	06	SURFACE	<100	150
P CLAIM 8-45-	KERR-MCGEE CORP.	JOHNSON	8	45 N	76.0 W	06	UNDERGRO	<100	150
RED ROCK 2	SOUTH PFAK MNG.	JOHNSON		N	0 W	06	SURFACE	<100	50
SNAKEITE 1-3	GEORESOURCES,INC	JOHNSON	3	43 N	77.0 W	06	SURFACE	100 - 1,000	50
T BONE	KERR-MCGEE CORP.	JOHNSON	31	42 N	75.0 W	06	SURFACE	<100	50
VANCE 1	ROSENBERGER,DUAN	JOHNSON		N	0 W	06	SURFACE	<100	50
YANK 4	SAN JUAN URAN,EX	JOHNSON		N	0 W	06	SURFACE	<100	50
ANTELOPE 4	SPACE AGE NUCLEA	NATRONA			0		SURFACE	<100	50
BAKFR 3,4,5+6	JONES +YANGILDER	NATRONA	5	39 N	77.0 W	06	SURFACE	1,000 - 100,000	50
BRIDGER TRAIL	HIGHLAND URAN,IN	NATRONA			0		SURFACE	<100	50
BUSS-RUSS	FEDERAL AMERICAN	NATRONA	27	33 N	89.0 W	06	UNDERGRO	1,000 - 100,000	350
DRY LAKE	GEORESOURCES,INC	NATRONA	33	35 N	79.0 W	06	SURFACE	<100	100
HEAP LEACH B-2	UNION CARBIDE CP	NATRONA			0		HL-DUMPS	<100	150

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 72

WINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** MINEING (CONT'D) *****									
MARS GROUP	FEDERAL AMERICAN	NATRONA	24	33 N	89.0 N	06	SURFACE	1,000 - 100,000	100
PAY - ALJIE	UNION CARBIDE	NATRONA	15	33 N	89.0 N	06	SURFACE	>100,000	100
PINE TREE GROUP	MINERALS EXPL.CO	NATRONA			0		SURFACE	<100	50
PIT #2	UNION CARBIDE CP	NATRONA			0		SURFACE	1,000 - 100,000	50
PIT #1	UNION CARBIDE CP	NATRONA			0		SURFACE	1,000 - 100,000	150
PIT #2	UNION CARBIDE CP	NATRONA	15	33 N	89.0 N	06	SURFACE	>100,000	200
RIDGE 1 + TETON	WEITZ, JOE	NATRONA	25	34 N	89.0 N	06	SURFACE	1,000 - 100,000	50
SHUT P	THIN ARROW PETRO	NATRONA			0		UNDERGRU	100 - 1,000	50
SPA SAGE	CUNNINGHAM MIN	NATRONA	34	37 N	89.0 N	06	SURFACE	1,000 - 100,000	50
TEL GROUP	FEDERAL AMERICA	NATRONA	72	33 N	89.0 N	06	SURFACE	>100,000	150
VICA PITS	FEDERAL AMERICAN	NATRONA	21	33 N	89.0 N	06	SURFACE	>100,000	100
ADFX 1	WOLFF, OTTO	NIOBRARA			0		SURFACE	<100	50
BOHEP 1	ASSOCIATED OREVEL	NIOBRARA			0		SURFACE	<100	0
CLAIM 1	WESTER URANIUM	NIOBRARA			0		SURFACE	<100	50
WFG 1	WFG, J.W.	NIOBRARA			0		SURFACE	<100	50
MILLER LEASE 7	WALKER, REEF	NIOBRARA			0		SURFACE	<100	0
MOORE 1	MURPHY, MURPHY	NIOBRARA	23	36 N	85.0 N	06	SURFACE	<100	50
NEVADA 1	JOHNSON + WALKER	NIOBRARA			0		SURFACE	100 - 1,000	50
W4 MOPE	BARPAUGH PAC-AN	NIOBRARA	24	34 N	87.0 N	06	SURFACE	<100	50
POTTER LEASE	BEKUS-FITZGERY	NIOBRARA	23	31 N	84.0 N	06	UNDERGRU	<100	50
PEER LEASE	SMITH, W.W.	NIOBRARA			0		SURFACE	<100	50
SILVER CLIFF PAL	LEMAN, MATTHEW H	NIOBRARA			0		UNDERGRU	100 - 1,000	50
WALT CLAYS	MARTIN, F. + PUSEY	NIOBRARA			0		SURFACE	100 - 1,000	50
WEIGAND LEASE	ROCKY MOUNTAIN, WILL.	NIOBRARA			0		SURFACE	100 - 1,000	50
PARD 4	UNKNOWN + CONTROL	SUNLETT			0		SURFACE	100 - 1,000	50
GOLDEN ARROW	LUCKY JOHN MNG.C	SAFFETATER	30	26 N	94.0 W	06	SURFACE	<100	50
LONG WOLF 1	BLACK SHEEP URAN	SAFFETATER			0		UNDERGRU	<100	50
LUCKY IMP 15	BLACK BUTTES URAN	SAFFETATER			0		SURFACE	<100	50
SEC. 36, 24, - 24.	FRYMAN URAN.CO.	SWEETWATER	36	24 N	94.0 N	06	SURFACE	100 - 1,000	50
ARLINGTON 1	PACIFIC, FIFER	WASHAKIE			0		SURFACE	<100	0
SCORPIO	S. + W. MNG. + EXPL	WASHAKIE			0		UNDERGRU	<100	50
SEC. 16-42, -60	WYOMING, STATE OF	WESTON	16	42 N	89.0 N	06	SURFACE	100 - 1,000	100
***** UNKNOWN *****									
BELT MINE	WICKERSON, ALBERT	UNKNOWN			0		UNDERGRU	<100	0
BLACK SHEEP	ARMINE, A.H.	UNKNOWN			0		SURFACE	<100	0
CRUP OUT	FULLER, BLANCHE	UNKNOWN			0		SURFACE	<100	0
CUP 1	WAGNER BANKS + HO	UNKNOWN			0		UNDERGRU	<100	0
DIEGO 11	C.U. DEVELOPMENT	UNKNOWN			0		UNDERGRU	<100	0
FRIDAY 1	LONGWILL, J.E., JR	UNKNOWN			0		UNDERGRU	<100	0
W4 ROCKS 1	WHITE KING URAN	UNKNOWN			0		UNDERGRU	<100	0
DE ROY BENSON	BENSON, LEROY	UNKNOWN			0		SURFACE	<100	0
RAFFINATE-CAYADA	SHEPPITT GORDON	UNKNOWN			0		MISC.-PB	100 - 1,000	0
RAFFINATE-CONGO	COMMERCIAL DI	UNKNOWN			0		MISC.-PB	1,000 - 100,000	0
STANLEY 1	SOUTHERN ILL. NAT	UNKNOWN			0		SURFACE	<100	0
URANIUM	AYERS, EVERETT	UNKNOWN			0		SURFACE	<100	0

INACTIVE URANIUM MINES IN THE UNITED STATES
SOURCE: DOE, GRAND JUNCTION, COLORADO

PAGE 73

MINE NAME	CONTROLLER NAME	COUNTY	SEC.	TOWNSHIP	RANGE	MERID.	MINING METHOD	TOTAL PRODUCTION (TONS AS OF 01/01/79)	DEPTH (FT.)
***** UNKNOWN	(CONT'D)	*****							
WM STAHR	F. C. EXPLORATIO	UNKNOWN			0		SURFACE	<100	0

APPENDIX G

GENERAL OBSERVATIONS OF URANIUM MINE SITES IN
COLORADO, NEW MEXICO, TEXAS, AND WYOMING

G.1. General observations on inactive uranium mine sites in Colorado,
New Mexico, Texas, and Wyoming

"Walk through" surveys were conducted at selected inactive uranium mines in Colorado, New Mexico, Texas, and Wyoming. The primary purpose of the "walk through" surveys was to note and describe the general environmental conditions of the mines. Limited gamma radiation rate measurements were made at each site and estimates were made of the volume and area of the mine wastes. In addition, each waste area was observed for indications of wind and water erosion and to see if the mine entry and vents were open to the atmosphere.

G.1.1 Colorado

The surveys were conducted at mining areas near Uravan and Boulder, Colorado. Each area is listed, and the survey results at each mine are discussed.

G.1.1.1 Uravan Area

The Uravan area lies within the Uravan Mineral Belt, which is situated on the Utah-Colorado border, encompassing parts of Mesa, Montrose, and San Miguel Counties in Colorado and Grand and San Juan Counties in Utah. Uranium has been mined in the belt from the Salt Wash Member of the Morrison Formation since 1900. About 150 mines were being worked in the belt in 1978. Three companies have announced their intention to build mills in the belt area (Wh78). Ore deposits are found mainly in sandstone lenses which are up to 1,600 m wide and average about 15 m thick. The ore deposits range in size from a few MT in the form of a fossil log to many thousands of MT. The ore deposits generally range in thickness from a few centimeters to 7.6 m. Irregularly shaped, they can be found almost anywhere within the sandstone lenses. One study of the Salt Wash deposits indicated that 70 percent of the deposits contained less than 2,700 MT of ore each (Wh78).

Since the ore bodies occur as rolls, pods, or tabular masses, their size precludes the use of a prearranged and uniform stoping system. Mining practices were to simply follow the ore and leave open stopes behind. Consequently, a large number of relatively small mines have been operated in this mineral belt. One mill processed ore from 200 mines which have produced ore ranging from 91 to 910,000 MT.

A substantial fraction of the inactive mines are located in or near the Uravan Mineral Belt and are listed by State and county. A total of 1860 inactive uranium mines, or about 57 percent of all inactive uranium mines in the entire United States, are located in or near the Uravan Mineral Belt.

<u>State</u>	<u>County</u>	<u>Inactive Mines</u>	
		<u>Surface</u>	<u>Underground</u>
Colorado	Mesa	77	109
Colorado	Montrose	76	404
Colorado	San Miguel	70	269
Utah	Emery	60	126
Utah	Grand	69	95
Utah	San Juan	<u>140</u>	<u>365</u>
	Total	492	1368

G.1.1.1.1 Mine 1

This mine had a vertical shaft which was barricaded to prevent livestock from falling into it; however, it remains open to the atmosphere. About 13,800 cubic meters of mine wastes were dumped on a downslope area adjacent to the shaft presently covering an area of about 0.1 hectare (Fig. G.1). There was evidence of wind and water erosion of the wastes. Exposure rates, measured 0.914 m above the wastes, ranged from 140-170 $\mu\text{R/hr}$ with several spots reaching 250 $\mu\text{R/hr}$. Several small adjacent waste piles had exposure rates of 110 $\mu\text{R/hr}$. No springs or standing water were observed near the wastes.

G.1.1.1.2 Mine 2

This rim mine (Fig. G.2) faces the San Miguel river valley and produced about 1,200 cubic meters of wastes, which were dumped down the canyon wall and presently cover an area of about 0.4 hectare. Exposure rates measured near the dump point were about 200 $\mu\text{R/hr}$; measurements on the road around the ore bins ranged from 50-150 $\mu\text{R/hr}$. The mine entry remains open to the atmosphere. There was evidence of wind and water erosion of the mine wastes.

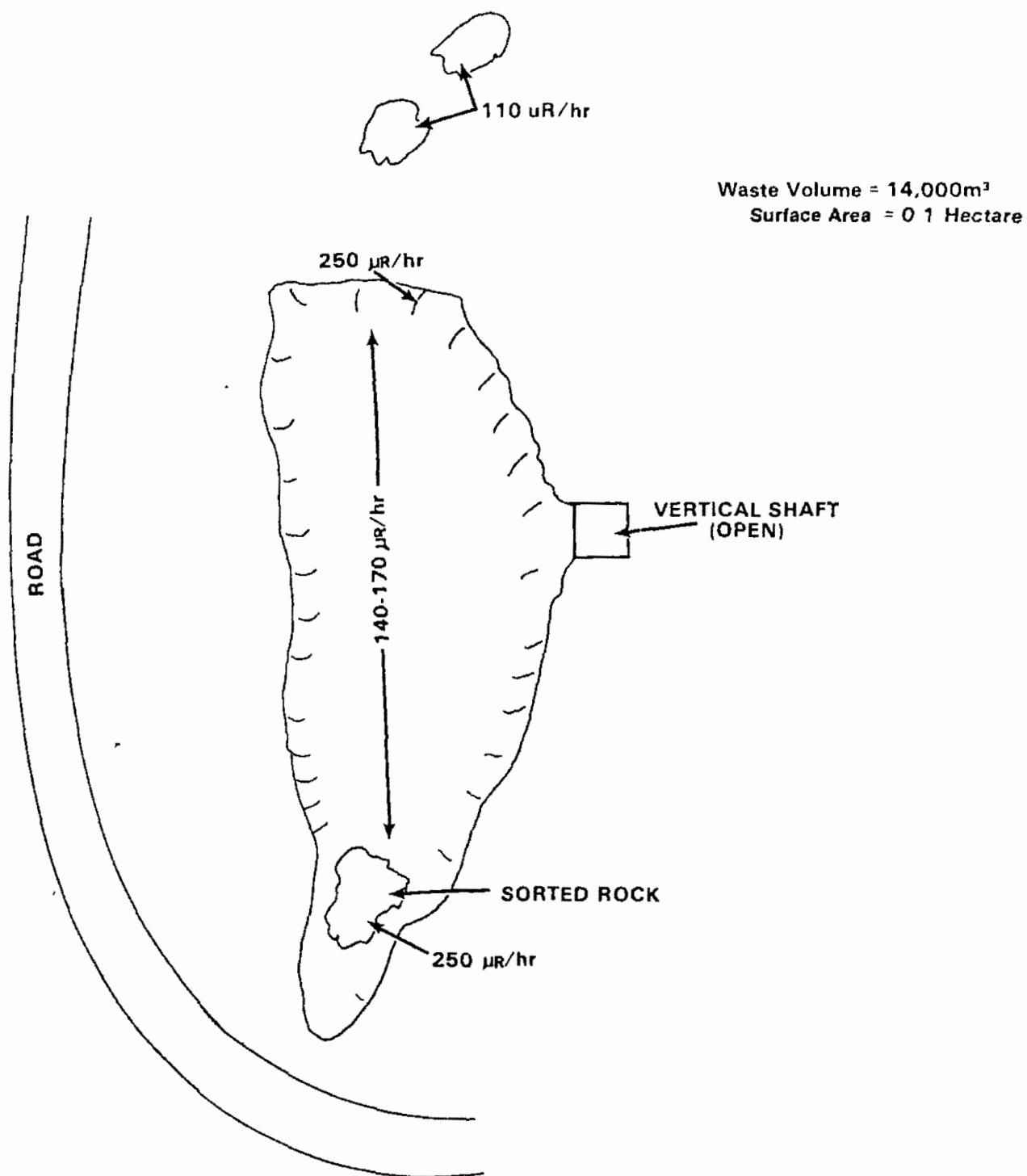


Figure G.1 Plan view of inactive underground uranium mine No. 1, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado

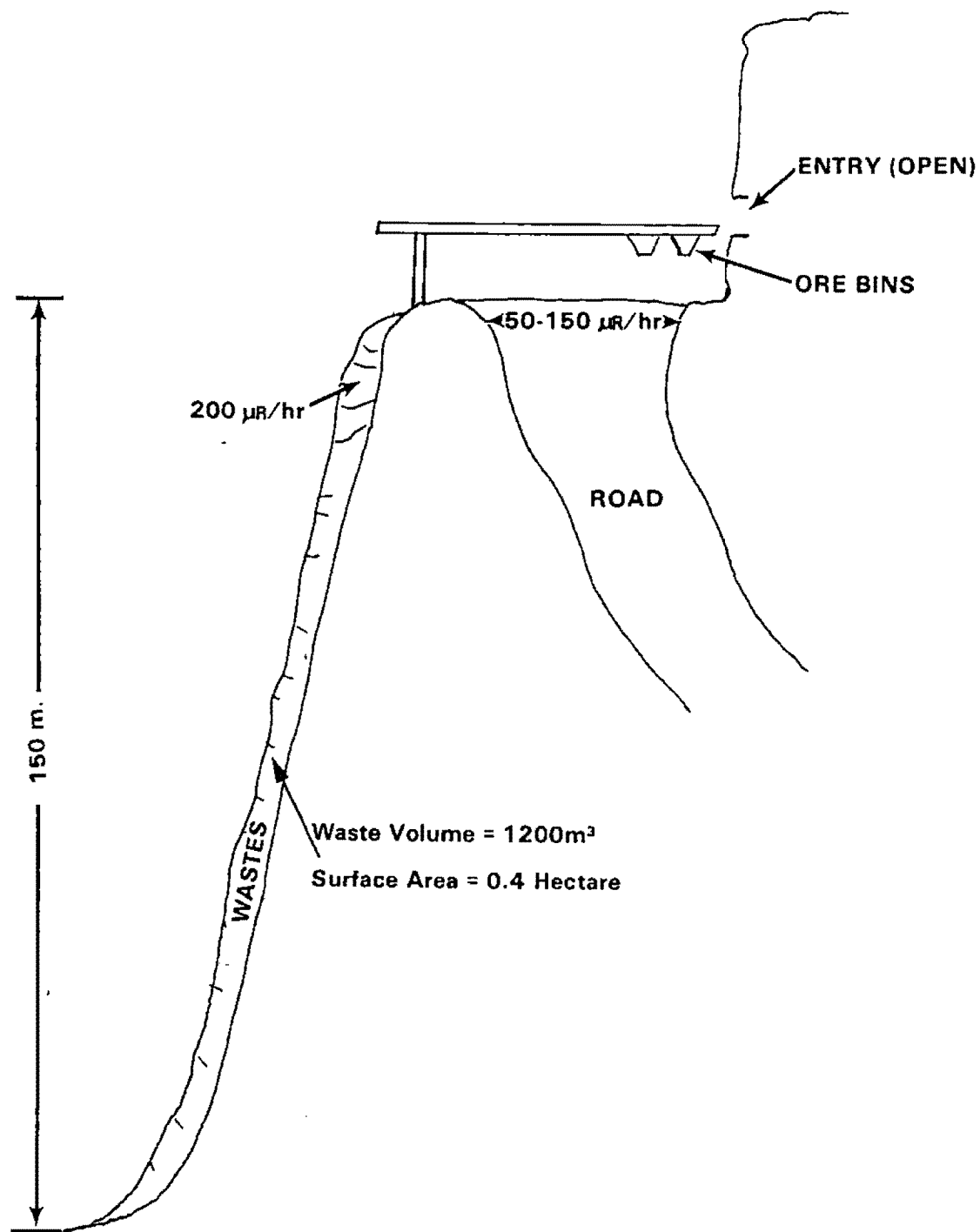


Figure G.2 Sectional view of inactive underground uranium mine No. 2, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado

G.1.1.1.3 Mine 3

This mine has an incline entry that remains open to the atmosphere. Two waste piles were built up to support ore bins (Fig. G.3). About 38,300 cubic meters of wastes, which have a surface area of about two hectares, are contained in the two piles. Part of the waste piles extend into an adjacent wash and are subject to water erosion. Wind erosion of the wastes was also evident. Gamma exposure rates on the waste piles ranged from 120-150 $\mu\text{R/hr}$, while areas adjacent to the piles were about 160 $\mu\text{R/hr}$. The exposure rate on the mine access road was 80 $\mu\text{R/hr}$.

G.1.1.1.4 Mine 4

This was a rim mine with a portal remaining open. The mine wastes volume was about 6,100 cubic meters covering an area of about 0.4 hectares (Fig. G.4). Wastes have eroded down the slope, through a drain pipe under the highway, and into the San Miguel River. Exposure rates on the access road and under the ore bins were about 70 $\mu\text{R/hr}$. Wind erosion of the wastes was also evident.

G.1.1.1.5 Mine 5

This mine contained a vertical shaft used for forced ventilation of connecting mines (Fig. G.5). An undetermined amount of low-grade ore had been dumped in small piles covering an area of about 5 hectares and was later removed for milling. The two remaining piles cover about 1.2 hectares and contain about 76,500 m^3 of protore and barren wastes including clean-up materials from the 5 hectare area. Gamma exposure rates over the former waste area ranged from 50-150 $\mu\text{R/hr}$. Exposure rates over the consolidated piles ranged from 50-220 $\mu\text{R/hr}$. Wind and water erosion were evident at both the former and present waste storage areas.

G.1.1.1.6 Mine 6

This mine had a vertical shaft used to force ventilate connecting active mines (Fig. G.6). About 45,900 cubic meters of wastes were dumped on a downslope adjacent to the mine shaft and now cover about 0.4 hectare of ground. Gamma exposure rates measured over the waste pile ranged from 180-220 $\mu\text{R/hr}$. Runoff and wind erosion of the wastes were evident. Some of the runoff appeared to have entered a nearby stock pond.

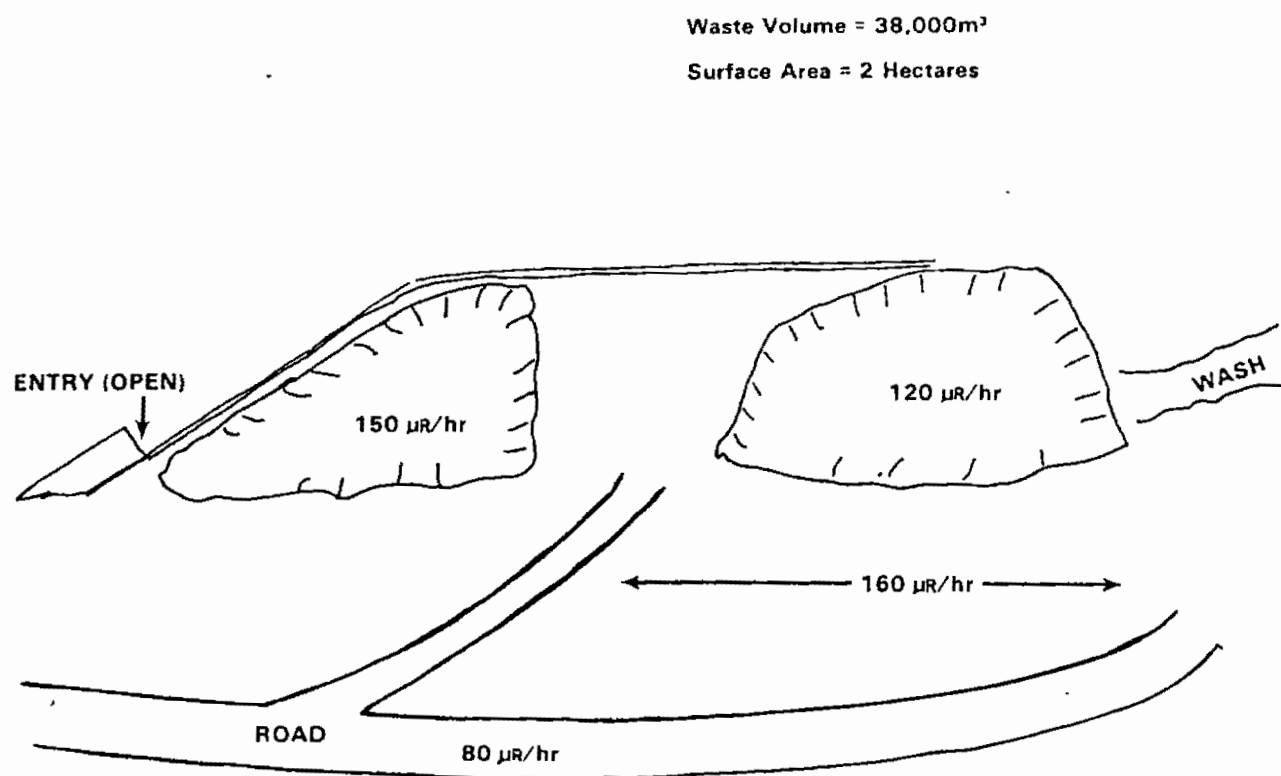


Figure G.3 Plan view of inactive underground uranium mine No. 3, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado

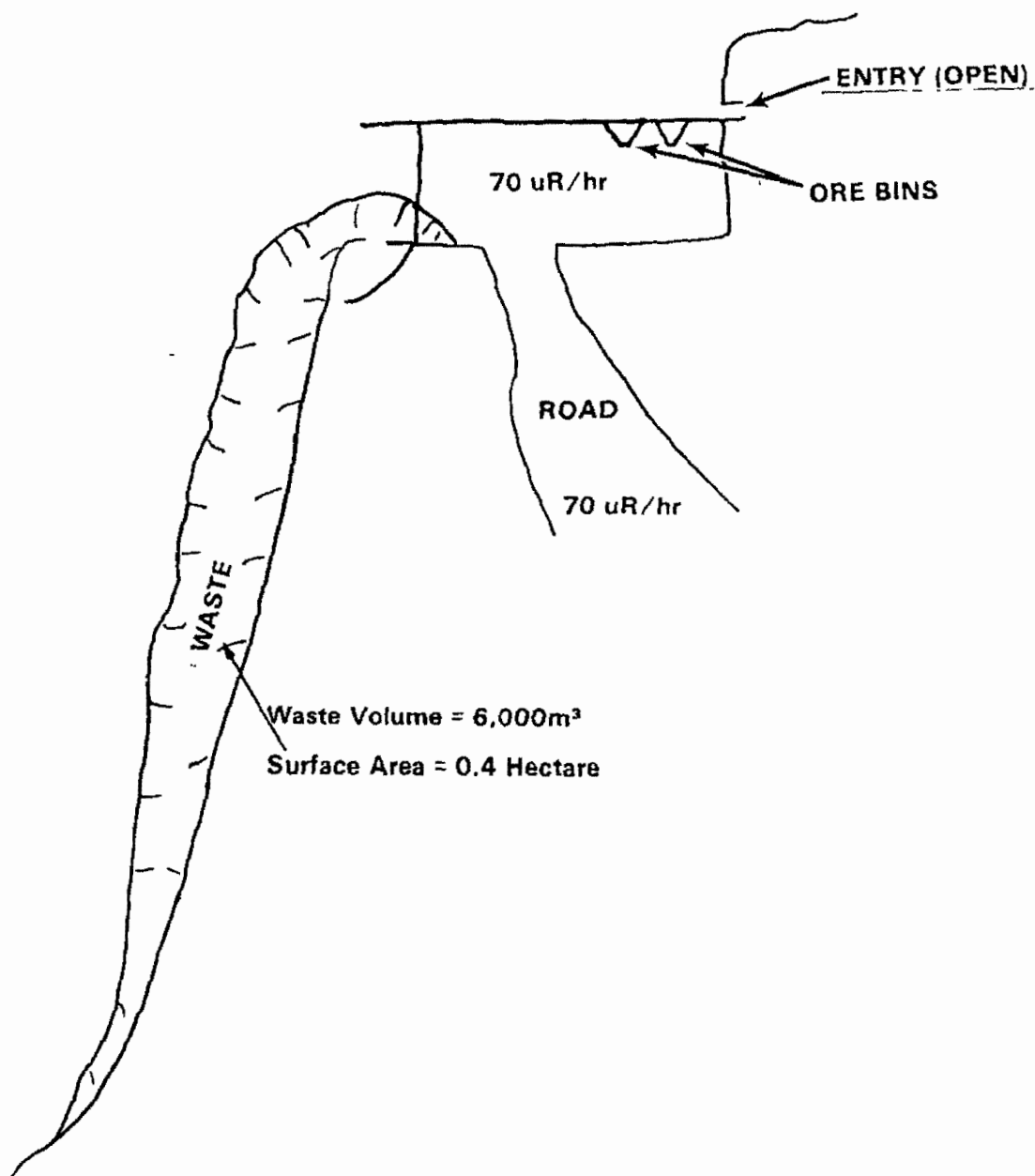


Figure G.4 Sectional view of inactive underground uranium mine No. 4, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado

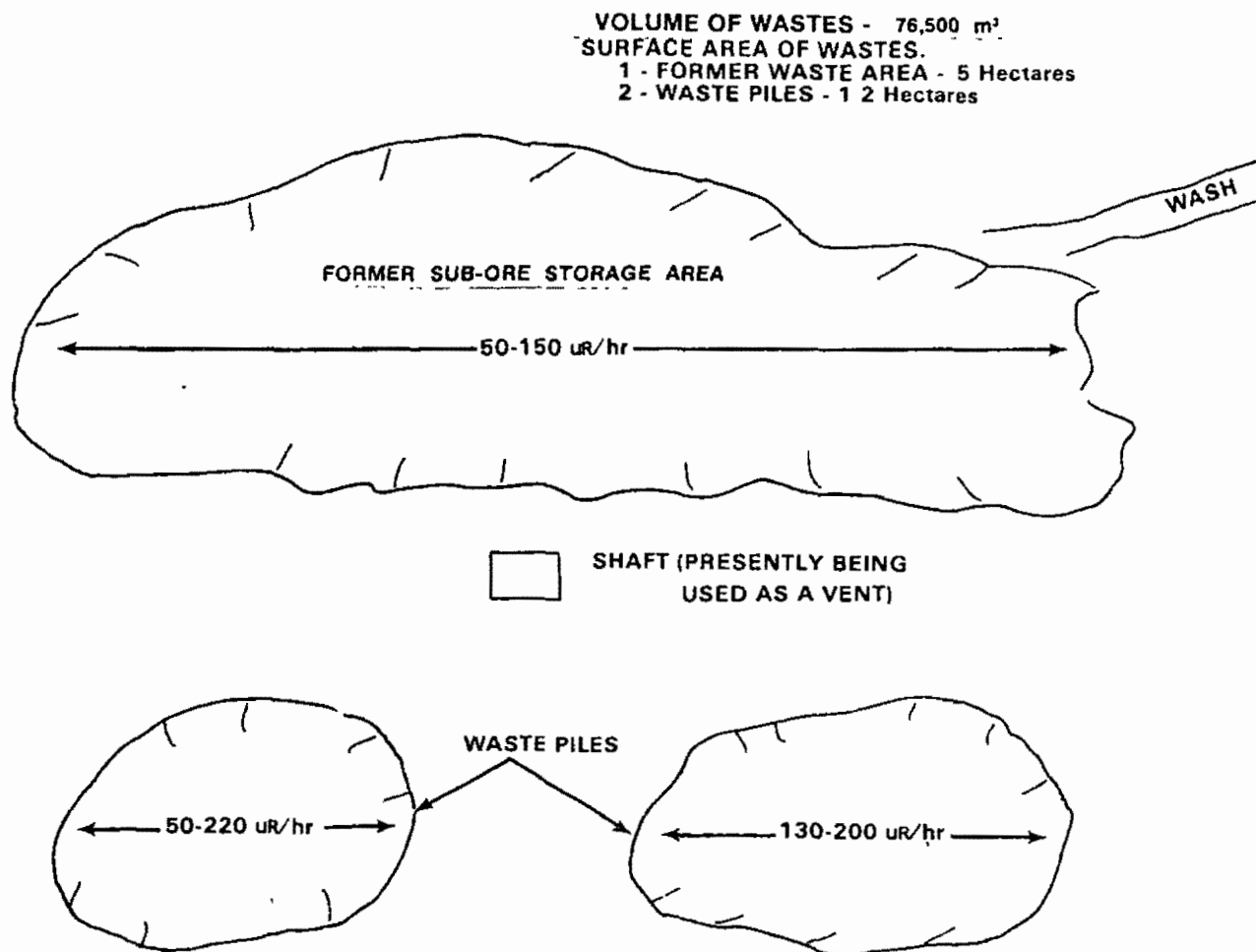


Figure G.5 Plan view of inactive underground uranium mine No. 5, related rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado

Waste Volume = 46,000m³

Surface Area = 0.4 Hectare

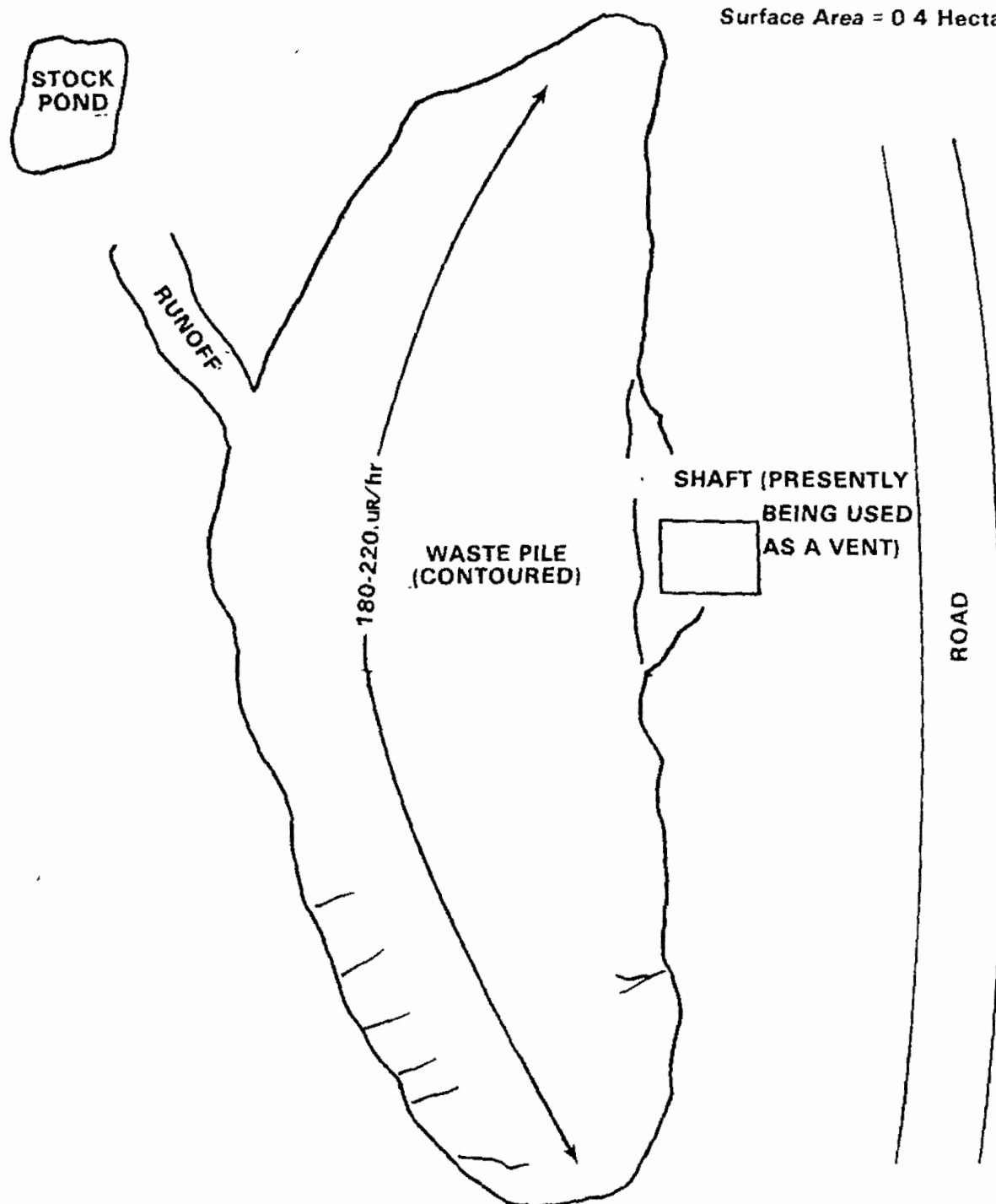


Figure G.6 Plan view of inactive underground uranium mine No. 6, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado

G.1.1.2 Jamestown Area

Uranium ore bodies in the Central City District area generally tend to be small but high in grade. They are mined in conjunction with precious and base-metal ores, particularly gold mining. Pitchblende, associated with all types of veins and shoots, occurs in small pods or lenses systematically arranged in some veins but erratically distributed in others (Si56). Small quantities of pitchblende ore have been shipped from the Central City District since 1872; however, most of the ore mined before 1917 was used as a source of radium. The fluorite ores of the Jamestown District contain small amounts of base metal sulfides and some uranium ore. The quantity of uranium ore was insufficient to be mined for uranium alone (Bu56). The surveys conducted in the Jamestown, Colorado area were made to evaluate some mining areas where uranium was recovered as a by-product.

G.1.1.2.1 Mine 7

This mine was relatively small and produced high grade ore (Fig. G.7). About 38 cubic meters of wastes remain around the shaft, and gamma exposure rates of 400 $\mu\text{R/hr}$ were measured. Erosion of the wastes into the nearby wash was evident. Wind erosion is probably minimal. The mine shaft remains open but filled with water.

G.1.1.2.2 Mine 8

This mine was principally a fluorspar producer; however, uranium ore was also produced and sold (Fig. G.8). The mine shaft remains open to the atmosphere. Mine wastes, adjacent to the shaft, occupy about 800 m^2 , estimated to be about 1,700 cubic meters. Gamma exposure rates on the waste pile ranged from 60-80 $\mu\text{R/hr}$. Extensive water erosion of the wastes has occurred and has produced exposure rates below the waste piles ranging from 40-100 $\mu\text{R/hr}$. Wind erosion of the wastes is probably minimal.

G.1.1.2.3 Mine 9

This mine (Fig. G.9) was located adjacent to the highway just south of Jamestown, Colorado. The mine entry has been covered by a landslide. About 460 cubic meters of wastes, an area of about 400 m^2 , are present on the site. Exposure rates near the entry were about 100 $\mu\text{R/hr}$ and ranged from 40-60 $\mu\text{R/hr}$ near the highway.

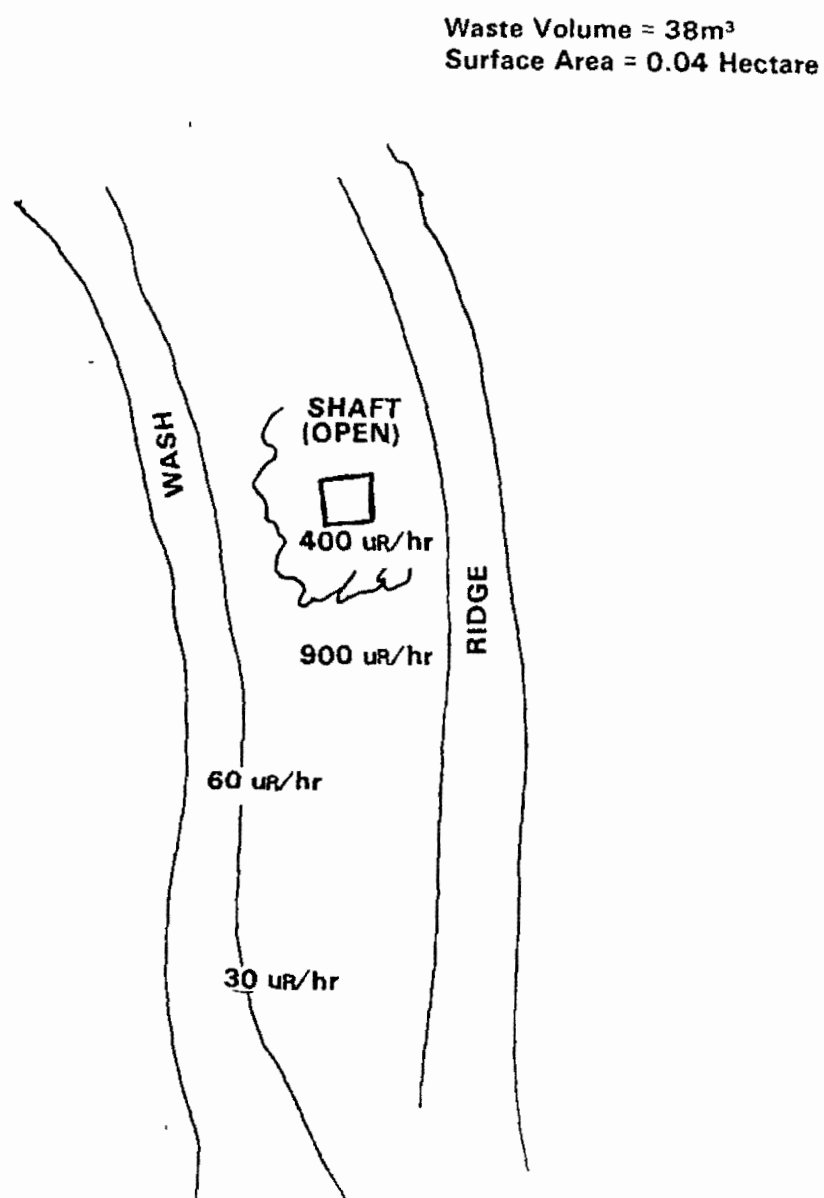


Figure G.7 Plan view of inactive underground uranium mine No. 7, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado

Waste Volume = 1700m³

Surface Area = 0.1 Hectare

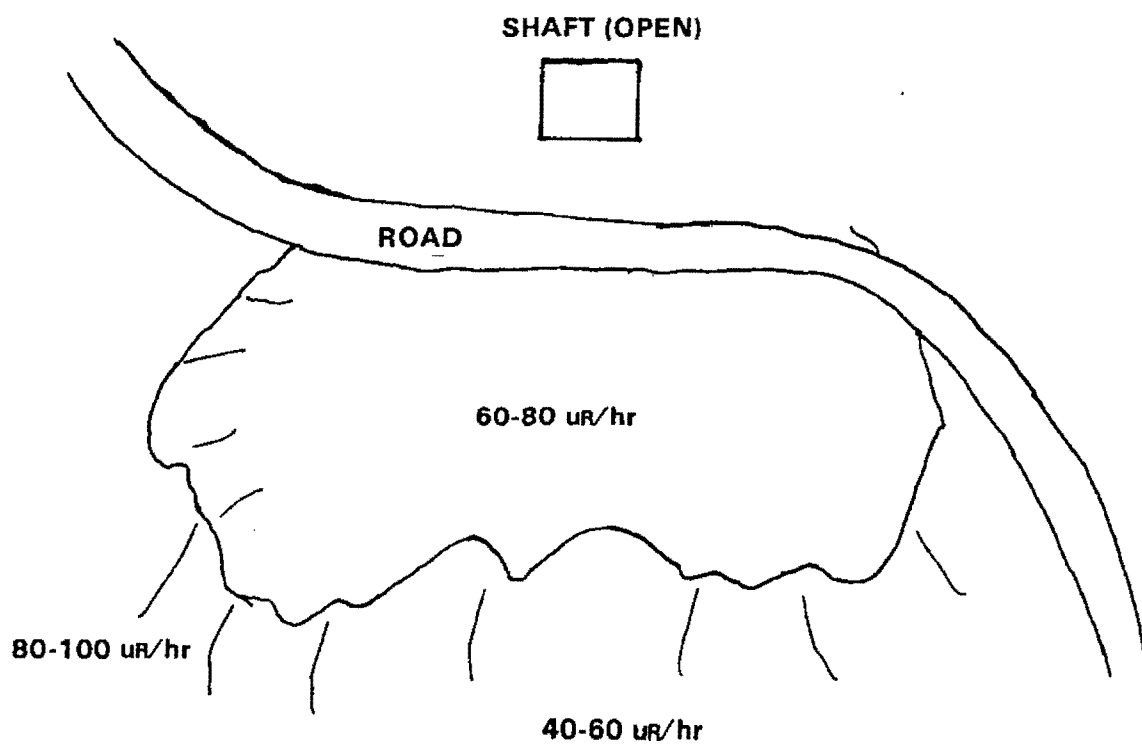


Figure G.8 Plan view of inactive underground uranium mine No. 8, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado

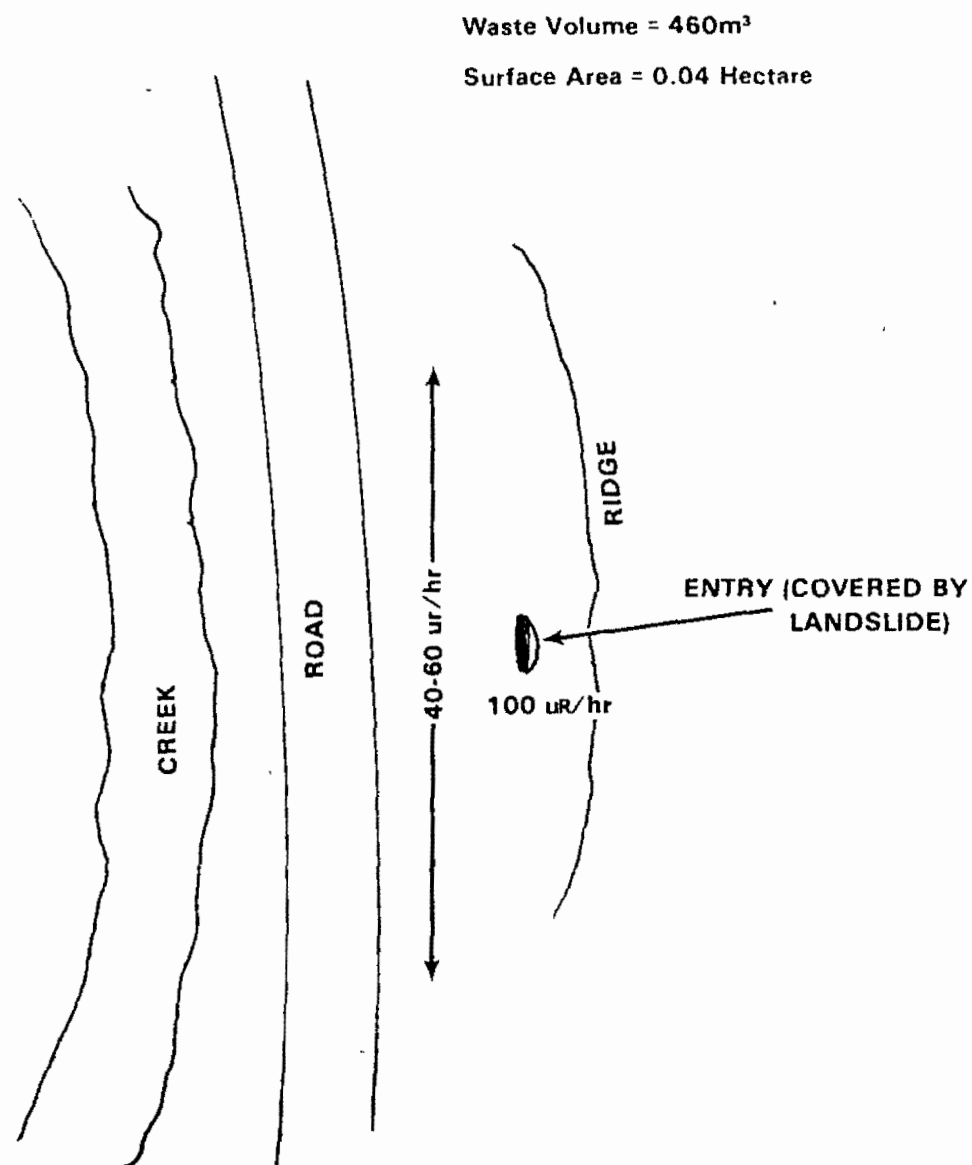


Figure G.9 Plan view of inactive underground flurospar-uranium mine No. 9, related waste rock piles, and surface gamma exposure rates, near Jamestown, Colorado

G.1.1.2.4 Mine 10

This mine was relatively small and the entry remains open (Fig. G.10). Exposure rates near the entry ranged from 100-600 $\mu\text{R/hr}$. Exposure rates on the mine access road were about 70 $\mu\text{R/hr}$. Piles containing mine wastes occupy about 0.1 hectare with a volume of 150 cubic meters. Water and wind erosion of the wastes was evident.

G.1.1.3 Summary

Those mines surveyed in the Uravan area are probably typical of the many inactive uranium mines in that area. Most of the mines are underground and relatively small. Wind and water erosion of the waste piles was evident at all of the mines having entryways open, except where noted. Information derived from these mine surveys is presented in Table G.1 below. Some typical mine waste piles are shown in Fig. G.11. Subsequent photographs depict a typical rim mine (Fig. G.12), an accumulation of wastes on a ledge from a typical rim mine (Fig. G.13), and a mine waste dump from a rim mine (Fig. G.14).

Table G.1 Uravan and Jamestown areas

Mine	Cubic Meters of Wastes	Surface Area of Wastes (Hectares)
<u>Uravan area</u>		
1	13,800	0.1
2	1,200	0.4
3	38,000	2.0
4	6,100	0.4
5	76,500	1.2
6	46,000	0.4
<u>Jamestown area</u>		
7	38	0.04
8	1,700	0.08
9	460	0.04
10	150	0.1

Waste Volume = 150m^3
Surface Area = 0.1 Hectare

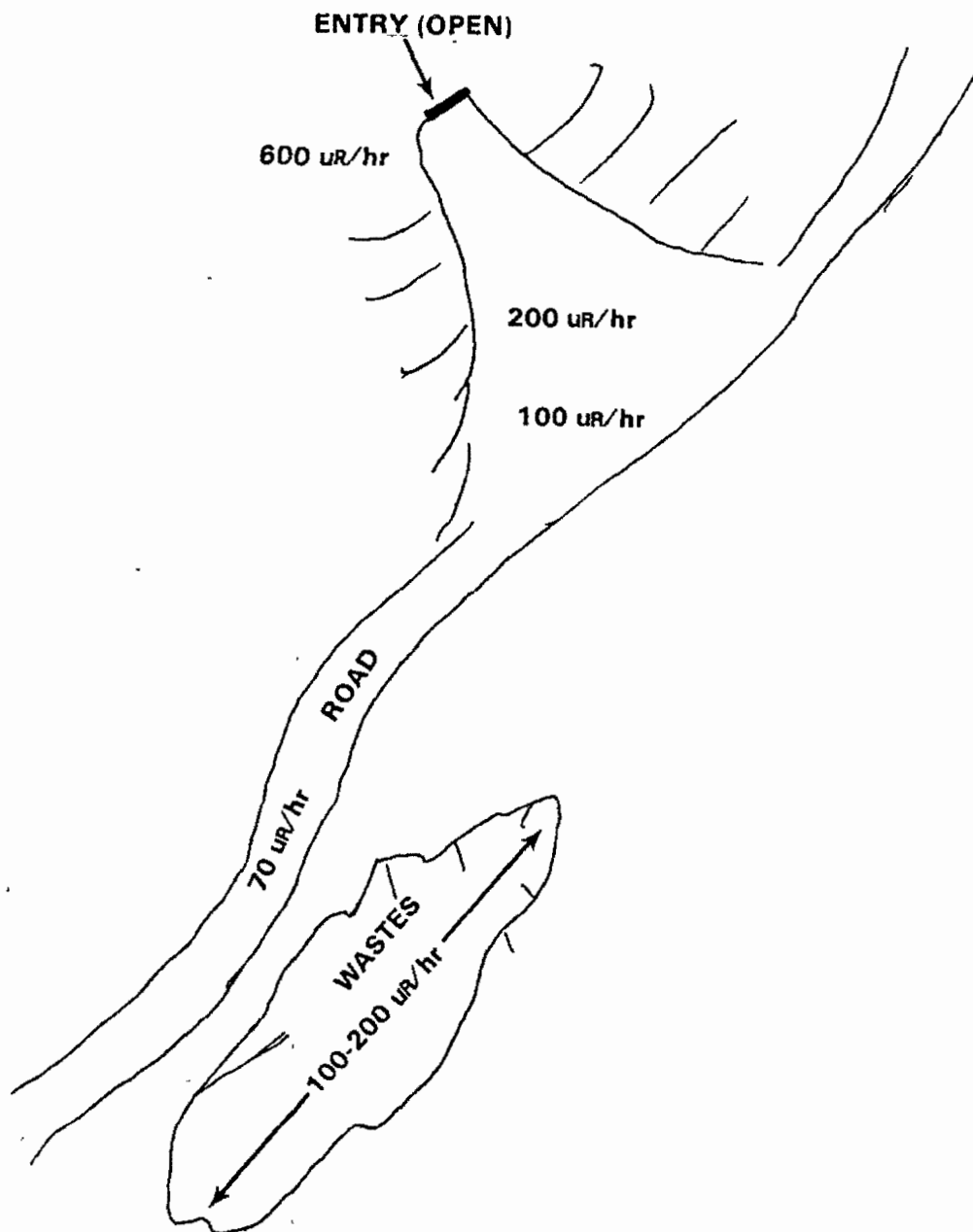


Figure G.10 Plan view of inactive underground uranium mine No. 10, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado



Figure G.11 Typical mine waste pile associated with a small- to medium-sized inactive underground uranium mine in the Uravan Mineral Belt, Colorado

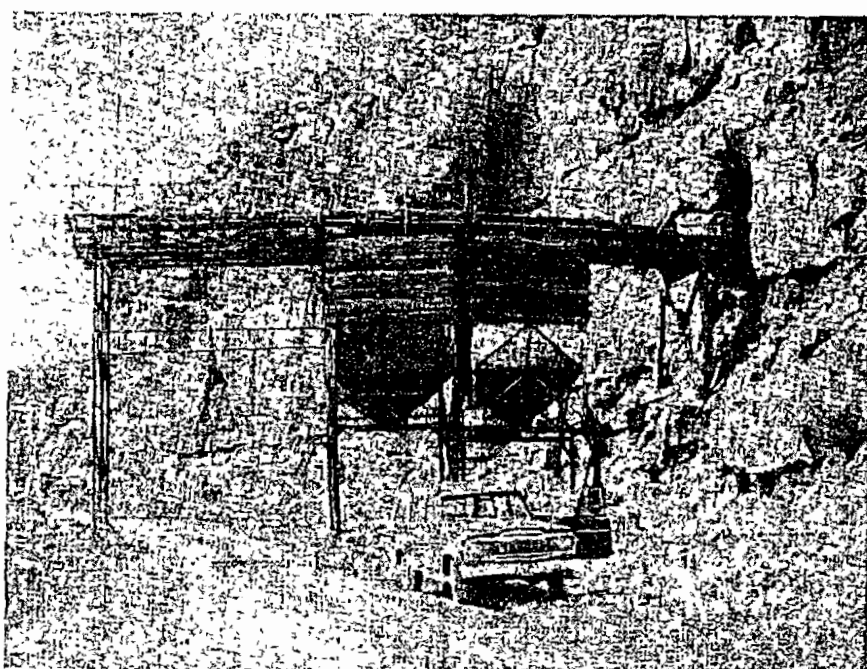


Figure G.12 Side view of a typical underground uranium mine located on the rim of a sandstone mesa in the Uravan Mineral Belt, Colorado

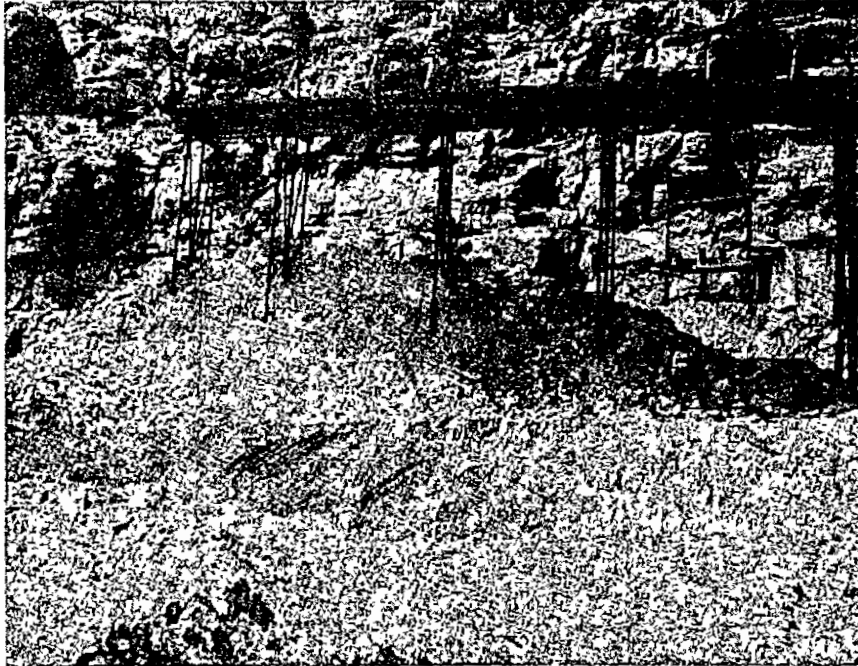


Figure G.13 Mine waste accumulations near the portal of a typical underground rim-type uranium mine in western Colorado

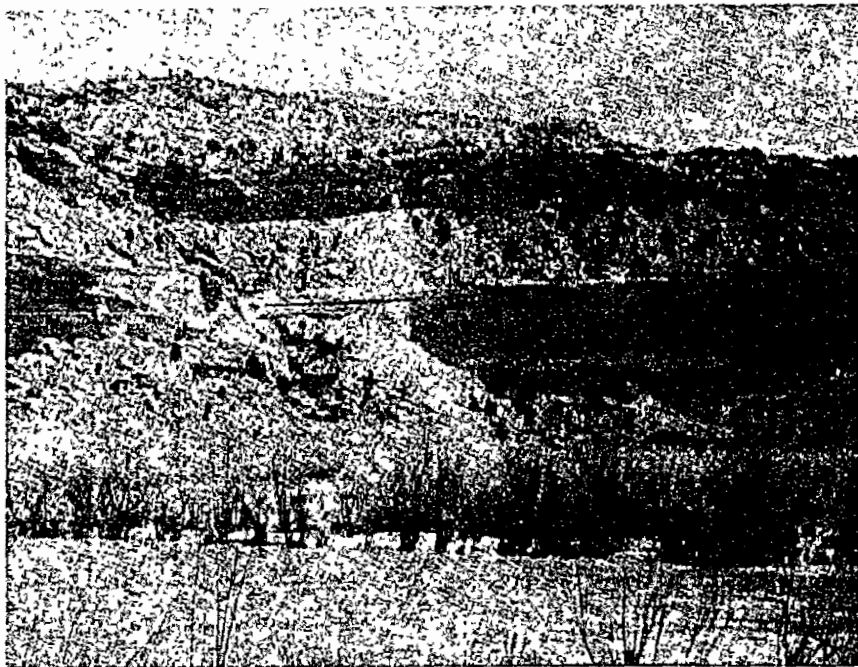


Figure G.14 Mine waste dump associated with a typical rim-type underground uranium mine in western Colorado

Ore-hauling losses have occurred along the mine access and public roads. In addition, mine wastes have been used for road ballast. The ore losses and mine wastes used for road construction probably became airborne with some of the large quantities of dust produced by ore-hauling equipment. The lands in the mining areas generally are used for grazing. Some enhanced uptake of radioactive materials and trace metals by cattle may be occurring. Exploratory drilling is abundant throughout the survey area. Little or no reclamation of abandoned drill sites was observed.

Mines 7 through 10 were surveyed near Jamestown, Colorado. This type of mine was prevalent in this area. Although uranium was not principally produced by many of the mines, the mine wastes generally contain radioactivity. Figure G.15 shows wastes at one such mine entering a stream. Mine wastes producing exposure rates of 40-100 μ R/hr in that area were used as a fill for the Jamestown Park (210 acres). Evidence indicates that some dwellings were built on or near the mine wastes. Surface and groundwater contamination from the mining activities is possible.

G.1.2 New Mexico

Uranium mining operations are continuing to expand production throughout the Grants Mineral Belt region of New Mexico. Underground mining is predominant in the Ambrosia Lake, Churchrock, and Crownpoint areas. Surface mining operations are also expanding at the Jackpile and St. Anthony mines near Paguete/Laguna. However, the majority of inactive uranium mines are located in the area around Grants; therefore, this area was selected for the reconnaissance and field study surveys in New Mexico.

G.1.2.1 Inactive Surface Mines

Two inactive surface mining areas were observed - the Poison Canyon and Zia strip mining areas. Both inactive sites appear to have been more of a shallow strip mining operation compared to the extensive and deeper open pit operations currently underway at the Jackpile and St. Anthony mines. Shallow ore pockets were removed at the Poison Canyon and Zia areas, leaving relatively small pits and waste piles scattered over several hectares. Field studies were completed at several of the Poison Canyon open pits; the radiological data obtained from these surveys are summarized below.

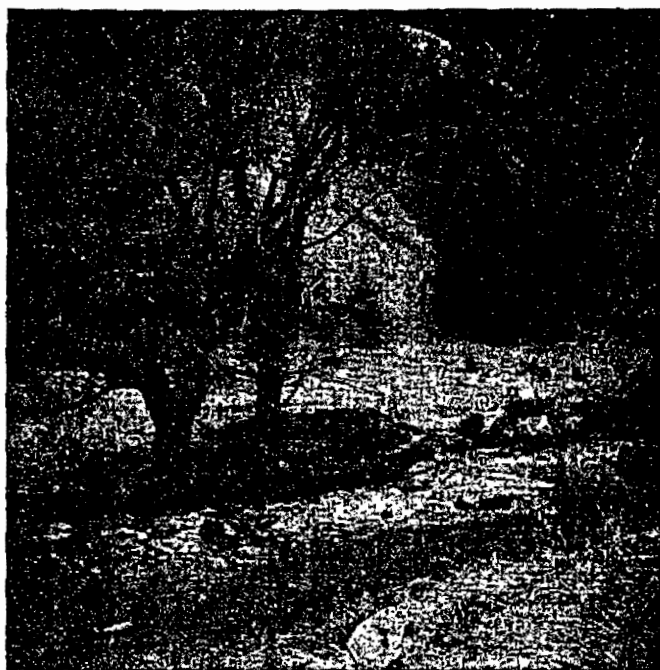


Figure G.15 Movement of fluorspar-uranium mine wastes from a tailings pile into a stream in the Jamestown area of Colorado

G.1.2.2 Inactive Underground Mines

Six underground mines (vertical shaft and incline mining) were observed on Mesa Montanosa: Beacon Hill, Davenport, Dog, Flea, Mesa Top, and Malpais. Several of these mines (e.g., Mesa Top and Malpais mines) were interconnected during their active mining periods. A few of these inactive mines are being used today as fresh air intake points for active mining operations in the area (e.g., mine vents of the inactive Gossett mine are being used in the ventilation system of the active Poison Canyon Mine).

Five underground mines were visited in the Poison Canyon area - Barbara Jane 1 and 3, Westvaco, Santa Fe, and Flat Top mines. Also visited were the Marcus, San Mateo, Anaconda F-33, Hogan, Dakota, and Dysart No. 1 mines. Table G.2 summarizes the reconnaissance and field study survey results.

G.1.2.3 Summary

In summary, the open pit (strip mining) areas of New Mexico have not been restored and numerous shallow open pits or trenches remain with their waste piles undergoing rainwater runoff and windblown contamination of surrounding areas. Most of the inactive underground mine sites have had the head frame and buildings removed and the portals sealed by timber or steel plates to prevent entry, but openings do allow radon exhaust via natural ventilation of the mine. Most mine sites have waste piles which are undergoing rainwater runoff and windblown contamination of surrounding areas. Most of the cased mine vents are not capped to prevent radon exhausting. No mine water drainage was apparent at any of the sites, and most of the mines appear to have collapsed or are flooded.

G.1.3 Texas

Compared to some other western states, the uranium production of south Texas is relatively insignificant, comprising 5 percent of the current United States annual total. However, the relative impact of the mining operations is of interest herein for several reasons: 1) geographic concentration of the actual mine operations, 2) close proximity of the mines to the general population, 3) effect of the high precipitation of the region on the relatively abundant toxic trace elements in the uranium ore and overburden, and 4) minimal land reclamation of some of the older mines which produced high grade ore. The location of uranium ore deposits in south Texas can be readily estimated from the mine locations shown in Fig. 2.4. The deposits

Table G.2 Inactive uranium mine sites surveyed in New Mexico

Mine	Township and Range	Description
Anaconda F-33	T12N R9W Sect. 33 and 34	Two portals sealed by steel doors, large waste piles, no water, runoff apparent, no vents found.
Barbara Jane 1	T13N R19W Sect. 30	No head frame, portal covered by steel plate, but open. Shaft appears to be open, no water, relatively small waste pile, runoff from waste pile, three open mine vents and one capped. Surveys completed at this site.
Barbara Jane 3	T13N R19W Sect. 30	Head frame remains, portal covered by steel plate but openings. Shaft appears to have collapsed. Water drainage from active mine flows through area, small waste piles cleaned off to surface. Several (about six) cased mine vents open to surface.
Beacon Hill	T13N R9W Sect. 20	Small head frame, incline shaft is collapsed. Waste piles in area, no water but runoff apparent. No vents found.
Dakota	T13N R10W Sect.4	Two open portals but mine not deep. Some waste piles, no vents found, no water, runoff apparent.
Davenport	T13N R9W Sect. 20	Open incline but roof collapsed about 60 yards into mine. No vents found. Waste piles in area, no water, runoff apparent. Surveys completed at this site.
Dog Mine	T13N R9W Sect.20	Head frame, incline shaft sealed but appears to have collapsed. No vents found. Large water drainage ditch, large waste piles, runoff apparent.
Dysart 1	T14N R10W Sect. 11	Head frame and buildings, open vents but used in nearby active mine system, waste piles, no water, no runoff.

Table G.2 (continued)

Mine	Township and Range	Description
Flat Top	T13N R9W Sect. 30	No head frame, timber and concrete slab over shaft. No water, waste pile cleaned to surface. No runoff. Several open mine vents found nearby.
Flea Mine	T13N R9W Sect. 20	No head frame, portal sealed with timber. No vents. No water but runoff apparent. Large waste piles.
Hogan	T13N R9W Sect. 14	No head frame, concrete pad covers shaft. No vents found, no water. Waste cleaned to surface, no runoff.
Malpais	T13N R9W Sect. 20	Mine shaft not found but believed to be covered by waste piles. No water but runoff apparent.
Marquez	T13N R9W Sect. 23	Building, portal sealed with timber, several open vents. San Mateo Creek flows through site. Waste piles and water runoff.
Mesa Top	T13N R9W Sect. 20	No head frame, shaft sealed with timber but open, large waste piles. Several open mine vents in area. No water runoff apparent. Surveys completed at this site.
Poison Canyon Strip Mines	T13N R10W Sects. 25, 26 and 139	Open pit or strip mining areas. No shafts or buildings, extensive waste piles and numerous pits, no vents. One abandoned water well, rain water in one pit. Runoff apparent, creek flows between several waste piles in Section 25. Surveys completed at this site.
San Mateo	T13N R8W Sect. 30	No head frame, shaft area collapsed. Large waste piles. Heap leach pile, mine water drainage areas. No water, extensive runoff. One open vent found. Surveys completed at this site.

Table G.2 (continued)

Mine	Township and Range	Description
Santa Fe	T13N R9W Sect. 24	Head frame and building, waste cleaned to surface. No water, no runoff, no vents found.
Westvaco	T13N R10W Sect. 25	No head frame, portal caved in to form hole, no vents found. Small waste pile, no water, runoff apparent.
Zia Strip Mines	T12N R9W Sect. 15	Shallow strip mining areas, no shafts or buildings. Waste piles, no water, no runoff apparent, no vents.

are in tuffaceous, zeolitic sandstone and mudstone beds that strike north eastward and dip gently southeastward (Ea75). Uranium is produced from a three-county area comprised of Karnes, Live Oak, and Duval counties. In each area the host rock is different and ranges in age from Eocene to Pliocene. The Catahoula Tuff is believed by many authors to be the principal source rock for uranium and other elements in the deposits.

Uranium ores currently mined in south Texas are generally of very low grade, the average being about 0.06 percent U_3O_8 . In the recent past many operations, now inactive, were mining ore of the range 0.20-0.25 percent U_3O_8 . The ore zone thickness, although variable, is seldom more than 3.05 m. The usual mining method is by open pit, however, in situ leaching is becoming commonplace and useful under certain conditions. Mine size and geometry are variable, depending on the period of mining activity, the depth of the ore zone, and the proximity (vertically and laterally) of other ore bodies. Many of the mines have a linear trend, paralleling the mineralized roll front. A typical open pit mine would be 30-100 m deep and cover approximately 250,000 m^2 . Currently, a stripping ratio of 35:1 is followed in this area. Any groundwater encountered is diverted to sumps and, from there, pumped to holding ponds.

The reclamation of the pit areas involves contouring the land surface such that all drainage is internal to an on-site holding pond. Topsoil cover is spread about evenly and then seeded with various grasses. It is uneconomical to backfill all of the overburden into the pits and, consequently, some pit remnants consisting of steep walls, etc. are usually left. Generally, reduced agricultural and grazing productivity can be expected in the immediate area of the pit and overburden piles, particularly in the case of older mines.

In situ leaching is common when the depth, size, water content, etc. of the ore body prevents economical open pit mining. It is carefully controlled by the State, especially with respect to monitoring requirements. With cessation of leaching it should be noted that local baseline water quality of the mined aquifer is never fully attained. Potential problem areas are locally increased mobility of trace metals and elevated ammonia levels in the leached ore zone. Tailings from in situ leaching operations can be stored on site or transferred to mill tailings piles.

A compilation of both active and inactive mines by location (county) and type of operation is presented in Table G.3 and Fig. 2.4, both of which are based on State data (Co78).

Table G.3 Status and location of uranium mines in Texas

County	Open Pit ^(a)			In Situ Leaching ^(a)			Total
	A	I	P	A	I	P	
Karnes	8	35	9				52
Karnes-Gonzales	2						2
Karnes-Atascosa	1						1
Gonzales	1						1
Atascosa		1					1
Bee				1			1
Live Oak	6	8		6			20
Duval	1			2		5	8
Webb-Duval				1			1
Webb				1			1
Total	19	44	9	11	0	5	88

^(a) A = active; I = inactive; P = planned.

G.1.3.1 Field Surveys

On May 24-30, 1979, active and inactive open pit uranium mines in Karnes County, Texas were visited in the company of Texas Health Department radiation specialists. The survey included mine wastes and pits in varying stages--active-mining underway, inactive-being reclaimed, inactive-reclaimed, and inactive-abandoned without reclamation. A gamma survey was conducted at one open pit mine that had just been regraded and covered with topsoil but not yet reseeded. The field survey results were supplemented with extensive gamma survey and environmental monitoring results from the Texas Health Department and the Texas Railroad Commission.

Uranium mining in Texas involves considerable and successive description of the land surface as ore bodies are first uncovered and then removed. Tremendous volumes of topsoil, overburden, and water must be relocated a number of times in the course of mining. Dewatering has reached the stage where off-site release is becoming necessary. In the past, mine waters were rarely discharged but were stored in temporary basins on site. Occasionally, mine water was pumped to stock ponds to augment other supplies, typically derived from rainfall runoff. Water quality had to meet accepted standards for stock use.

Figures G.16 and G.17 are of a typical large open pit mine in Texas as of 1972 and 1978. Note the extensive changes in the landscape as unmined land is stripped, mined, and then reclaimed. Much of the overburden is left adjacent to the mines, and standing water remains in most pits. The water originates as groundwater seepage and overland flow from precipitation. The Galen mine, in the right foreground, was abandoned without stabilization about 10 years ago. Natural vegetation is very thin owing to the lack of topsoil and probable toxic effects of trace elements in the wastes. The pile is deeply eroded in places as was shown previously in Figs. 6.10 and 6.11.

Current Texas requirements for stabilization specify that the gamma radiation dose rate must be no more than 0.5 rem per year. For continuous exposure, this corresponds to $57 \mu\text{R/hr}$ above background (about $5 \mu\text{R/hr}$). Results of 21 mine surveys in Texas (Co77) indicated that gamma-ray exposures in excess of $62 \mu\text{R/hr}$ were found at 16 of the mines surveyed. Contributing causes are mineralized overburden (15 of 16), ore pads not properly decontaminated (8 of 16), and mineralized rock in the pit (4 of 16).

G.1.3.2 Summary

In summary, uranium mining has caused radiation levels at some abandoned uranium mines to exceed natural background levels. On approximately one-tenth of the mined area of south Texas, exposure rates could exceed $60 \mu\text{R/hr}$ (the equivalent of 0.5 rem per year for continuous exposure). Although no one is believed to be receiving an exposure in excess of 0.5 rem per year now, the area being mined is increasing, and so is the State's population; hence, the potential for increased population exposures is becoming greater. Individuals occupying a dwelling built on abandoned mine



Figure G.16 1972 aerial photograph of the Galen and Pawelek open pit mines, Karnes County, Texas



Figure G.17 1978 aerial photograph of the Galen and Pawelek open pit mines, Karnes County, Texas

areas could receive excessive lung exposures from radon and its progeny, as well as gamma ray exposures exceeding 0.5 rem per year.

Reclamation by the mining company can reduce radiation levels on mines. One of the most effective methods is to fill in the pit area with the remaining ore, sub-ore, and overburden material and then cover this area with natural dirt or rock of low radioactivity content.

G.1.4 Wyoming

The second largest producer of uranium in the United States is Wyoming. With higher uranium prices, the mining of many low-grade ore deposits would become economical, classifying Wyoming as the largest uranium reserve in the United States. Currently there are many new and expanded operations being planned. Both surface and underground mining methods are used; however, in situ leaching is also underway. Generally, the ore host rocks are arkosic sandstones and conglomerates. Currently, unoxidized ores are being mined, whereas, in the past, shallow oxidized ore bodies were worked. As a result, newer mines are discharging considerably more water to formerly ephemeral streams and, in one case, to a dry lake bed. Within Wyoming, there are 14 major uranium districts, 4 of which are currently producing. These four districts, which are detailed below, exemplify the overall geology of Wyoming uranium occurrences.

G.1.4.1 Highland Flats - Box Creek District

Currently, the largest producing area is the Highland Flats - Box Creek district, located in central Converse County. Host rocks for this deposit are arkosic sandstones of fluvial origin lying within the Fort Union Formation. The ore occurs in roll-type, tabular, and dish-shaped deposits. The largest and most significant of these are the roll-type deposits, varying between 1.5 m and 6.1 m thick. All types are generally associated with each other, occurring from about 46-91 m below the land surface. The ore grade ranges from 0.1 to 0.15 percent U_3O_8 . Former mining operations in this area were in the overlying Wasatch Formation and produced ore of approximately 0.22 percent U_3O_8 .

G.1.4.2 Crooks Gap District

The second largest producer of uranium, the Crooks Gap district, is located in the Green Mountains of Fremont County. Operations began there in 1954. The host rocks are arkosic sandstones in the Battle Spring Formation. The ore bodies are of tabular, stratiform, and roll-type occurrence modes and are concentrated in narrow zones at the margins of the altered sandstone. Those currently mined are at or below the water table and are unoxidized. Ore grade ranges from 0.18 to 0.23 percent U_3O_8 . Formerly, smaller, near-surface ore bodies were mined.

G.1.4.3 Gas Hills District

The Gas Hills district has produced the most uranium in Wyoming and has the largest number of mine and mill operations. Large-scale continuous production has occurred since 1960. It is located in eastern Fremont County. The host rocks are arkosic sandstones in the Wind River Formation. Within this region there are four types of deposits, the roll-type being the most important. These are found at depths of about 30 m to 150 m below the surface and up to 122 m below the water table. The ore zones are 0.3 m to 3.1 m thick, occasionally ranging from 6 m to 10 m thick. The current ore grade is approximately 0.1 to 0.15 percent U_3O_8 . Also within the district are small, high-grade residual deposits behind the main solution front deposits.

G.1.4.4 Shirley Basin District

The Shirley Basin district in northwest Carbon County has been actively mined since 1960 and is expected to expand considerably. The host rocks are arkosic sandstones within the Wind River Formation and the deposits are of the roll-type. Found at the leading edge of the tongue of the roll-front, the ore bodies tend to be large, about 15 m wide by 760 m long. Smaller ore bodies are found along the top and bottom of the roll-front. Overall, the ore bodies vary from a few hundred to a few thousand MT, at depths from 45 m to 90 m below the surface. Main ore bodies lie below the groundwater table, sometimes to depths of 90 m. The ore grade ranges from 0.2 to 0.6 percent U_3O_8 .

G.1.4.5 Summary

Approximately 90 percent of the ore produced comes from surface mining operations. Overburden thickness ranges from 30.5 m to 137 m below ground surface. Spoils bank accumulation is on the order of $764 \times 10^3 \text{ m}^3/\text{month}$ for an average lifetime of 15 years. This material is stockpiled for later reclamation pursuant to the State Environmental Quality Act of 1973; however, to date, land reclamation has not begun at any surface mine site. It is evident that the mine waste volume from an underground mining operation is much less than that generated by an open pit mine; therefore, it is estimated that about two hectares per portal would be sufficient to dispose of the waste rock.

The majority of the inactive mines are located in the Gas Hills and Shirley Basin mining districts, located in Fremont and Carbon Counties, respectively (Personal communication with UNC staff 1979). There are some inactive but not abandoned mines at every production area. Any increase in the value of U_3O_8 will lead to the reopening of many mines. Therefore, it is difficult to select an inactive mine that would be considered abandoned (mined out).

The Morton Ranch leasehold, described as typical of smaller inactive and perhaps abandoned operations, was the site of the radiological survey. The topographical and climatological parameters of the area are similar in practically any portion of central Wyoming. Precipitation ranges from 30 cm to 36 cm per year, with June being the wettest month, November the driest. The wind blows constantly at variable frequencies up to 129 km/hr (NUREG 0438). The topography is dominated by plains, low-lying hills, and table lands interrupted by stream channelways.

In 1973, an inactive pit, 1601, was very briefly mined to determine the metallurgical qualities of the underlying ore. A pit 110 m x 238 m x 12.2 m deep remains. Adjacent to the pit, piles containing $237,000 \text{ m}^3$ [$73,000 \text{ m}^3$ of ore and $164,000 \text{ m}^3$ of spoil (sub-ore)] of spoil material occupies less than 2.5 hectares.

Results of the gamma survey performed along radials originating from the center of the pit appear in Fig. G.18. The near-surface ore body complicated field results from the survey; therefore, soil samples were taken at every 366 m to determine the presence of wind-blown material or surface ore outcropping. Additional soil samples, 75 cm profiles, were taken at erosional occurrences and in drainages.

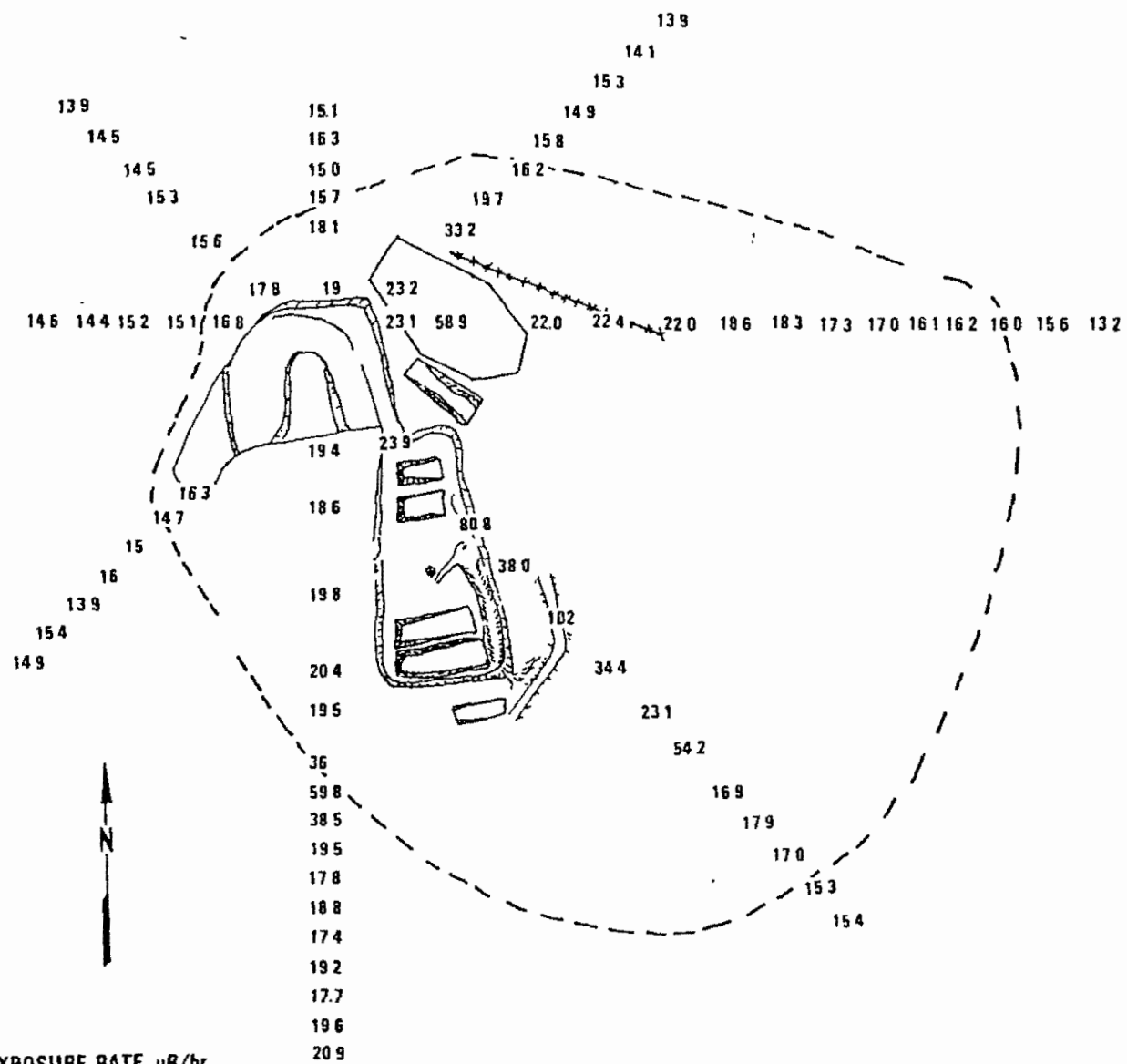


Figure G.18 Results of gamma exposure rate survey at the 1601 pit and environs, Morton Ranch uranium mine, Converse County, Wyoming

G.2 Aqueous Transport of Mining Wastes in New Mexico and Wyoming

G.2.1 Description of Field Studies

Field studies were conducted to investigate the transport of trace elements and radionuclides from inactive mining areas to off-site locations in New Mexico and Wyoming. These areas were selected because extensive uranium mining has occurred to date and is likely to continue. Since the mid 1950's, these States have produced the majority of domestic U_3O_8 .

Samples of surface soils, stream sediments, mine drainage water, and surface water were collected. Interpretation of the data is complex since wind and water erosion work together at different seasons of the year to transport the mine material stored above ground. Compounding the problem is the semiarid environment of Wyoming and New Mexico where precipitation averages 13 to 31 centimeters per year and occurs primarily in the spring. The short-duration "flash flood" summer thundershower will move large quantities of material in increments rather than a gradual erosional pattern.

Sampling was conducted during April and May 1979 at a site within the South Powder River Basin in Wyoming, at two sites in Poison Canyon drainage, and at the San Mateo mine areas in New Mexico. Most of the soil samples were obtained in well-defined runoff gullies where mine and mine spoil drainage intersected stream beds. The arid nature of the locations did not provide much opportunity to observe and to measure surface runoff characteristics.

Sampling locations at the Morton Ranch property in Wyoming are summarized in Fig. G.19. The types of samples are identified in the legend. Samples taken at the San Mateo mine in New Mexico are depicted in Fig. G.20.

At three locations at Morton Ranch, a 75 cm profile was taken which consisted of 15 consecutive 5 cm segments. The potential variability of trace metals and radionuclides at depth may be related to the solubility of the species and the amount of surface water residence time. Understanding the fractionation of surface contaminants in the soil column is important in evaluating the transport of various species by redissolution or leaching.

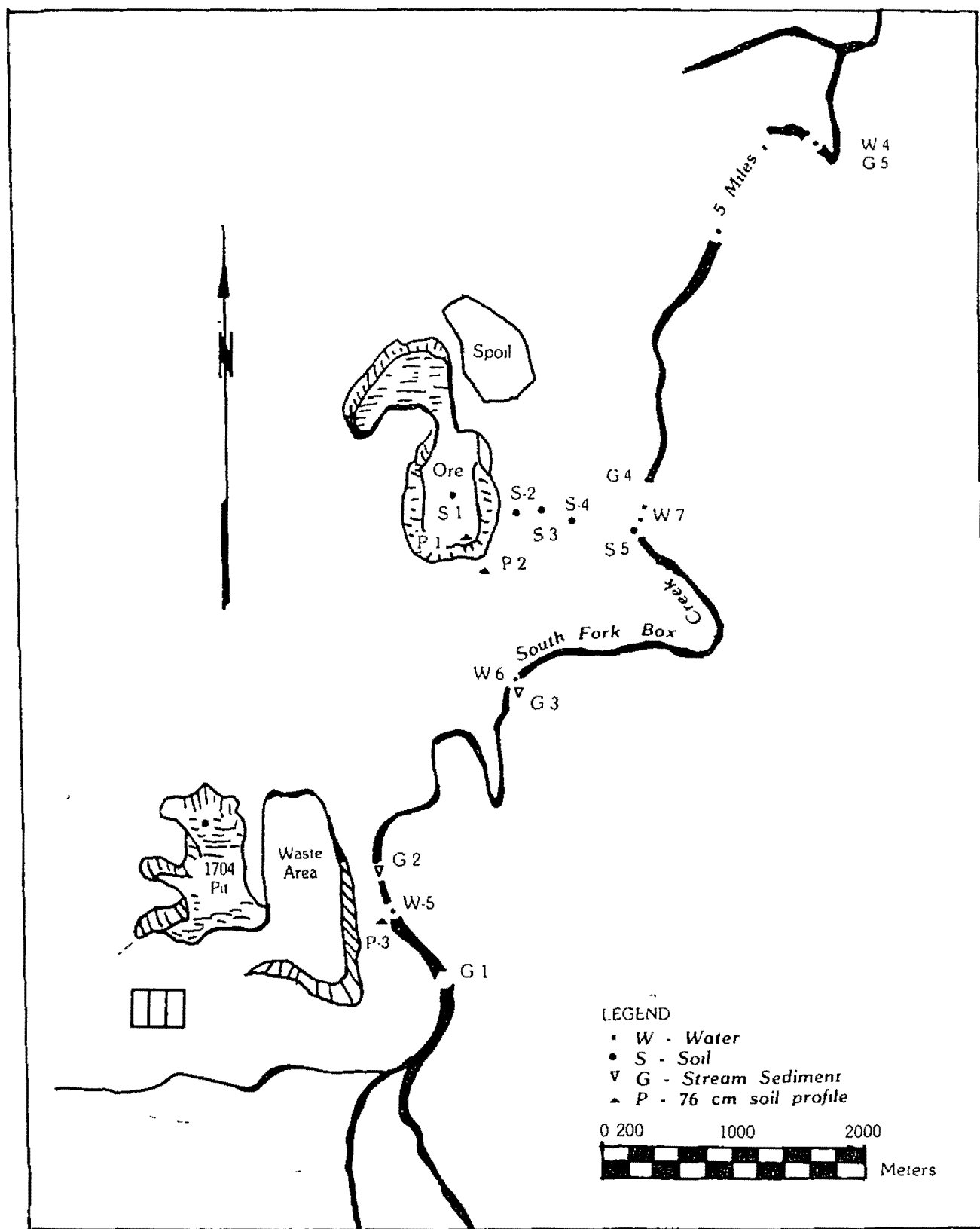


Figure G 19 Location of sampling sites at the Morton Ranch mine South Powder River Basin Wyoming

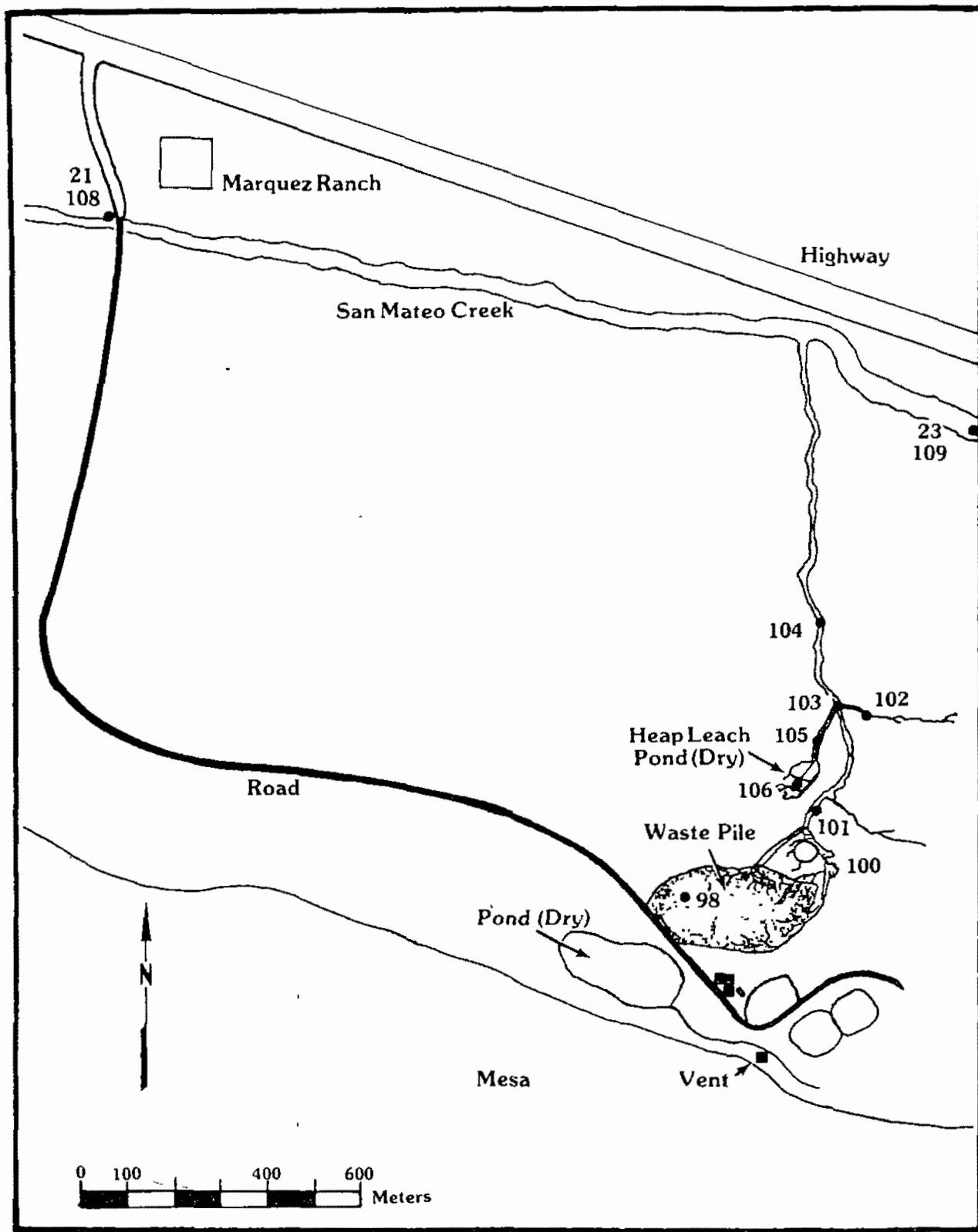


Figure G.20 Sample locations for radionuclides and select trace metals in sediments, San Mateo Mine, New Mexico

G.2.2 Discussion of Results

The radiochemical and trace metal analyses for the stations shown in Fig. G.19 are contained in Tables G.4, G.5, G.6, and G.7 (Written Communication from N.A. Wogman, Battelle Pacific Northwest Laboratory, 1979).

Table G.4 Trace elements and radionuclides in water in the South Fork of Box Creek drainage at UNC Morton Ranch lease

Location ^(a)	pCi/g		μg/l			
	Ra-226	U-238	Ba	Se	V	Mn
W-3 H ₂ O	15.5	1220	63	334	8	77
Filter	18.7	33.7	22	38	47	22
W-5 H ₂ O	4.2	287	76	78	5	9
Filter	4.48	2.97	16	0.6	8.1	29
W-6 H ₂ O	0.6	34.2	11	<4	10	91
Filter	0.091	0.209	<4	<.1	<.6	11
W-7 H ₂ O	0.5	38.7	86	<4	7	22
Filter	0.18	0.976	4	<.1	1.0	18
W-4 H ₂ O	0.13	10.9	44	<4	6	149
Filter	0.045	0.188	<3	<.1	1.7	6.7
Acid Blank H ₂ O	N.D.	N.D.	1	<4	<5	<3
Filter	0.59	0.073	<3	<.12	<.4	<.2

(a) See Figure G.19.

Note. -- United Nuclear Corporation (UNC) has recently transferred its interest to the Tennessee Valley Authority.

Table G.5 Radionuclides and trace metals in sediments in the South Fork of Box Creek at UNC Morton Ranch lease

Location ^(a)	pCi/g		μg/g				
	Ra-226	U-238	Ba	Se	V	Mn	As
G-1	1.3	0.41	750	<1	<39	70	< 1.6
G-2	1.8	1.04	690	<1	<42	114	2.9
G-3	6.8	7.46	550	7	90	3390	10
G-4	1.4	1.45	750	<1	<51	100	<1.7
G-5	1.2	0.87	720	<1	<50	107	3.4

^(a) See Figure G.19.

The results in Tables G.4 through G.7 may indicate dispersal patterns of mine drainage and waste in a semiarid environment. Intermittent runoff from surface or underground mining spoils or an irregular low-volume discharge of mine water compounds the difficulty in attributing the impacts to either suspended or dissolved species. The analyses performed on samples from the South Fork of Box Creek cannot be used to describe typical impacts from larger operations; yet the complexity of stream sedimentation and dilution is evident and can signal precautions in monitoring any site of mining operation.

For instance, water samples collected from Box Creek represent an intermittently flowing stream without a constant mine discharge. Even in spring after most of the snow has melted and recharged the upper reaches, streams such as Box Creek do not flow but rather contain shallow impoundments. There seems to be a movement of water beneath the stream sediments which supports these impoundments without observable surface flow. Each of the water and sediment samples was from impoundments, except for sediment sample G-1 which was from a dry stream bed. The results in Table G.4 show that radium-226 and uranium-238 in solution decrease in concentration with downstream distance. Station W-3 is hydrologically separate from the drainage basin at the present time, but represents mine discharge when dewatering begins.

Table G.6 Radionuclides and trace metals in soils near the 1601 open pit mine, UNC Morton Ranch lease, Wyoming

Location ^(a)	pCi/g			μg/g					
	Ra-226	U-238	Th-230	Ba	Mn	V	Zn	Se	As
S-1	119	188	161	760	205	110	29	17	7.5
S-2	33	7.21	38.6	800	250	160	85	25	11
S-3	3.8	1.87	3.71	760	200	70	36	<1	6.8
S-4	1.9	1.78	3.54	530	190	100	58	<1	6.0
S-5	2.5	1.64	1.91	810	140	<50	27	<1	4.2
S-6 ^(b)	1.86	1.62	1.74	680	180	100	33	<1	3.9
S-7	3.4	1.60	2.71	570	180	70	32	<1	5.6

(a) See Figure G.19.

(b) Background sample.

In conjunction with the decreasing soluble species with distance downstream, there is a possible trend of increased levels in sediments (see Table G.5). This may be, in part, due to the sampling locations (ponded water). It is possible that the soluble species precipitate in standing water either because of evaporation increasing the solute concentration or the increase in pH. The pH of the 1704 mine water increased from 5.2 to 7.2 eight km downstream in Box Creek. Another possibility is that these ponds receive surface runoff from several hundred meters of stream bed and, therefore, have higher sediment loading.

The data in Table G.6 are from a series of soil samples obtained from the natural drainage leading from the inactive 1601 open pit to the South Fork of Box Creek. This mine is dry and therefore it has no liquid discharge to Box Creek. Over 1,100 m separate the mine spoils material from the creek bank. The soil samples were collected at 183, 366, 567, and 1,100 m from the spoils bank. If there had been runoff from cloudbursts or snow melt, sheet erosion across this plane could move the spoils material to the creek. This mine was operated for a very short time eight years ago; therefore, the full

impact of the long-term erosion could not be measured. Also, wind erosion complicates the predictive value of these results since this channel is aligned with the predominant wind direction.

Table G.7 Radionuclides and trace metals in soil profiles at the open pit mines, UNC Morton Ranch lease, Wyoming

Location ^(a)	Depth (cm)	pCi/g			μg/g			
		U-238	Ra-226	Th-230	Mn	Se	Ba	V
P-1	0-2	188	119	161	205	17	760	110
	4-6	1.62	2.8	1.79	120	1.2	720	< 50
	10-12	2.12	1.8	1.87	59	< 1	470	150
	22-24	1.21	1.4	1.0	86	1.9	550	130
	28-30	1.26	1.5	1.16	57	< 1	590	70
P-2	0-2	11.0	33.2	31.4	130	2.7	690	90
	4-6	5.39	13.0	18.0	94	1.1	760	< 50
	10-12	1.57	2.6	2.15	130	< 1	770	60
	22-24	11.0	146	238	190	1.3	560	120
	28-30	5.73	485	481	72	< 1	730	90
P-3	0-2	3.3	8.2	5.5	120	1.3	701	130
	4-6	3.0	52.0	11.1	110	1.1	693	70
	10-12	2.69	1.86	1.35	210	1.4	591	130
	22-24	1.97	1.62	1.78	270	1.1	618	90
	28-30	3.51	2.1	1.5	350	1.1	660	70

(a) See Figure G.19

The trend of decreasing radioactive species with increasing distance from the spoils bank is evident. The background results for S-6 are composite samples of at least six locations where the gamma survey indicated no surface contamination. The results show that Ra-226, U-238, and Th-230 are measurable and decreasing towards the creek. These concentrations above

the background samples can be attributed either to wind or water erosion from the 1601 spoils bank.

Water erosion of the mine waste pile is documented by the gullying scars on the sloped surfaces. Data in Table G.6 indicate definite migration of this material at least 360 m from the largest gully. Yet it is possible that this contamination may have occurred while the mine was active. The ore body is sandstone, the fines of which could be resuspended by vehicle traffic or equipment operation.

In Fig. G.20 the drainage pattern for the San Mateo mine in New Mexico is depicted. As in the case of the 1601 mine in Wyoming, nearby drainage courses are dry most of the year. Runoff in San Mateo Creek lasts for several months as a result of snow melt, and is nil the rest of the year, except for brief runoff from storms. Radium-226 data for the sediments (see Table G.8) reflect decreasing concentration with distance to San Mateo Creek, especially within 350 to 460 m from the waste pile. The radium in San Mateo Creek downstream from the intersection of the mine drainage is higher than the gully data would indicate. Data for sample 104 seem to indicate that contamination has not moved from the gully wash to the creek beds, but this is considered most unlikely on the basis of known erosion and obvious topographic relations.

Barium and selenium trends, for the most part, follow a similar pattern of decreasing concentration with distance from the waste pile. Arsenic concentrations are an exception. Sediments show increasing arsenic to 350 to 460 m downstream and then decrease from that point to San Mateo Creek. In the creek sediments, arsenic and barium concentrations are higher than would be expected if mine wastes were the sole source of these elements.

Sediment and water samples (Tables G.8 and G.9, respectively) from San Mateo Creek indicate that barium and manganese concentrations are either equal to or higher upstream of the gulley intersection than below it. This apparent anomaly and the higher-than-expected concentrations for radium and arsenic in sediments cannot be readily explained.

In summary, the data collected during April and May 1979 at the San Mateo mine do not indicate that mine waste has reached San Mateo Creek; yet downstream sediments show anomalies for certain elements. The transport of mine waste is measurable approximately 370 to 460 meters from the waste pile.

Water erosion is the likely reason for the contamination, but wind may also be dispersing material. Meteorological data, particularly wind roses, are unavailable at this time.

Table G.8 Radionuclides and trace metals in sediments from the drainage of the San Mateo mine and from San Mateo Creek, New Mexico

Location (a)	pCi/g		$\mu\text{g/g}$					Remarks
	Ra-226	Th-232	Ba	Se	V	Mn	As	
98	117	0.86	-	-	-	-	-	Waste pile
100	55	0.64	566	3.9	78	176	3.1	Base of pile
101	36	0.66	484	3.7	114	179	5.1	100 m from pile
103	1.6	0.43	383	1.2	<50	191	6.2	400-500 m from pile
104	0.77	0.54	434	<1	<42	146	3.6	600-700 m from pile
105	1.2	0.80	517	<1	<52	186	3.8	Heap leach pile
102	0.77	0.55	562	1.2	102	473	5.5	Background
109	0.38	0.39	695	<1	<51	152	2.2	2 km upstream
108	8.1	0.53	597	<1	55	157	5.2	Downstream

(a) See Figure G.20.

Table G.9 Radium-226 and trace elements in water from San Mateo Creek near San Mateo mine discharge point

Location (a)	pCi/l	$\mu\text{g/l}$						
	Ra-226	Ba	Se	V	Mn	As	Mo	Zn
23 (upstream)	---	79	< 4	8	55	9	23	56
21 (downstream)	12.5	26	21	21	10	9	170	150

(a) See Figure G.20.

Data from three soil profiles near the spoils areas of the 1601 and 1704 mines at Morton Ranch were used to investigate possible downward migration of soluble species. In semiarid environments, small precipitation events usually result in little noticeable runoff. In many areas, the surface soil is very porous and rain immediately infiltrates. The disturbed spoils material is probably of even higher porosity; therefore, soluble species could, over a period of time, begin to migrate to greater depths. Based upon solubility and frequency of rewetting, a fractionation of species could occur.

The overburden at 1601 and 1704 mines contains a high percentage of clay. The resultant spoils bank is a homogeneous mixture of clay and sandstone. It was observed during the collection of the soil samples that the clay was moist, highly plastic, and obviously of low permeability. Very slow downward migration of surface water would be expected; consequently, not much fractionation of the species would result.

The results in Table G.7, however, show contradictory evidence of isotopic disequilibrium. The isotopes of uranium, radium, and thorium are near equilibrium conditions throughout the profile P-1 taken above the slope of the 1601 spoils bank. However, the lower layers of profile P-2 show a marked departure of uranium from the radium and thorium. Since uranium is more soluble than the other species, it is difficult to explain the anomaly.

In profile number 1, the top segment probably contains ore material eroded to the spoil surface. The segments at greater depth show species equal to or less than the data reported as background on the surface in Table G.6 (S-6). This would imply that no downward migration had occurred on the surface of the spoils area. Note that the clay-like matrix of the spoils may have prevented much surface infiltration, which accounts for the shallow erosional scars on horizontal surfaces.

Profile number 2 was obtained in an alluvial fan below the spoil bank slope. The data suggest that the profile did not extend below the alluvium and into the spoil material. The anomaly at greater depth cannot be explained, but, perhaps, irregular wind and water erosion of the spoils material caused this layering effect. Still, the isotopic disequilibrium is not explained.

Profile number 3 was a saturated clay column which contained, from the odor, a percentage of organic material that was undergoing bacterial decomposition. Sheep in the area use the pond nearby and this may account for the source of the organics. The results for profile 3, again, show little above background concentrations at greater depth, except for slight increases in manganese, selenium, and uranium. The profile was in the sediments of Box Creek close to the runoff observed from the 1704 spoils bank. The radium and uranium data show that concentrations in sediments in the first 30 cm are higher than what was observed further downstream at point G-2 (Table G.5). This is undoubtedly the result of erosion from the spoils bank. This clay sediment matrix could be subject to transport further down Box Creek but only under high flow conditions.

G.2.3 Conclusions

In summary, the field studies attempted to identify and quantify the transport of mine spoils material. There is evidence that contamination is measurable up to 370 to 460 m in a gully draining the San Mateo pile and, perhaps, 570 m in the natural sloping plane in which the 1601 Morton Ranch mine waste is located. The mechanism of transport is complex because of the semiarid environments. There is a good chance of wind and water erosion combining to move spoils material these distances in under 10 years.

The data also support the conclusion that it is not evident that radio-nuclides or trace metal species have reached intermittent streams, either as soluble or particulate material. Anomalies were uncovered in San Mateo creek which cannot be explained from the samples collected so far. The water and sediment samples in the South Fork of Box Creek show decreasing concentrations of radium and uranium isotopes with downstream distance, but not so clearly for the trace metals. The levels of radium and uranium are not conclusive of major off-site movement of the spoils material.

The soil profiles obtained at the Morton Ranch leasehold do not support the conjecture of downward migration. Perhaps the clay-like consistency of the spoils material allows for too little fractionation after only eight years. The isotopic_ratio discrepancy at greater depth in the profile of an alluvial fan of spoils is not understood at this time.

G.3 References

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APPENDIX H

INFLUENCE OF MINE DRAINAGE ON SEEPAGE TO GROUNDWATER AND SURFACE WATER OUTFLOW

H.1.0 Influence of Mine Drainage on Seepage to Groundwater and Surface Water Outflow

H.1.1. Surface Mines

Mine water discharge from each open pit mine in the Wyoming model area is estimated to be $3.00 \text{ m}^3/\text{min}$. Discharge is to natural ephemeral streams which will, in some cases, become perennial as a result of the discharge. Effects of mine drainage on water quality, stream biota, and downstream potable water uses are largely a result of dilution of suspended and dissolved load. The first step in the analysis is to determine what the net outflow of water from the sub-basin will be and then to route this water downstream where contact with man or the food chain is possible. It is assumed that suspended and dissolved chemical loads not leaving the sub-basin because of infiltration and evaporation remain on the stream bed and are available for subsequent transport in flood flows, which are calculated in Section 3.3.3.

Numerous assumptions are made in the analysis to follow, which is patterned after that in NUREG-0511 (Generic Environmental Impact Statement on Uranium Milling), with certain corrections and modifications. The calculations are based on the operation of three surface mines dewatering at a rate of $9.00 \text{ m}^3/\text{min}$ (3 mines, $3.00 \text{ m}^3/\text{min}$ each) into a sub-basin of 11.4 km^2 area. This will transform 22.7 kilometers of channel (in the sub-basin and basin) into perennial streams. Note that $7.04 \text{ m}^3/\text{min}$ will discharge from the sub-basin. The remainder of the flow ($1.96 \text{ m}^3/\text{min}$) is lost by infiltration and seepage in the sub-basin.

Mine discharge is assumed to enter into a hydrographic area described as follows (Table H.1) and shown in Fig. H.1:

Table H.1 Characteristics of the sub-basin containing the model mines

	Symbol	Value	Units
Slope of region		0.01	%
Substrate hydraulic conductivity (vertical)	K_v	6×10^{-6}	m/min
Substrate hydraulic conductivity (horizontal)	K_h	6×10^{-4}	m/min
Composite hydraulic conductivity	K	6×10^{-5}	m/min
Total outflow of a stream section	Q_T	Calculated	m^3/min
Total loss due to infiltration (seepage) and evaporation	Q_L	Calculated	m^3/min
Seepage loss	Q_s	Calculated	m^3/min
Evaporative loss	Q_e	Calculated	m^3/min
Annual evaporation rate	E	2×10^{-6}	m/min
Length of stream section	L	Variable	m
Reach of stream section with perennial flow	R	Variable	m
Channel dimensions (see Figure H.2)	a, b	Variable	m
Cross sectional area of channel calculated from Q_{in} and V	A	Variable	m^2
Water input from mine drainage	Q_{in}	3.00 per mine	m^3/min
Wetted perimeter of stream bed	B	Variable	m
Velocity of flow	V	36	m/min

$$\begin{aligned}
 K &= (K_h \cdot K_v)^{1/2} \\
 &= (6 \times 10^{-6} \times 6 \times 10^{-4})^{1/2} \\
 &= 6 \times 10^{-5} \text{ m/min}
 \end{aligned}$$

$$Q_L = (Q_s + Q_e)$$

$$B = a + 2b$$

$$Q_s = KB \times f(L) \quad \text{Assuming that } B \text{ and } a \text{ are dependent only on } Q_{in},$$

$$Q_e = Ea \times f(L) \quad V \text{ is assumed constant at } 36 \text{ m/min, and } f \text{ is a}$$

$$V = 36 \text{ m/min} \quad \text{function of } L.$$

$$A = \frac{Q_{in}}{V}$$

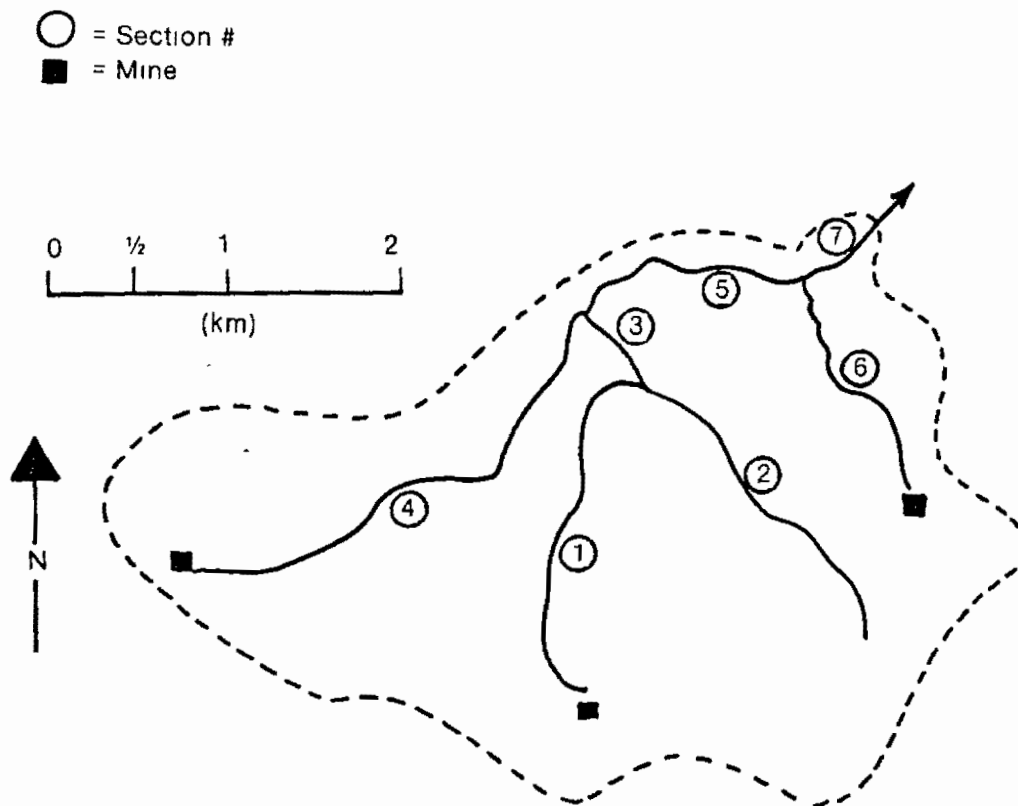


Figure H.1 Wyoming model area sub-basin drainage system

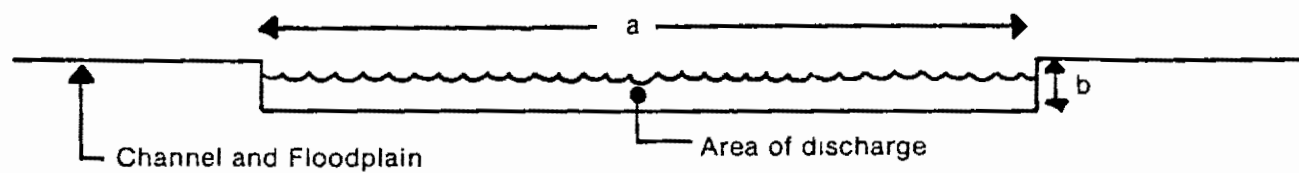


Figure H.2 Model area stream cross section

To find the total outflow (Q_T) from a stream section of length L :

$$\frac{dQ_T}{dL} = -KB - Ea$$

$$\int dQ_T = -(KB + Ea) \int dL$$

$$Q_T = -(KB + Ea)L + C$$

At $L = 0$, $Q_T = Q_{in}$, so $C = Q_{in}$:

$$Q_T = Q_{in} - (KB + Ea)L$$

To find the total loss (Q_L) associated with a stream of length L :

$$\text{Since } Q_T = Q_{in} - Q_L,$$

$$Q_L = (KB + Ea)L$$

$$A = a \times b$$

Assuming a ratio of $b/a = 0.01$,

$$a = \left(\frac{Q_{in}}{0.36} \right)^{1/2}$$

$$B = 1.02a$$

$$\text{so } Q_L = (1.02 K + E) aL$$

$$= [(1.02 (6 \times 10^{-5} \text{ m/min})) + (2 \times 10^{-6} \text{ m/min})] aL$$

$$= (6.32 \times 10^{-5} \text{ m/min}) aL$$

$$= (6.32 \times 10^{-5} \text{ m/min}) \left(\frac{Q_{in}}{0.36} \right)^{1/2} L$$

To find the reach (R) of perennial stream created by discharge (if there is no net outflow from the section):

$$\text{If } Q_T = 0 \text{ then } 0 = Q_{in} - (KB + Ea)L$$

$$L_Q = 0 = R = \left(\frac{Q_{in}}{KB + Ea} \right)$$

Assuming a ratio of $b/a = 0.01$,

$$a = \left(\frac{Q_{in}}{0.36} \right)^{1/2}$$

$$B = 1.02a$$

$$\text{so } R = \frac{Q_{in}}{(1.02K + E) \left(\frac{Q_{in}}{0.36} \right)^{1/2}} = \frac{(0.36 Q_{in})^{1/2}}{1.02K + E} = 9493.67 \times Q_{in}^{1/2}$$

The basin drainage channels are assumed to be considerably larger than those of the sub-basin and therefore effect a greater evaporation and seepage loss on the system. Consequently, loss rates due to seepage were doubled, producing the following equations for estimating flows within the basin channels:

$$Q_L = (1.24 \times 10^{-4} \text{ m/min}) \left(\frac{Q_{in}}{0.36} \right)^{1/2}$$

$$R = 4823.15 \times Q_{in}^{1/2}$$

Table H.2 contains the calculations of seepage and evaporation loss and cumulative discharge for the sub-basin and contiguous areas of the basin.

Infiltration of mine discharge to ephemeral streams was not calculated separately but instead was solved as a combined loss for evaporation and infiltration. Both losses are a function of surface area. Infiltration takes place in both the sub-basin and the basin and at different rates. The final calculated infiltration percentage represents a combination of the sub-basin and basin losses, which were calculated separately. When 3 mines are operating, the full reach of perennial stream created is 22.7 km.

Infiltration losses in the sub-basin can be calculated as follows:

$$Q_L = [1.02 (6 \times 10^{-5}) + (2 \times 10^{-6})] aL \quad (H.1)$$

where Q_L = flow rate or loss as infiltration plus evaporation, m^3/min
 a = width of stream, meters
 L = length of stream, meters
 Q_s = $6.12 \times 10^{-5} \times aL$
 = infiltration loss, m^3/min
 Q_e = $2.0 \times 10^{-6} \times aL$
 = evaporation loss, m^3/min

Table H.2 Seepage and outflow calculations for the Wyoming model mine drainage system

Section Number ^(a)	Section Length L (m)	Q_{in} (3.00 m ³ /min per mine)	Total Q_{in} (m ³ /min)	Q_L (m ³ /min)	$Q_{in} - Q_L$ (m ³ /min)	R (m)	Cumulative Discharge (m ³ /min)
1	2400	3.00	3.00	0.44	2.56	-	
2	2100	----	----	----	----	-	
3	600	----	2.56	0.10	2.46	-	
4	3000	3.00	3.00	0.55	2.45	-	
5	1400	----	4.91	0.33	4.59	-	
6	1600	3.00	3.00	0.29	2.71	-	
7	900	----	7.30	0.26	7.04	-	7.04
Basin	141000	7.04	7.04	7.04	----	12797	0

(a) See Fig. H.1.

$$\text{Therefore, } \frac{Q_s}{Q_e} = 30.6 \quad (\text{H.2})$$

$$\begin{aligned} \text{Since total loss} &= Q_s + Q_e = 1.96 \text{ m}^3/\text{min} \\ \text{and } Q_s &= Q_e \times 30.6 \\ \text{then } Q_s &= 1.96 - Q_e \text{ and } Q_e = \frac{1.96}{31.6} = 0.062 \text{ m}^3/\text{min} \end{aligned}$$

$$\begin{aligned} \text{Then loss due to infiltration in the sub-basin:} \\ &= 1.96 - 0.062 \\ &= 1.898 \text{ m}^3/\text{min} \end{aligned}$$

Infiltration losses in the basin can be calculated as follows:

$$Q_L = [1.02 (1.2 \times 10^{-4}) + (2 \times 10^{-6})] \text{ aL} \quad (\text{H.3})$$

$$\begin{aligned} \text{where } Q_s &= 1.22 \times 10^{-4} \times \text{aL} \\ Q_e &= 2 \times 10^{-6} \times \text{aL} \end{aligned}$$

$$\text{Therefore, } \frac{Q_s}{Q_e} = 61.0 \quad (\text{H.4})$$

$$\text{Total Loss} = Q_s + Q_e = 7.04 \text{ m}^3/\text{min}$$

$$\begin{aligned} \text{Then loss due to infiltration in the basin:} \\ &= 7.04 - 0.114 \\ &= 6.926 \text{ m}^3/\text{min} \end{aligned}$$

Therefore, total inflow equals $9 \text{ m}^3/\text{min}$ or $4.73 \times 10^6 \text{ m}^3/\text{yr}$, and total annual infiltration loss equals $4.65 \times 10^6 \text{ m}^3$. Restated, 98.2 percent of the discharge infiltrates and the remainder evaporates.

H.1.2 Underground Mines

Mine water discharge from underground mines in the New Mexico model area averages slightly under $2 \text{ m}^3/\text{min}$. Flow characteristics parallel those in the Wyoming model area, as does the methodology applied in calculating the infiltration and evaporation losses and the net outflow. The calculations are based on the operation of 14 mines dewatering at a rate of $28 \text{ m}^3/\text{min}$ ($2.00 \text{ m}^3/\text{min}$ per mine) into a sub-basin of 246 km^2 area. This will result in transformation of 24.8 km of channel (in the sub-basin only) into perennial streams. Note that there is no discharge from the sub-basin to the basin.

Mine discharge is assumed to enter into a small sub-basin hydrographic unit shown in Figure H.3. The sub-basin characteristics used as model input parameters are the same as those for Wyoming (Table H.1) with the exception of those presented in Table H.3.

Table H.3 Characteristics of the sub-basin hydrographic unit in the model underground uranium mine area

Parameter	Symbol	Value	Units
Substrate hydraulic conductivity (vertical)	K_v	6×10^{-5}	m/min
Substrate hydraulic conductivity (horizontal)	K_h	6×10^{-3}	m/min
Substrate hydraulic conductivity	K	6×10^{-4}	m/min
Annual evaporation rate	E	4.0×10^{-6}	m/min
Water input from mine discharge	Q_{in}	2.00 per mine	m^3/min

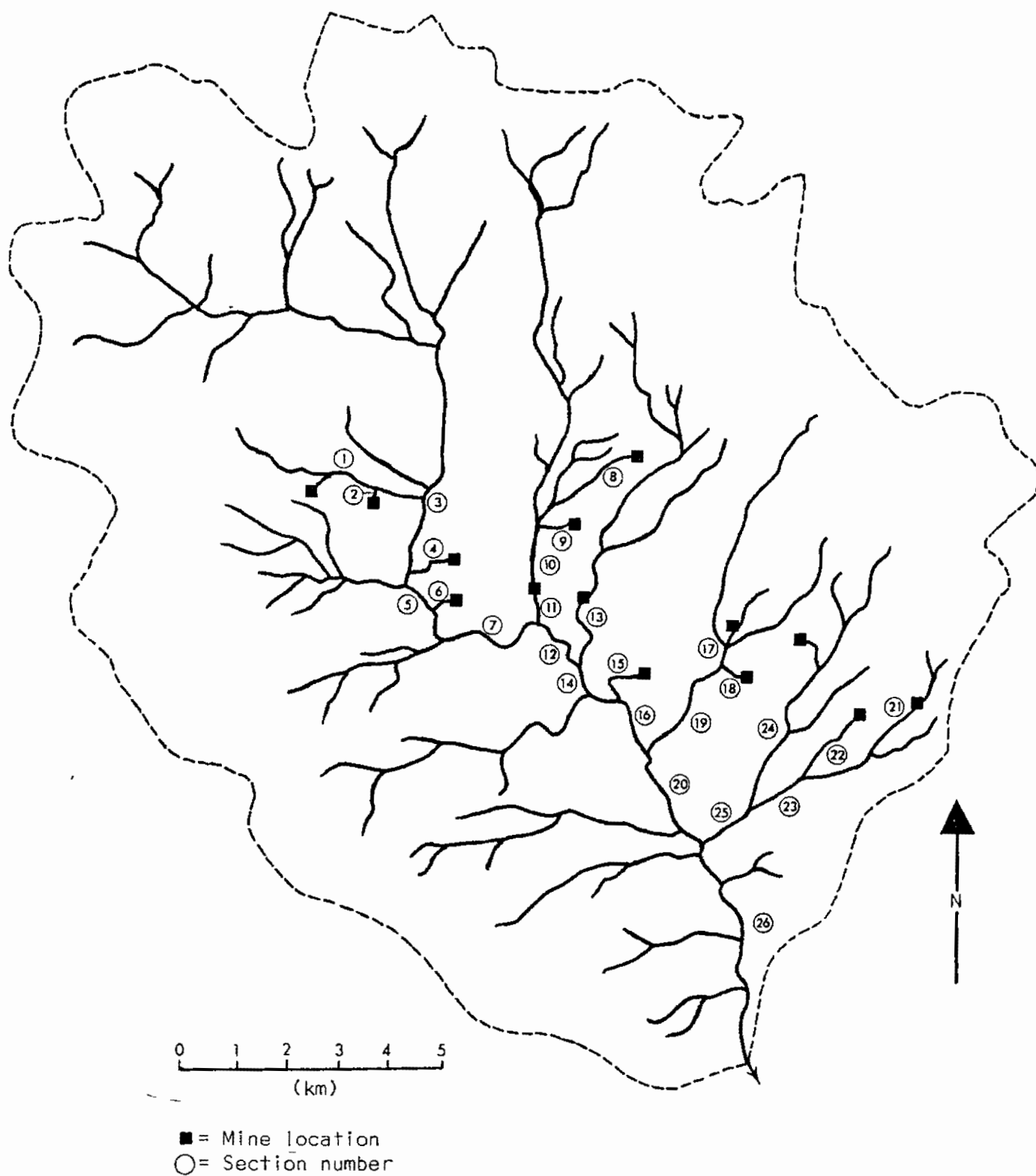


Figure H.3 New Mexico model area sub-basin drainage system

The loss and outflow formulas are of the same format and derivation as the Wyoming equations (see Surface Mines), but have different final forms due to the change in infiltration and evaporation rates. The final equations for the New Mexico model area are as follows:

To find the total loss (Q_L) associated with a stream segment of length L :

$$\begin{aligned} Q_L &= (1.02K + E)aL & (H.5) \\ &= [(1.02 (6 \times 10^{-4} \text{ m/min})) + (4.0 \times 10^{-6} \text{ m/min})] aL \\ &= (6.16 \times 10^{-4} \text{ m/min}) aL \\ &= (6.16 \times 10^{-4}) \left(\frac{Q_{in}}{0.36} \right)^{1/2} L \end{aligned}$$

To find the reach (R) of perennial stream created by discharge (if there is no net outflow from the section):

$$\begin{aligned} R &= \frac{Q_{in}}{(1.02 K + E) \left(\frac{Q_{in}}{0.36} \right)^{1/2}} & (H.6) \\ &= \frac{(0.36 Q_{in})^{1/2}}{(1.02 K + E)} \\ &= 974.03 \times (Q_{in})^{1/2} \end{aligned}$$

Since no net outflow from the sub-basin occurs, equations modified for the basin channel characteristics are not necessary for this model area. Table H.4 presents the calculations of infiltration and evaporation loss and cumulative discharge within the sub-basin.

Infiltration of mine discharge to ephemeral streams was not calculated separately but instead was solved as a combined loss for evaporation and infiltration. Both losses are a function of surface area. Infiltration takes place only in the sub-basin as there is no net outflow. When 14 mines are operating, the full reach of the perennial stream created is 24.8 km.

Infiltration losses can be calculated as follows:

$$Q_L = [(1.02 (6 \times 10^{-4})) + (4.0 \times 10^{-6})] aL \quad (H.7)$$

Table H.4 Seepage and outflow calculations for the New Mexico model mine area drainage system

Section Number ^(a)	Section Length L(m)	Q_{in} (2.00 m ³ /min per mine)	Total Q_{in} (m ³ /min)	Q_L (m ³ /min)	$Q_{in} - Q_L$ (m ³ /min)	R (m)	Cumulative Discharge (m ³ /min)
1	1500	2.00	2.00	-	-	1948	
2	200	2.00	2.00	0.29	1.71	-	
3	2300	0	1.71	-	-	1665	
4	1000	2.00	2.00	1.45	0.55	-	
5	800	0	0.55	-	-	534	
6	500	2.00	2.00	0.73	1.27	-	
7	3000	0	1.27	-	-	1241	
8	3400	2.00	2.00	-	-	1948	
9	1000	2.00	2.00	1.45	0.55	-	
10	700	0	0.55	0.53	0.02	-	
11	500	2.00	2.02	0.73	1.29	-	
12	1100	0	1.29	1.28	0.01	-	
13	1500	2.00	2.00	-	-	1948	
14	1400	0	0.01	-	-	10	
15	1200	2.00	2.00	1.74	0.26	-	
16	1100	0	0.26	-	-	251	
17	1000	2.00	2.00	1.45	0.55	-	
18	600	2.00	2.00	0.87	1.13	-	
19	1900	0	1.68	-	-	1635	
20	2100	0	0	-	-	-	
21	3200	2.00	2.00	-	-	1948	
22	2100	2.00	2.00	-	-	1948	
23	1200	0	0	-	-	-	
24	3900	2.00	2.00	-	-	1948	
25	1300	0	0	-	-	-	
26	4900	0	0	-	-	-	0

(a) See Fig. H.3.

where Q_L = flow rate or loss as infiltration plus evaporation, m^3/min
 a = width of stream, meters
 L = length of stream, meters
 $Q_s = 6.12 \times 10^{-4} \times aL$
 $=$ infiltration loss, m^3/min
 $Q_e = 4.0 \times 10^{-6} \times aL$
 $=$ evaporation loss, m^3/min

$$\text{Therefore, } \frac{Q_s}{Q_e} = 153 \quad (H.8)$$

Since total loss = $Q_s + Q_e = 28 m^3/min$

and $Q_s = Q_e \times 153$

then $Q_s = 28 - Q_e$ and $Q_e = \frac{28}{154} = 0.18 m^3/min$

Then loss due to infiltration in the sub-basin:

$$= 28 - 0.18$$

$$= 27.82 m^3/min$$

Therefore, total inflow equals $28.0 m^3/min$ or $1.47 \times 10^7 m^3/yr$, and total annual infiltration loss equals $1.46 \times 10^7 m^3$. Restated, 99.3 percent of the discharge infiltrates and the remainder evaporates.

APPENDIX I
COMPUTATION OF MASS EMISSION FACTORS
FOR WIND EROSION

I.1.0 Computation of Mass Emission Factors for Wind Erosion

Mass emission factors due to wind erosion of overburden, waste rock, and sub-ore piles were computed by the equation (Hu76)

$$EF = aIKCLV \quad (I-1)$$

where,

EF = emission factor, MT/hectare-yr,

a = the portion of total wind erosion losses that would be measured as suspended particulates - a has the value of 0.025 for rocky, gravelly surfaces (Hu76),

I = soil erodibility - I has the value of 85 MT/hectare-yr for rocky, gravelly surfaces (Hu76),

K = surface roughness factor, assumed to be 1.0,*

C = climatic factor - reported to be 1.0 for New Mexico and 0.40 for Wyoming mining regions (Hu76),

L = unshielded field width factor - assumed to be 1.0,* and

V = vegetative cover factor - assumed to be 1.0*.

Substituting the assigned values into the equation yields an emission factor of 2.12 MT/hectare-yr for New Mexico mines and 0.850 MT/hectare-yr for Wyoming mines. These factors are applied with appropriate parameters in Sections 3.3.4.1 and 3.4.4.3 to estimate the average annual contaminant emissions due to wind erosion.

The mass emission factors due to wind erosion of the ore stockpiles were computed by the equation (Bo78)

$$EF = 0.025 \frac{(S)}{1.5} \frac{(D)}{90} \frac{(d)}{235} \frac{(f)}{15} \quad (I-2)$$

*Dale, J.T., 1979, Air Program Branch, U.S. Environmental Protection Agency, Region VIII, Denver, CO, Memo Concerning Uranium Resources Development Company's Mining Operation in San Juan County, Utah - PDS Permit Requirements.

where,

- EF = emission factor, kg of dust per annual MT of material put through storage cycle,
- S = silt content - assumed to be 3.0,*
- D = duration of storage - 41 days,
- d = dry days per year - reported to be 273 days at Casper, WY and 306 days at Albuquerque, NM (DOC77), and
- f = percent of the time the wind speed exceeds 19.3 km/hr - reported to be 49 percent at Casper, WY and 20 percent at Albuquerque, NM (DOC51-60).

Substituting the assigned values into Equation I-2 provides emission factors related to ore storage piles at mines in New Mexico and Wyoming of 0.040 kg/MT and 0.086 kg/MT, respectively. These factors are applied with the appropriate parameters in Sections 3.3.4.1 and 3.4.4.3 to estimate the average annual contaminant emissions from ore stockpiles due to wind erosion.

I.2.0 References

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- DOC51-60 Department of Commerce, U.S. Weather Bureau, 1951-1960, "Climatography of the United States Series 82 -- Decennial Census of the United States Climate."
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- Hu76 Hubbard, S.J., 1976, "Evaluation of Fugitive Dust Emissions from Mining: Task 1 Report - Identification of Fugitive Dust Sources Associated with Mining," Report prepared by PEDCO-Environmental Specialists, Inc. for the U.S. Environmental Protection Agency, Las Vegas, NV.

APPENDIX J

AQUATIC DOSIMETRY AND HEALTH EFFECTS MODELS
AND PARAMETER VALUES

J.1 Introduction

This appendix describes the methodology and the parametric values used to estimate dose equivalents and health risks for maximum and average individuals and for the population within the assessment area surrounding the generic underground uranium mine site in New Mexico and the generic surface mine site in Wyoming. In the appendix, the term dose equivalents refers to the following:

For radionuclides inhaled or ingested by an individual, dose equivalents are the annual committed dose equivalents that will be accumulated over 70 years following intake for an adult. For external exposure to an individual, dose equivalents are the annual dose equivalents to an adult for a radionuclide buildup time in the environment of 8.5 years (one-half the assumed period of mine operation of 17 years). For radionuclides inhaled or ingested by the population, dose equivalents are the annual collective dose equivalents that will be accumulated over 70 years following intake for an adult. For external exposure to a population, dose equivalents are the annual collective dose equivalents to an adult for a radionuclide buildup time in the environment of 8.5 years (one-half the assumed period of mine operation of 17 years).

Simple models are used throughout these calculations. The maximum individual calculations are performed independently from the population calculations. However, the average individual calculations are obtained by dividing the population calculations by the population size.

J.2 Pathways Considered

The aquatic pathway analyses consider the general surface stream flow pattern shown in Fig. J.1. The generic mine is dewatered by pumping water into the third order stream. Much of the year the only water flowing in the third order stream is mine water discharge. As the mine water travels downstream in the third order stream, it either percolates into the soil beneath the stream or evaporates. At some location downstream before the third order stream enters the second order stream, the third order stream dries up so that, for a large part of the year, the radioactive discharges in the mine water do not reach the second nor the first order streams. Some of the discharged radioactivity moves into the soil beneath the third order stream

bed and the rest is probably deposited in the sediment on the stream bottom. The radioactivity deposited in the sediment is subject to resuspension and transport to the second order and first order streams during periods of flooding. However, during these floods, the resuspended radioactivity would be subject to the large dilution volumes associated with the flood. Some of the radioactivity that percolates into the soil could eventually reach shallow groundwater, but many of the radionuclides would be subject to large removal factors because of ion-exchange interactions between radionuclides and components of the soil. These interactions would cause groundwater concentrations of radionuclides to be greatly reduced when compared to the original surface water concentrations.

An effort has been made to use realistic computational methods and parametric data in these analyses whenever possible. However, when actual data did not exist, some conservative assumptions were made. The continuous stream flow assumption discussed in the following paragraph is the major conservative assumption in these aquatic analyses.

To correctly analyze the effects of radionuclides discharged from the mine to third order streams, one would have to have extensive hydrologic information on stream flow rates over a long period of record in order to predict the quantities of radionuclides reaching the second order and first order streams. One would also need extensive information on soil types and ion-exchange characteristics of the soils within the stream bed areas. Because these data are not obtainable for use in these analyses, the simplifying assumption is made that the third order stream discharges mine water continuously into the second order stream and that the second order stream discharges continuously into the first order stream. Thus, the water concentrations of radionuclides in the first order stream are computed by dividing the annual radionuclide discharge from one mine by the average-annual flow rate in the first order stream. Population and average individual dose equivalents and health effects estimates are calculated using these water concentrations. Maximum individual dose equivalents and health effects are computed using the water concentrations in the second order stream which are computed by dividing the annual radionuclide discharges from one mine by the annual average flow rate for the second order stream. These

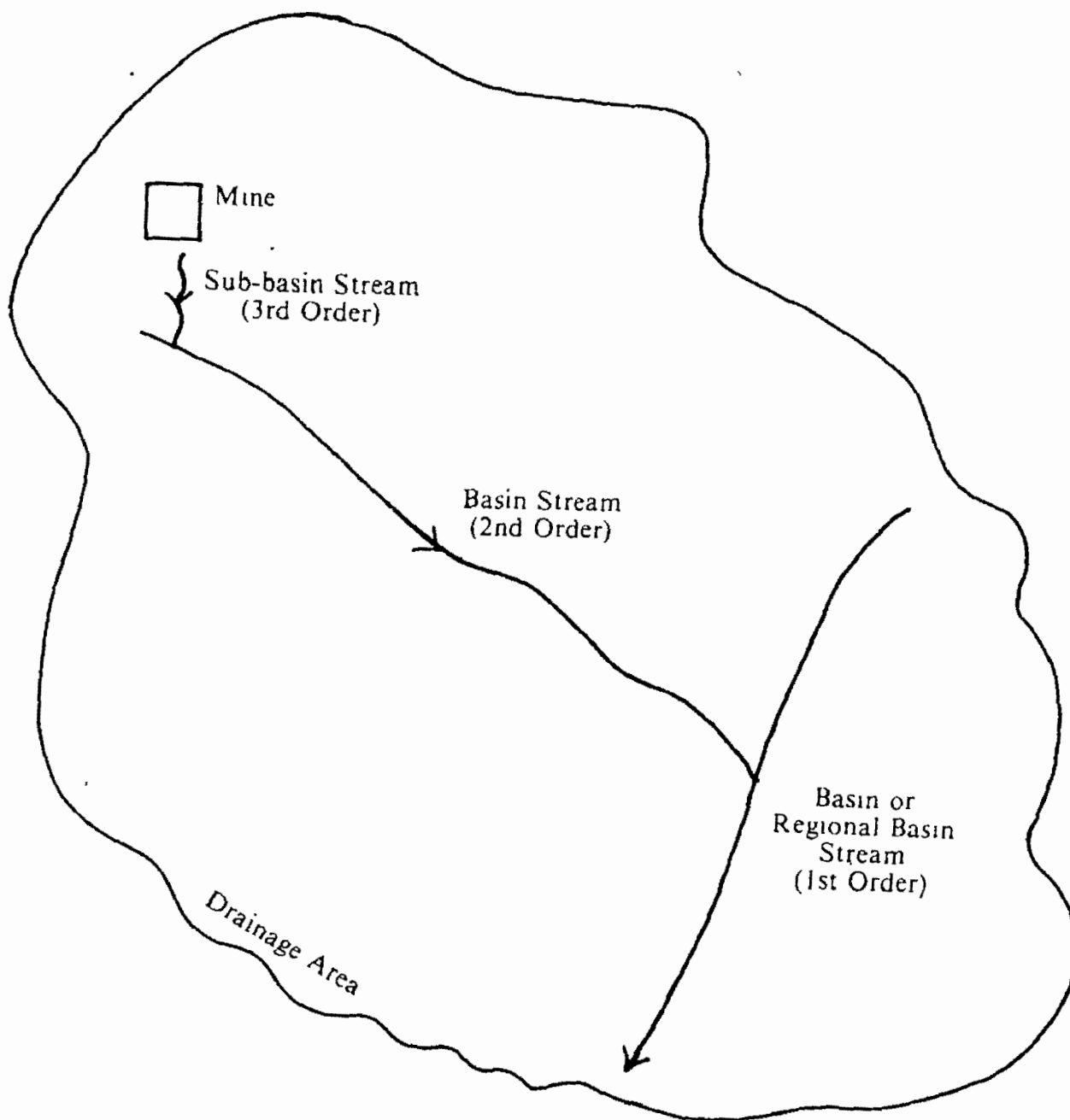


Figure J.1 Surface stream flow pattern within drainage area

techniques for predicting water concentration should be quite conservative since they do not account for periods of no-flow in the third order streams nor for loss of radioactivity from the water by percolation into the soil.

The environmental transport pathways which are examined as potential contributors of dose equivalents and health effects to individuals and to the population are listed in Table J.1. It was found that essentially all potable drinking water in both the New Mexico and the Wyoming assessment areas is taken from groundwater supplies. Public water supplies are taken from aquifers below the elevation of the aquifers which could be affected by recharge from the mine surface water. For this reason, drinking water (pathway 1) is not considered as a pathway of exposure to the population. Drinking water could be a significant pathway of exposure for the maximum individual living near a uranium mine if he were drinking surface water downstream of the mine discharge point or water from a shallow aquifer which had been recharged by the mine surface water. However, this pathway was not assessed because it was not possible to quantify radionuclide concentrations in potable groundwater from mine discharges with available information. Also, it is believed that the occurrence of direct consumption by individuals of surface water containing mine discharges would be infrequent or non-existent.

Through telephone conversations, it was learned that almost all milk cows in the assessment areas obtain their drinking water from groundwater supplies. In addition, most of the dairy cattle are located 50 km or more away from the uranium mines (Romo, T., Valencia County Agent, Los Lunas, NM, 1979, personal communication and Loper, R., Soil Conservation Service, U.S. Department of Agriculture, Douglas, WY, 1979, personal communication). For these reasons, consumption of milk produced by cows drinking contaminated water is not considered in the population dose equivalent calculations. Thus, the pathways listed in Table J-1 which are considered for the population dose equivalent calculations are 2 through 8 and 10. For the maximum individual dose equivalent calculations, the pathways considered are 2 through 10.

For these generic site analyses, detailed data are not readily available regarding fish catch and surface stream water usage as a function of distance downstream from the mine for the third, second, and first order streams.

Thus, a simplified approach is taken in computing the population dose equivalents and health effects. The population assessment area is defined as the areas draining into the streams shown in Fig. J.1. These drainage areas are discussed in more detail in Sections 3.3.3 and 3.4.3*. To compute the population dose equivalents and health effects, the assumption is made that the water concentrations in the first order stream are representative of the radionuclide concentrations to which persons within the assessment area could be exposed. The resulting river water concentrations of radionuclides are applied to each of the eight pathways for which population dose equivalents are calculated.

The mathematical models used for the population and maximum individual dose equivalents and health effects calculations will be discussed in the following paragraphs. Average individual dose equivalents and health effects are computed by dividing the population values by the number of persons in the assessment area. For each pathway, dose equivalents are calculated for endosteal cells (bone), red bone marrow, lung, liver, stomach wall, lower large intestine (LLI) wall, thyroid, kidney, muscle, ovaries, testes and a weighted mean dose equivalent which is weighted over all organs. For the external exposure pathways (pathways 7 and 8) these dose equivalents are used to estimate health impact by applying a dose-to-health-impact conversion factor. This factor is a function of organ and is independent of the nuclide involved. However, for all of the internal exposure pathways (pathways 1 through 6, 9, and 10) dose equivalents are calculated and reported only for the purpose of supplying this information to the reader. The health impact for each pathway is estimated by computing the quantity of each radionuclide taken in by the population or a maximum individual and applying an intake-to-health-impact conversion factor. This factor is a function of the nuclide involved but is not a function of organ. When estimates of health impacts have been computed for each pathway and radionuclide involved, they may be summed over pathways and radionuclides to obtain an estimate of total health impacts to the population, and the average and maximum individuals.

*For the Wyoming site, the assessment area is the 13,650 km² regional basin drainage area of the Cheyenne River discussed in Section 3.3.3.1.2. For the New Mexico site, the assessment area is the 19,037 km² basin drainage area of the Rio San Jose-Rio Puerco Rivers discussed in Section 3.4.3.1.2.

Table J.1 Aquatic environmental transport pathways examined

Pathway Number	Pathway
1	Drinking water ingestion
2	Freshwater fish ingestion
3	Above-surface crops ingestion -- Irrigated Cropland
4	Milk ingestion -- cows consuming forage raised on irrigated pastures
5	Beef ingestion -- cows consuming forage raised on irrigated pastures
6	Inhalation of resuspended material deposited during irrigation
7	External exposure due to ground contamination by material deposited during irrigation
8	External exposure due to air submersion in resus- pended material originally deposited during irrigation
9	Milk ingestion -- cows drinking contaminated surface water
10	Beef ingestion -- cows drinking contaminated surface water

J.3 Freshwater Fish Ingestion

The computational equation for intake of a radionuclide by the maximum individual is

$$II_{np} = \frac{Q_n \cdot CF_n \cdot I_f}{R_{ind}} \quad (J.1)$$

where,

II_{np} = annual intake of radionuclide n through pathway p for the maximum individual (Ci/y),

Q_n = release rate of radionuclide n to the third order stream (Ci/y),

CF_n = concentration factor for freshwater fish for radionuclide n (Ci/kg per Ci/l),

I_f = freshwater fish annual consumption rate for an individual (kg/y),
and

R_{ind} = flow rate in 2nd order stream (l/y).

The maximum individual dose equivalent may be computed from the equation

$$DI_{nop} = II_{np} \cdot D_{nop} \quad (J.2)$$

where,

DI_{nop} = dose equivalent rate to the maximum individual for nuclide n, organ o, and pathway p (rem/y), and

D_{nop} = dose equivalent conversion factor for nuclide n, organ o, and pathway p (rem/Ci intake unless specified otherwise).

The maximum individual increased health risk may be computed using the equation

$$IR_{np} = II_{np} \cdot HI_{np} \quad (J.3)$$

where,

IR_{np} = increased annual health risk* rate to the maximum individual for nuclide n and pathway p (increase in risk per year of release), and
 HI_{np} = health risk conversion factor for nuclide n and pathway p (increase in risk/Ci intake).

Equation J.3 may be applied for computing either fatal cancer risk or genetic risk to future generations by applying HI_{np} for fatal cancer risk or genetic risk, respectively.

The equation used to compute total intake of a radionuclide by the population was

$$IP_{np} = \frac{Q_n \cdot CF_n \cdot I_f \cdot P_{ff}}{R}, \quad (J.4)$$

where,

IP_{np} = annual intake of radionuclide n through pathway p for the population (person Ci/y),
 R = flow rate in 1st order stream (l/y), and
 P_{ff} = population eating freshwater fish taken from streams in the assessment area (persons).

The population dose equivalent may be computed from the equation

$$DP_{nop} = IP_{np} \cdot D_{nop} \quad (J.5)$$

where,

DP_{nop} = annual dose equivalent to the population for nuclide n, organ o, and pathway p (person-rem/y);

*The term health risk is used to describe the increase in fatal cancer risk to an individual during his lifetime for somatic risks. For genetic risk, the term health risk refers to the increased chance for genetic defects in all the descendants of an exposed individual.

and the annual health effects to the population may be computed using the equation

$$PR_{np} = IP_{np} \cdot HI_{np} \quad (J.6)$$

where,

PR_{np} = annual health effects to the population for nuclide n and pathway p (health effects /y of release).

Equation J.6 may be applied for computing either fatal cancers or genetic effects to future generations by applying the proper value for HI_{np} , as discussed above.

The annual dose equivalent and increased health risk to an average individual may be computed using equations J.7 and J.8, respectively.

$$DAI_{nop} = DP_{nop}/P_T \quad (J.7)$$

$$AIR_{np} = PR_{np}/P_T \quad (J.8)$$

where,

DAI_{nop} = annual dose equivalent to the average individual within the assessment area for nuclide n, organ o, and pathway p (rem/y),

AIR_{np} = increased annual risk to the average individual for nuclide n and pathway p (increase in risk per year of release), and

P_T = size of population residing within the assessment area (persons).

J.4 Above-Surface Crops, Milk, and Beef Ingestion - Irrigated Crop Land

The equation used to compute intake of a radionuclide by the maximum individual was

$$II_{np} = \frac{Q_n \cdot W \cdot RI_{np}}{R_{ind}} \quad (J.9)$$

where,

W = irrigation rate of irrigated farmland ($\ell/m^2\text{-y}$), and

RI_{np} = intake rate of radionuclide n by standard man for above-surface crops and for a continuous deposition rate to the surface for root uptake due to ditch irrigation (C_i intake/y per $C_i/m^2\text{-y}$ deposited).

The approach for calculation of RI_{np} uses techniques described in Regulatory Guide 1.109 (NRC77) and AIRDOS-EPA (Mo79) to compute intake rate by receptors per unit deposition rate (by ditch irrigation) to the ground surface. Basically, equations 49 (vegetation), 51 (milk), or 52 (beef) from the AIRDOS-EPA document (Mo79) are utilized to predict concentrations of radionuclides in the foodstuffs at equilibrium. Only the root uptake portion of these equations is used since essentially all irrigation in the assessment areas is ditch irrigation (Romo, T., Valencia County Agent, Los Lunas, NM., 1979, personal communication, and Loper, R., Soil Conservation Service, U.S. Department of Agriculture, Douglas, WY, 1979, personal communication). The effects of removal of radionuclides from the soil root zone by leaching are added to the equations used to predict concentrations of radionuclides in foodstuffs since this is an important removal mechanism for the long-lived radionuclides considered in this analysis. The concentrations are multiplied by the annual intake rate of the foodstuff by an individual and divided by the annual deposition rate of radionuclides to the ground surface to yield the quantity RI_{np} . The maximum individual dose equivalents may be computed using equation J.2 and the increased health risk to the maximum individual may be computed using equation J.3.

The equation to use in computing total intake of a radionuclide by the population is

$$IP_{np} = \frac{Q_n \cdot W \cdot RI_{np} \cdot P_p}{R} \quad (J.10)$$

where,

P_p = population consuming foodstuffs raised on irrigated land (persons).
 P_p can be determined using the equation

$$P_p = CP_p \cdot f_p \cdot A_I \quad (J.11)$$

and the ratio can be written

$$\frac{W/R}{A_I \cdot R} = \frac{R_I}{A_I \cdot R} = \frac{f_R}{A_I} \quad (J.12)$$

where,

CP_p = persons per unit area that can be fed from foodstuff p raised on irrigated land (persons fed/m²),

f_p = fraction of irrigated land used to raise foodstuff p,

A_I = irrigated land area within the assessment area (m²),

R_I = total flow of irrigation water (ℓ/yr), and

$f_R = \frac{R_I}{R}$ = fraction of river flow used for irrigation.

Substituting equations J.11 and J.12 into J.10 and cancelling like terms yields

$$IP_{np} = Q_n \cdot f_R \cdot RI_{np} \cdot CP_p \cdot f_p. \quad (J.13)$$

The population annual dose equivalents may be computed using equation J.5 and the annual increased health effects to the population may be computed using equation J.6. The annual dose equivalents and increased health risks to an average individual may be computed using equations J.7 and J.8.

J.5 Inhalation of Resuspended Material Deposited During Irrigation

In determining the equations to use to model resuspension of radioactive materials deposited by irrigation water, it is assumed that the resuspended material does not disperse beyond the irrigation area. This assumption should be acceptable for dose equivalent and health effects calculations where the irrigation area is large compared to a point resuspension source and where the population density does not vary greatly within an assessment area. Both of these criteria are met for the New Mexico and the Wyoming assessment areas.

Figure J.2 shows, pictorially, the conservation of mass relationship used in the resuspension model. The differential equation which expresses the change in soil surface concentration as a function of time is

$$\frac{d\Omega_n}{dt} = \frac{Q_n \cdot W}{R} + v_{gn} \cdot \chi_n - \lambda_R \Omega_n - \lambda_{Dn} \Omega_n - \lambda_{sn} \Omega_n \quad (J.14)$$

where,

Ω_n = ground concentration of nuclide n at time t (Ci/m^2),

t = time after release of material to surface stream (y),

v_{gn} = deposition velocity from air to land surface (m/y),

X_n = air concentration of nuclide n (Ci/m^3),

λ_R = rate constant for resuspension of radionuclides from soil to air (y^{-1}),

λ_{Dn} = radioactive decay constant for radionuclide n (y^{-1}),

λ_{sn} = rate constant for transfer of radionuclides from available to unavailable status in soil (y^{-1}),

and the other terms are as previously defined.

If it is further assumed that, at equilibrium, the material resuspended from the ground surface is equal to the material redeposited to the ground surface (i.e., $v_{gn}X_n = \lambda_R\Omega_n$), the differential equation can be simplified to yield

$$\frac{d\Omega_n}{dt} = \frac{Q_n \cdot W}{R} - (\lambda_{Dn} + \lambda_{sn})\Omega_n \quad (\text{J.15})$$

Equation J.15 would rigorously hold only for equilibrium conditions. It can be shown that it is conservative to apply the nonequilibrium initial conditions,

$\Omega_n = 0$ at $t = 0$, in solving equation J.15 to yield

$$\Omega_n = \frac{Q_n \cdot W}{R(\lambda_{Dn} + \lambda_{sn})} [1 - e^{-(\lambda_{Dn} + \lambda_{sn})t}]$$

When using equation J.16 in computations involving the maximum individual, R is replaced by R_{ind} .

As mentioned previously, assuming a fairly large uniformly contaminated area, the air concentration of radionuclides due to resuspension can be expressed as

$$X_n = RF \cdot \Omega_n \quad (\text{J.17})$$

where,

$RF = \lambda_R/v_{gn}$ = resuspension factor (m^{-1}).

Combining equations J.16 and J.17 yields

$$X_n = \frac{Q_n \cdot W \cdot RF}{R(\lambda_{Dn} + \lambda_{sn})} [1 - e^{-(\lambda_{Dn} + \lambda_{sn})t}] \quad (\text{J.18})$$

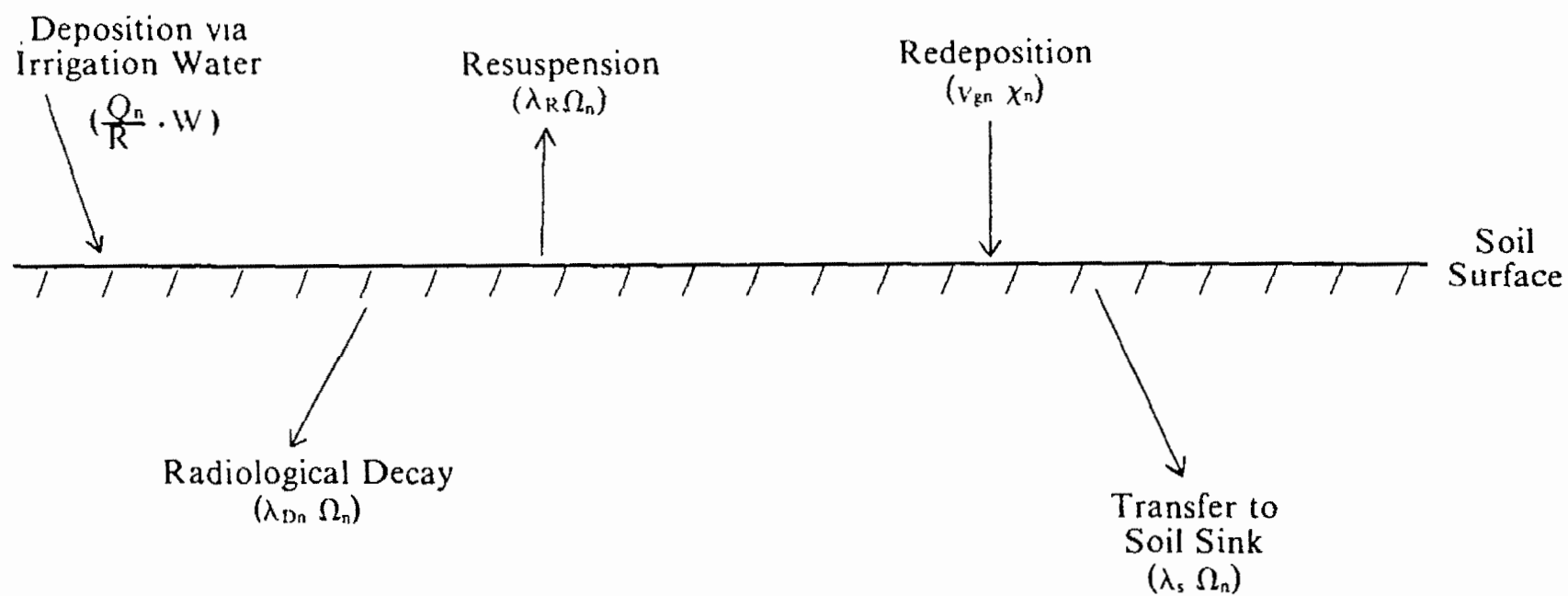


Figure J.2 Conservation of mass relationship for resuspension model

This equation is appropriate to apply for the population dose equivalent calculations. For calculations involving the maximum individual, R is replaced by R_{ind} .

The equation used to compute intake of a radionuclide by the maximum individual is

$$II_{np} = X_n I_B \quad (J.19)$$

where,

I_B = breathing rate for standard man (m^3/y).

Combining equations J.18 and J.19 yields

$$II_{np} = \frac{Q_n \cdot W \cdot RF \cdot I_B}{R_{ind} (\lambda_{Dn} + \lambda_{sn})} [1 - e^{-(\lambda_{Dn} + \lambda_{sn})t}] \quad (J.20)$$

The maximum individual annual dose equivalents may be computed using equation J.2 and the maximum individual annual increased health risks may be computed using equation J.3.

The equation to use in computing total intake of a radionuclide by the population is

$$IP_{np} = X_n \cdot I_B \cdot p \cdot A_I \quad (J.21)$$

where,

p = population density within the assessment area ($person/m^2$).

After substitution of equations J.18 and J.12, we have

$$IP_{np} = \frac{Q_n \cdot f_R \cdot RF \cdot I_B \cdot P}{(\lambda_{Dn} + \lambda_{sn})} [1 - e^{-(\lambda_{Dn} + \lambda_{sn})t}]. \quad (J.22)$$

The annual population dose equivalents may be computed using equation J.5 and the annual increased health effects to the population may be computed using equation J.6. The annual dose equivalents and increased health risks to an average individual may be computed using equations J.7 and J.8.

J.6 External Exposure Due to Ground Contamination by Material Deposited from Irrigation Water

As was discussed earlier, health risks are computed from dose equivalents for the external exposure pathways. The equation used in computing annual dose

equivalents to the maximum individual is

$$DI_{nop} = \Omega_n \cdot D_{nop} \cdot SOF \quad (J.23)$$

where,

D_{nop} = dose equivalent factor for external ground contamination for nuclide n, organ o, and pathway p (rem/y per Ci/m²), and
 SOF = household shielding and occupancy factor (dimensionless).

Upon substitution of equation J.16 we have (substituting R_{ind} for R)

$$DI_{nop} = \frac{Q_n \cdot W \cdot D_{nop} \cdot SOF}{R_{ind} (\lambda_{Dn} + \lambda_{sn})} [1 - e^{-(\lambda_{Dn} + \lambda_{sn})t}]. \quad (J.24)$$

The increase in annual health risks for the maximum individual can be computed using the equation

$$IR_{nop} = DI_{nop} \cdot HE_{op} \quad (J.25)$$

where,

IR_{nop} = increased annual risk to the maximum individual for nuclide n, organ o, and pathway p (increase in risk/year of release), and
 HE_{op} = health risk conversion factor for external doses for organ o and pathway p (increase in risk/rem).

The equation for computing annual external population dose equivalents due to uniform ground contamination is

$$DP_{nop} = \Omega_n \cdot D_{nop} \cdot SOF \cdot p \cdot A_I \quad (J.26)$$

Substitution of equations J.16 and J.12 into this equation yields

$$DP_{nop} = \frac{Q_n \cdot f_R \cdot D_{nop} \cdot SOF \cdot p}{(\lambda_{Dn} + \lambda_{sn})} [1 - e^{-(\lambda_{Dn} + \lambda_{sn})t}]. \quad (J.27)$$

For equations J.24 and J.27, the exposed persons are represented as a point receptor 1 m above a plane surface with a uniform distribution of radioactivity. The increase in health risks for the population are estimated using the equation

$$PR_{nop} = DP_{nop} \cdot HE_{op} \quad (J.28)$$

where,

PR_{nop} = increased annual health effects to the population for nuclide n, organ o, and pathway p (health effects /y of release).

The annual dose equivalents and increased health risks to an average individual may be computed using equations J.7, and J.29, respectively:

$$AIR_{nop} = PR_{nop}/P_T \quad (J.29)$$

where,

AIR_{nop} = annual increased health risk to the average individual for nuclide n, organ o, and pathway p (increase in risk/year of release).

J.7 External Exposure Due to Air Submersion in Resuspended Material Originally Deposited During Irrigation

The equation applied in calculating maximum individual annual external dose equivalents due to submersion is

$$DI_{nop} = X_n \cdot D_{nop} \cdot SOF \quad (J.30)$$

where,

D_{nop} = dose equivalent conversion factor for external air submersion for nuclide n , organ o , and pathway p (rem/y per Ci/m³).

Upon combining equations J.30 and J.18, the dose equivalent rate equation becomes

$$DI_{nop} = \frac{Q_n \cdot W \cdot RF \cdot D_{nop} \cdot SOF}{R_{ind} (\lambda_{Dn} + \lambda_{sn})} [1 - e^{-(\lambda_{Dn} + \lambda_{sn})t}]. \quad (J.31)$$

The increase in annual health risks for the maximum individual can be computed using equation J.25.

The equation for computing external population annual dose equivalents due to air submersion is

$$DP_{nop} = X_n \cdot D_{nop} \cdot SOF \cdot p \cdot A_I \quad (J.32)$$

Substitution of equations J.18 and J.12 into equation J.32 yields

$$DP_{nop} = \frac{Q_n \cdot f_R \cdot RF \cdot D_{nop} \cdot SOF \cdot p}{(\lambda_{Dn} + \lambda_{sn})} [1 - e^{-(\lambda_{Dn} + \lambda_{sn})t}]. \quad (J.33)$$

Equations J.31 and J.33 are derived assuming a point receptor immersed in a semi-infinite hemispherical cloud of air in which the distribution of activity is spatially uniform.

The increase in annual health risks for the population are estimated using equation J.28. The annual dose equivalent and increased health risk to an average individual may be computed using equations J.7 and J.29, respectively.

J.8 Milk Ingestion--Cows Drinking Contaminated Surface Water

The computational equation for intake of a radionuclide by the maximum individual is

$$II_{np} = \frac{Q_n \cdot I_{wm} \cdot F_{mn} \cdot I_{mF}}{R_{ind}} \quad (J.34)$$

where,

- I_{wm} = milk cow drinking water ingestion rate (ℓ/d),
- F_{mn} = concentration of radionuclide n in the milk per unit daily intake of the radionuclide via cattle drinking water (Ci/ℓ milk per Ci/day), and
- I_{mF} = adult consumption rate of milk (ℓ/y).

The maximum annual individual dose equivalents may be computed from equation J.2 and the maximum annual individual increased health risks may be computed using equation J.3. Annual population dose equivalents and increased health impacts were not calculated for this pathway since it was determined that consumption of contaminated drinking water by milk cows would be infrequent or nonexistent.

J.9 Beef Ingestion--Cows Drinking Contaminated Surface Water

The computational equation for intake of a radionuclide by the maximum individual is

$$II_{np} = \frac{Q_n \cdot I_{WB} \cdot F_{Bn} \cdot I_{BF}}{R_{ind}} \quad (J.35)$$

where,

- I_{WB} = beef cattle drinking water ingestion rate (ℓ/d),
- F_{Bn} = concentration of radionuclide n in beef per unit daily intake of the radionuclide via cattle drinking water (Ci/kg beef per Ci/day), and
- I_{BF} = adult consumption rate of beef (kg/y).

The maximum individual annual dose equivalents may be computed from equation J.2 and the maximum individual annual increased health risks may be computed using equation J.3.

For computing total intake of a radionuclide by the population, the appropriate equation is

$$IP_{np} = \frac{Q_n \cdot I_{WB} \cdot F_{Bn} \cdot I_{BF} \cdot P_{BW}}{R} \quad (J.36)$$

where,

P_{BW} = number of persons eating beef from cows which drink contaminated water (persons).

The annual population dose equivalents may be computed from equation J.5 and the annual increased health effects to the population may be computed using equation J.6. The annual dose equivalents and increased health risks to an average individual may be computed using equations J.7 and J.8, respectively.

J.10 Generic Sites

Two generic sites were chosen to represent locations where uranium mines may be located. A generic site in Converse County, Wyoming, was chosen to represent typical surface mine sites and one in Valencia County, New Mexico, was chosen to represent typical underground mining sites. Figure J.1 shows the general assessment area for both generic sites. These sites are described more fully in Sections 3.3.3 and 3.4.3 and in Subsection J.2 of this appendix. Some of the characteristics of these sites used in the dose equivalent and health effects calculations are listed in Table J.2.

Table J.2 Characteristics of the generic sites

	New Mexico	Wyoming
Annual rainfall (cm)	20	28
Total population in assessment area	64,950	16,230
Population density in assessment area (persons/m ²)	3.41×10^{-6}	1.19×10^{-6}
Assessment area size (km ²)	19,037	13,650
Streams within assessment area ^(a) (avg. annual flow rate, g/yr)	Arroyo del Puerto-San Mateo Creek	Not named
Third order	(small)	(small)
Second order	Rio San Jose (5.83×10^9)	Lance Creek (2.18×10^{10})
First order	Rio Puerco (4.26×10^{10})	Cheyenne, Dry Fork (5.64×10^{10})
Number of persons eating fish containing radionuclides from mine discharges	6,495	1,623
Annual irrigation rate within assessment areas (m)	1.07(b)	0.59
Fraction of annual-average first order stream flow used for irrigation	0.30	0.29
Land area irrigated within assessment area (km ²)	12 ^(b)	28
Number of persons eating beef from cows drinking contaminated water	38,510	3,454

(a) See Sections 3.3.3 and 3.4.3 for a full discussion of assessment area streams and hydrology.

(b) Some groundwater used for irrigation. See discussion in Subsection J.13.

J.11 Population and Population Density in Assessment Areas

For both generic sites, the population within the assessment area is determined by computing the population density for a county containing a significant part of the assessment area and then scaling this population density up to the assessment area size. For New Mexico, Valencia County data are used (Romo, T., Valencia County Agent, Los Lunas, NM, 1979, personal communication). The current population of Valencia County is about 50,000 persons. The county area is $14,650 \text{ km}^2$. The population density, based on these data, is $3.41 \times 10^{-6} \text{ persons/m}^2$. Considering the size of the assessment area to be $19,037 \text{ km}^2$ (Section 3.4.3), the estimated total population in the assessment area is 64,950 persons. For Wyoming, Converse County data are applied (Zaborac, J., Converse Area Planning Office, Douglas, WY, 1979, personal communication). The estimated current population of Converse County is 13,000 persons and the county area is $10,930 \text{ km}^2$. Thus, the population density is calculated to be $1.19 \times 10^{-6} \text{ persons/m}^2$. The assessment area contains $13,650 \text{ km}^2$ (Section 3.3.3); therefore, the total population within the Wyoming assessment area is estimated to be 16,230.

J.12 Population Consuming Fish Containing Radionuclides Discharged from Mines

Information on fish catch specific to the assessment areas is not available. However, three sources agreed that there is very little fishing activity in the streams in these areas (Patterson, R., New Mexico Game and Fish Department, Santa Fe, NM, 1979, personal communication; Baughman, J., Wyoming Game and Fish Division, Cheyenne, WY, 1979, personal communication; and Kaufmann, R., U.S. Environmental Protection Agency, Office of Radiation Programs, Las Vegas Facility, Las Vegas, NV, 1979, personal communication). Considering these data and the lack of specific information, it is assumed that 10 percent of the population within each assessment area consumes fish taken from streams within the assessment area. Based on the hydrologic characteristics of these areas, this is probably a conservative assumption.

J.13 Irrigation Within the Assessment Areas

For the New Mexico site, the following irrigation information was

obtained for Valencia County (Romo, T., Valencia County Agent, Los Lunas, NM, 1979, personal communication). Valencia County is a very large county and the Rio Grande flows through the eastern part of the county at a distance of about 100 km from the uranium mine site. Since it is known that a large amount of irrigation within the county occurs along the Rio Grande and since the Rio Grande was not included in the assessment area, the effect of the irrigation along the Rio Grande was extracted from the county data in estimating the amount of irrigated land within the assessment area. The calculational procedure is described below.

The three major streams for irrigation in Valencia County are the Rio San Jose (second order), the Rio Puerco (first order), and the Rio Grande. The average irrigation rate is 1.07 m/yr. Within Valencia County, 11,234 hectares are irrigated with surface water only, and 5,306 hectares use a combination of surface water and groundwater for irrigation. It is assumed that for this 5,306 hectares 50 percent of the irrigation water is surface water and 50 percent of this is groundwater that does not contain mine-related radionuclides. To calculate the equivalent acres of land totally irrigated from the Rio San Jose and the Rio Puerco within Valencia County we assume that the amount of irrigation existing along a stream is directly proportional to the product of the annual-average flow rate and the length of the stream within the county. Thus a ratio is established to predict the fraction of total land irrigated in Valencia County that is irrigated from the Rio San Jose and the Rio Puerco. The data applied in calculating this ratio are listed in Table J.3.

The ratio is

$$\frac{47.5(4.3 \times 10^7) + 132.5(5.8 \times 10^6)}{47.5(4.3 \times 10^7) + 132.5(5.8 \times 10^6) + 45(8.7 \times 10^8)} = 0.067.$$

Then the equivalent land irrigated using surface water from the Rio San Jose and Rio Puerco in Valencia County is $11,234 (0.067) + 5,306 (0.5)(0.067) = 930$ hectares. Scaling this up to the assessment area size using the ratio of assessment area size to Valencia County size yields:

Table J.3 Stream data for Valencia County

Stream	Length Within Valencia County (km)	Annual Average Flow Rate (m ³ /yr)
Rio San Jose	132.5	5.8x10 ⁶
Rio Puerco	47.5	4.3x10 ⁷
Rio Grande	45	8.7x10 ⁸

$$930 \text{ hectares} \times \frac{19,037}{14,650} \times \frac{0.01 \text{ km}^2}{\text{hectares}} = 12.1 \text{ km}^2.$$

The surface water usage to irrigate this land (1.07 m/y) is $1.29 \times 10^{10} \text{ } \ell / \text{y}$ and the fraction of the river flow in the first order stream that the surface water irrigation represents is $1.29 \times 10^{10} / 4.26 \times 10^{10} = 0.30$.

For the Wyoming site, the area of land irrigated within the assessment area is given in Section 3.3.3 as 2,800 hectares (28 km²). This land is irrigated almost entirely with surface water (Loper, R., Soil Conservation Service, U.S. Department of Agriculture, Douglas, WY, 1979, personal communication). The average irrigation rate within the area is $0.588 \frac{\text{m}^3}{\text{m}^2 \cdot \text{y}}$ (WSG77).

Thus the estimated total irrigation water usage within the assessment area is

$$(0.588 \frac{\text{m}}{\text{y}})(2.8 \times 10^7 \text{ m}^2) = 1.65 \times 10^7 \frac{\text{m}^3}{\text{y}} = 1.65 \times 10^{10} \text{ } \ell / \text{yr}.$$

Then, the fraction of the first order stream flow that is used for irrigation is $1.65 \times 10^{10} / 5.6 \times 10^{10} = 0.29$.

J.14 Population Consuming Beef from Cattle Drinking Water Containing Radionuclides Discharged from the Uranium Mine

For the New Mexico site, information on beef consumption is taken from the USDA (DOA73), (Herman, J., Statistician-in-Charge, New Mexico Crop and Livestock Reporting Service, U.S. Department of Agriculture, Las Cruces, NM, 1979, personal communication). The term "beef" is a misnomer in that total red meat consumption is actually considered in these calculations. Since well over 50 percent of the red meat production is beef, the calculations are simplified by assuming that all meat production, excluding poultry, is beef for the purpose of this assessment. Table J.4 shows the computational procedure used to estimate the total meat production for Valencia County for 1977. Using this information, the total edible meat production for the assessment area can be estimated as

$$\frac{19,037 \text{ km}^2}{14,650 \text{ km}^2} (5.038 \times 10^6 \text{ kg/y}) = 6.547 \times 10^6 \text{ kg/y}.$$

It is estimated that about 50 percent of the water drunk by meat producing animals in the assessment area is surface water and 50 percent uncontaminated groundwater (Kaufmann, R., U.S. Environmental Protection Agency, Office of Radiation Programs, Las Vegas Facility, Las Vegas, NV, 1979, personal communication). Then the weight of edible meat from animals drinking surface water containing mine effluents is $3.273 \times 10^6 \text{ kg/y}$. Since it is estimated that an adult eats 85 kg/y of meat (Mo79), the number of persons eating meat from animals raised in the assessment area which drink water containing mine effluent is 38,510 persons.

For the Wyoming site, information on meat consumption was obtained from the USDA (DOA79). The information listed in Table J.5 shows the computational procedure used to estimate the meat production for Converse County for 1976. Thus, the total edible meat production for the assessment area can be estimated as

$$\frac{13,650 \text{ km}^2}{10,930 \text{ km}^2} (469,700 \frac{\text{kg}}{\text{y}}) = 5.866 \times 10^5 \frac{\text{kg meat}}{\text{y}}.$$

J.4 Estimation of meat production in Valencia County for 1977

Animal	1977 NM State Live Weight Slaughter ^(a) (kg)	Edible Fraction Live Weight ^(b)	1977 NM State Edible Meat, Com- puted from Col. 2 and 3 (kg)	Total Animals on Pasture Jan. 1, 1973 ^(c)		Ratio of Animals on Pasture, Valencia County+ New Mexico State (Computed from Col. 5 and 6)	Estimated Edible Meat Production for Valencia County (Computed from Col. 4 and 7) (kg/y)
	2	3	4	NM State	Valencia County	7	8
Cattle	260,813,700	.55	143,447,500	1,615,000	54,000	0.0334	4,791,000
Hogs	8,862,900	.65	5,760,880	63,000	1,500	0.0238	137,100
Sheep	4,487,460	.55	2,468,110	743,000	30,000	0.0404	99,700
Total Edible Meat Valencia County							5,027,800

(a) Herman, J., Statistician-in-Charge, New Mexico Crop and Livestock Reporting Service, U.S. Department of Agriculture, Las Cruces, NM, 1979, personal communication.

(b) Walsh, P., Statistician-in-Charge, Alabama Crop and Livestock Reporting Service, U.S. Department of Agriculture, 1979, personal communication.

(c) DOA73.

Table J.5 Estimation of meat production in Converse County, Wyoming for 1976

Animals	1976 Wyoming State Red Meat Production ^(a) (kg)	Total Animals on Pasture Jan. 1, 1976 ^(a)		Ratio of Animals on Pasture, Converse County + Wyoming State (Computed from Col. 3 and 4)	Estimated Edible Meat Production for Converse County (Computed from Col. 2 and 5) <u>(kg)</u> y
		Wyoming State	Converse County		
1	2	3	4	5	6
Cattle	--	1,580,000	74,000	0.0468	--
Red Meat Animals (Cattle, hogs, and sheep)	10,028,200	--	--	0.0468 ^(b)	469,300

(a) D0A79.

(b) Applied value for cattle since cattle furnish the majority of the red meat production for Converse County and for the State of Wyoming.

As for New Mexico, it was estimated that about 50 percent of the water drunk by meat producing animals in the assessment area is surface water and 50 percent uncontaminated groundwater (Loper, R., Soil Conservation Service, U.S. Department of Agriculture, Douglas, WY, 1979, personal communication). Thus, the estimated weight of edible meat from animals drinking surface water containing mine effluents is 293,500 kg meat/y. Using the 85 kg/y adult meat consumption rate (Mo79) used for New Mexico, the number of persons eating meat from animals raised in the assessment area which drink water containing mine effluent is 3,454 persons.

J.15 Radionuclide Releases

For both the New Mexico and the Wyoming sites, radionuclide releases are given for "total uranium" and radium-226 in Tables 3.44 and 3.25, respectively. The total uranium releases are kg per year. This "total uranium" will be almost totally U-238 by weight. For this reason, it is assumed that the "total uranium" release is entirely U-238. Further assumptions are that U-234 is in secular equilibrium with U-238 but that Th-230 precipitates out of the mine water. Also, it is assumed that Rn-222, Pb-214, Bi-214, Pb-210, and Po-210 are in secular equilibrium with Ra-226. The radionuclides Pa-234m, Pa-234, Po-218, At-218, Po-214, Tl-210, Bi-210 and Tl-206 are not included in the analysis because they are not dosimetrically significant or they have very low branching ratios.

The total uranium release rate for New Mexico is listed as 1,480 kg/y-mine. The conversion from kg to Ci for U-238 is 3.336×10^{-4} Ci/kg. Using this factor, the estimated release rate of U-238 is 0.494 Ci/y-mine. The release rate for Ra-226 is 0.0144 Ci/y-mine. The total uranium release rate at the Wyoming site is listed as 110 kg/y-mine. Using the conversion factor, this release rate can be stated as 0.0367 Ci/y-mine. The Ra-226 release rate is 0.0065 Ci/y-mine.

As discussed in Section 3.3.3.1.4, radium-226 is strongly sorbed onto stream sediments and is subject to precipitation. For these reasons, it is assumed that only 10% of the Ra-226 released in mine discharges is still available in surface water in the second and first order streams. Thus, the

"effective" annual release of Ra-226 is 10% of the actual releases in mine water. Using the assumptions regarding secular equilibrium stated above, the radionuclide release rates used in the analyses for both sites are listed in Table J.6.

J.16 Fish Concentration Factors

The fish concentration factors express the ratio of radionuclide level in freshwater fish (Ci/kg) per unit concentration in water (Ci/l). The values used for this parameter are suggested by Thompson (Th72) and are listed in Table J.7.

J.17 Fish Consumption and Air Inhalation Rates

The freshwater fish consumption for an individual is taken as 1.0 kg/y which is the value used in the report by the United Nations Scientific Committee on the Effects of Atomic Radiation (UN77). This value is not specifically stated in Regulatory Guide 1.109 (NRC77). The breathing rate for an individual of 8030 m³/y (Mo79) is used for this analysis. This value is in close agreement with the value of 8,000 m³/y listed in Regulatory Guide 1.109.

J.18 Stream Flow Rates

As was noted in Section J.2, second order stream flow rates were used in computing maximum individual dose equivalents. These flow rates are listed in Table J.2. For the New Mexico site, the second order stream flow rate is 5.83×10^9 l/y and for the Wyoming site it is 2.18×10^{10} l/y. First order stream flow rates are considered to be more representative for computations involving the population and an average individual. These flow rates are also listed in Table J.2 and are 4.26×10^{10} l/y for the New Mexico site and 5.64×10^{10} l/y for the Wyoming site.

J.19 Normalized Human Intake Rate Factors

The normalized human intake rate factors, RI_{np} , express the intake rate of radionuclide n by standard man from consumption of above-surface crops, milk, and beef for a continuous deposition rate to the surface. For this

Table J.6 Annual radionuclide release rates to
streams for active uranium mines

Nuclide	Release Rates for 1 Mine (Ci/y-mine)	
	New Mexico Site	Wyoming Site
U-238	0.494	0.0367
U-234	0.494	0.0367
Th-230	0	0
Ra-226	0.00144	0.00065
Rn-222	0.00144	0.00065
Pb-214	0.00144	0.00065
Bi-214	0.00144	0.00065
Pb-210	0.00144	0.00065
Po-210	0.00144	0.00065

Table J. 7 Freshwater fish concentration factors

Nuclide	Concentration Factor (Ci/Kg + Ci/l)
U-238	2
U-234	2
Ra-226	50
Rn-222	57
Pb-214	100
Bi-214	15
Pb-210	100
Po-210	500

pathway, the source of deposition to the surface is irrigation of farmland. Only the quantity of radionuclides taken up through the root systems of plants is considered in deriving these factors since essentially all irrigation in the assessment areas is ditch irrigation. The method used to calculate these factors was discussed in Section J.4 of this appendix and is taken from the AIRDOS-EPA computer code. The values for various parameters used in computing RI_{np} for above-surface crops, milk, and beef are discussed in the AIRDOS-EPA manual (Mo79). The normalized human intake rate factors are tabulated in Table J.8.

J.20 Persons Fed from Foodstuffs Raised on Irrigated Land and Irrigated Land Usage

The number of persons who can be fed from a unit area of irrigated land for above-surface crops, milk, and beef is determined from data contained in the AIRDOS-EPA computer code (Mo79). The values used for these analyses are, in units of persons fed/m², 3.69×10^{-3} for above-surface crops, 1.76×10^{-3} for milk, and 1.16×10^{-4} for beef. In both New Mexico and Wyoming, irrigated farmland is used for raising above surface foods for direct consumption by humans and for raising silage for consumption by both milk and beef cows. After telephone conversations with both the New Mexico and Wyoming county agents, the fractions of irrigated land supporting above-surface crops, dairy farming, and beef farming (see Table J.9) were determined (Romo, T., Valencia County Agent, Los Lunas, NM, 1980, personal communication and Henderson, F. Converse County Agent, Douglas, WY, 1980, personal communication).

J.21 Resuspension Factor

The irrigated areas within the assessment sites are assumed to be relatively large, uniformly contaminated areas. For this situation, the resuspension factor (RF) is defined as the ratio of air concentration above a surface to ground surface concentration. It can be shown that this ratio is $RF = \lambda_R / v_{gn}$. The resuspension rate constant, λ_R , can vary over a wide range of values between 10^{-7} and $10^{-11} \text{ sec}^{-1}$ (Ne78). Since resuspension for this analysis is confined to irrigated land which should have a relatively damp surface, it is believed that a low resuspension rate should be used. The resuspension rate constant chosen for this analysis is $\lambda_R = 10^{-11} \text{ sec}^{-1}$. The deposition velocity to the ground surface (v_{gn}) can vary from values as low as 0.001 m/sec to as high as 0.1 m/sec (S168). For generic analyses where

Table J.8 Normalized human intake rate factors for
radionuclide uptake via plant root systems

Nuclide	RI_{np} (Ci/day intake per Ci/m ² -day deposited)		
	Above Surface		
	Crops	Milk	Beef
U-238	3.21×10^{-2}	2.02×10^{-4}	1.75×10^{-6}
U-234	3.21×10^{-2}	2.02×10^{-4}	1.75×10^{-6}
Ra-226	1.90×10^{-2}	3.93×10^{-3}	2.53×10^{-3}
Rn-222	6.39×10^{-1}	4.61×10^{-1}	3.49×10^{-1}
Pb-214	3.68×10^{-2}	8.40×10^{-4}	6.66×10^{-3}
Bi-214	8.43×10^{-1}	1.51×10^{-2}	3.92×10^{-1}
Pb-210	2.98×10^{-2}	7.51×10^{-4}	5.23×10^{-3}
Po-210	1.97×10^{-3}	3.42×10^{-5}	1.88×10^{-3}

Table J.9 Irrigated land usage

Type of food	Fraction of irrigated land used to raise each type of food	
	New Mexico	Wyoming
Above-surface crops	0.70	0.10
Milk	0.15	0.45
Beef	0.15	0.45

the site specific value is not known, a commonly chosen value for v_{gn} is 0.01 m/sec, and this is the value used in this analysis. The value for RF inferred from the values chosen for λ_R and v_{gn} is

$$RF = \lambda_R / v_{gn} = \frac{10^{-11} \text{ sec}^{-1}}{0.01 \text{ m sec}^{-1}} = 10^{-9} / \text{m}$$

J.22 Soil Removal Rate Constant

The soil removal rate constant from available to unavailable soil (λ_{sn}) expresses the rate of movement of radionuclides from the plant root zone in soil to the soil below the root zone. The values used in this analysis are based upon a method described by Baes (Ba79). The soil removal rate constants were computed using the data suggested by Baes except that values for the distribution coefficients for the nuclides not discussed by Baes were taken from a report by the Arthur D. Little Company (EPA77). The values used for the soil removal rate constant in this analysis are listed in Table J.10.

J.23 Radionuclide Decay Constants

The radionuclide decay constants express the rate of radioactive decay for the nuclides considered in this analysis. The values of this parameter for these nuclides are listed in Table J.10 and are derived using the half-lives given in the Radiological Health Handbook (HEW70). Since secular equilibrium was assumed (see J.15), the radionuclide decay constant for Ra-226 was used for its short-lived daughter products.

J.24 Shielding and Occupancy Factor

The shielding and occupancy factor is used to account for shielding of persons by buildings during the time that they spend indoors. It is also used to account for time spent away from the radiation exposure area. The shielding and occupancy factor used in this analysis is 0.5 and is taken from Regulatory Guide 1.109 (NRC77).

Table J.10 Soil removal rate constants and radioactive decay constants

Nuclide	Soil Removal Rate Constants (y^{-1})	Radioactive Decay Constants, λ Dn (y^{-1})
U-238	2.58×10^{-4}	1.54×10^{-10}
U-234	2.58×10^{-4}	2.81×10^{-6}
Ra-226	7.74×10^{-3}	4.33×10^{-4}
Rn-222	4.93	6.62×10^1
Pb-214	1.94×10^{-4}	1.36×10^4
Bi-214	7.63×10^{-2}	1.85×10^4
Pb-210	1.94×10^{-4}	3.30×10^{-2}
Po-210	3.26×10^{-3}	1.83×10^6

Table J.11 Milk and beef concentration factors

Nuclide	Milk Concentration Factor (Ci/g milk per Ci/d intake)	Beef Concentration Factor (Ci/kg beef per Ci/day intake)
U-238	1.40×10^{-4}	1.60×10^{-6}
U-234	1.40×10^{-4}	1.60×10^{-6}
Ra-226	5.90×10^{-4}	5.00×10^{-4}
Rn-222	2.00×10^{-2}	2.00×10^{-2}
Pb-214	8.70×10^{-5}	9.10×10^{-4}
Bi-214	5.00×10^{-4}	1.70×10^{-2}
Pb-210	9.90×10^{-5}	9.10×10^{-4}
Po-210	1.20×10^{-4}	8.70×10^{-3}

J.25 Milk and Beef Ingestion Rates by Humans and Milk and Beef Cow Drinking Water Rates

The milk ingestion rate by humans is assumed to be 112 l/y which is the value used in AIRDOS-EPA (Mo79). In computing dose equivalents for the beef ingestion pathway, hogs and sheep are lumped into the beef pathway. This simplified the calculations, and it is believed that the assumption is reasonable since cattle account for well over 50 percent of the meat production in both the New Mexico and Wyoming assessment areas (DOA73, DOA79). The meat ingestion rate by humans is assumed to be 85 kg/y, which is the value listed for adults in AIRDOS-EPA (Mo79). This ingestion rate is employed in determining the number of persons who could be fed from meat produced within the assessment areas, as discussed in Section J.14.

The milk cow drinking water intake used in these calculations is 60 l/d; for beef cattle, the value is 50 l/d. Both values are the ones suggested in Regulatory Guide 1.109 (NRC77).

J.26 Radionuclide Concentration Factors for Milk and Beef

The radionuclide concentration factors for milk and beef express the concentration of radionuclides in milk or beef per unit daily intake of radionuclides by cattle drinking water. The values for these parameters are taken from AIRDOS-EPA (Mo79) and are listed in Table J.11.

J.27 Dosimetry Factors

Internal and external dosimetry factors are derived from the RADRISK data library which is being developed by Oak Ridge National Laboratory (Du80). The organs listed are considered to be the more dosimetrically significant organs. Breast dosimetry factors were used for the calculations for muscle since the muscle factors were not listed in RADRISK and since the breast and muscle factors should be similar in magnitude. The dosimetry factors tabulated as "weighted mean" were obtained by summing over organs the product of the organ dosimetry factors and a relative-risk weighting factor for the organ. These dosimetry factors are not the same as total-body dosimetry factors but are an attempt to express an overall dose that has been weighted for risk due to exposing each individual organ. Weighting factors were chosen to have a sum of 1.

For the inhalation pathway, the dosimetry factors incorporate the "Task Group Lung Model" (TGLD66). All nuclides are assumed to be of Class W solubility and an AMAD of $1.0\mu\text{m}$ is used. Class W solubility is assumed rather than Class Y because it is believed that in order for radioactive compounds to be soluble in mine water they must be in a more soluble chemical form than Class Y compounds. The dosimetry factors used in these assessments are listed in Table J.12.

J.28 Health Effects Conversion Factors

Health effects conversion factors for the internal and external pathways are derived from the RADRISK data library (Du80). Internal health effects conversion factors are needed for both inhalation and ingestion. For these internal pathways, the conversion factors are specific to each radionuclide and express the potential health impact per unit radionuclide intake. External health effects conversion factors are a function of organ but are radionuclide independent. The factors relate potential health impact to external radiation dose equivalent. Separate sets of health effect conversion factors are needed to estimate potential somatic health effects (fatal cancers) and potential genetic effects (genetic defects in the offspring of the exposed persons). The health effects conversion factors for the internal pathways are listed in Table J.13, and those for the external pathways are listed in Table J.14. Additional discussion of the philosophy of health impact determination used to obtain the data in the RADRISK data library is included in Section L.1 of Appendix L.

TABLE J.12
DOSE EQUIVALENT CONVERSION FACTORS

(INHALATION AND INGESTION=REM/CI INTAKE		AIR SUBMERSION=REM/Y PER CI/M**3					GROUND CONTAMINATION=REM/Y PER CI/M**2)						
NUCLIDE	PATHWAY	ORGAN											
		ENDOST. CELLS	RED MARROW	LUNG	LIVER	STOMACH WALL	LLI WALL	THYROID	KIDNEY	WEIGHTED MEAN	MUSCLE	VARIES	TESTES
U-238	INHALATION	1.29E+07	4.54E+05	8.40E+07	6.40E+04	8.11E+03	1.02E+05	6.30E+04	6.63E+06	2.48E+07	6.31E+04	6.00E+04	6.01E+04
	INGESTION	1.15E+07	4.02E+05	4.99E+03	5.30E+04	8.56E+03	1.40E+05	5.58E+04	5.90E+06	3.61E+05	5.59E+04	5.30E+04	5.34E+04
	EXT. AIR SUBMERSION	3.47E+02	3.08E+02	1.26E+02	2.86E-03	1.07E+02	8.18E+01	1.58E+02	9.06E+01	1.83E+02	2.37E+02	9.60E+01	2.62E+02
	EXT. GROUND CONTAM.	2.71E+01	2.32E+01	8.59E+00	2.97E+00	7.26E+00	8.41E+00	7.05E+00	3.04E+00	1.84E+01	3.29E+01	5.32E+00	3.89E+01
U-234	INHALATION	1.58E+07	4.66E+05	9.52E+07	7.04E+04	8.33E+03	8.19E+04	7.04E+04	7.46E+06	2.82E+07	7.04E+04	7.04E+04	7.04E+04
	INGESTION	1.41E+07	4.24E+05	5.53E+03	6.26E+04	9.63E+03	1.52E+05	6.26E+04	6.64E+06	4.19E+05	6.26E+04	6.26E+04	6.26E+04
	EXT. AIR SUBMERSION	1.08E+03	9.80E+02	4.68E+02	3.76E+02	3.72E+02	2.90E+02	6.18E+02	3.75E+02	5.91E+02	6.15E+02	2.96E+02	6.85E+02
	EXT. GROUND CONTAM.	4.82E+01	4.22E+01	1.77E+01	1.00E+01	1.45E+01	1.46E+01	1.85E+01	9.92E+00	3.06E+01	4.71E+01	1.07E+01	5.53E+01
RA-226	INHALATION	2.54E+07	1.46E+06	9.55E+07	5.97E+05	3.91E+03	1.81E+05	6.37E+05	5.98E+05	2.87E+07	6.37E+05	6.38E+05	6.37E+05
	INGESTION	2.26E+07	1.30E+06	1.91E+03	5.32E+05	5.45E+03	3.35E+05	5.67E+05	5.33E+05	8.07E+05	5.68E+05	5.69E+05	5.67E+05
	EXT. AIR SUBMERSION	5.87E+04	5.49E+04	3.37E+04	2.91E+04	2.49E+04	2.09E+04	3.81E+04	2.76E+04	3.65E+04	3.42E+04	2.00E+04	4.68E+04
	EXT. GROUND CONTAM.	1.30E+03	1.21E+03	7.43E+02	6.41E+02	5.49E+02	4.61E+02	8.41E+02	6.09E+02	8.05E+02	7.57E+02	4.41E+02	1.02E+03
RN-222	INHALATION	3.32E+01	5.46E+00	1.03E+03	7.32E+00	4.88E-01	1.43E-02	1.02E+00	4.46E+01	3.03E+02	1.02E+00	1.01E+00	1.00E+00
	INGESTION	3.40E+04	6.00E+03	1.17E+02	7.39E+03	1.48E+04	6.87E+05	4.50E+02	1.40E+04	2.61E+04	1.06E+03	5.62E+03	9.12E+02
	EXT. AIR SUBMERSION	2.59E+03	2.48E+03	2.00E+03	1.85E+03	2.12E+03	1.56E+03	1.77E+03	1.86E+03	2.09E+03	2.14E+03	7.96E+02	2.40E+03
	EXT. GROUND CONTAM.	5.33E+01	5.12E+01	4.12E+01	3.80E+01	4.36E+01	3.21E+01	3.65E+01	3.83E+01	4.30E+01	4.40E+01	1.64E+01	4.94E+01
PB-214	INHALATION	7.37E+03	1.15E+03	9.34E+04	1.49E+03	1.03E+02	3.02E+00	1.55E+02	9.11E+03	2.78E+04	1.56E+02	1.52E+02	1.50E+02
	INGESTION	2.17E+03	3.19E+02	7.70E+00	4.18E+02	3.14E+03	1.03E+02	2.51E+01	1.01E+03	2.79E+02	3.55E+01	6.44E+01	2.70E+01
	EXT. AIR SUBMERSION	1.86E+06	1.76E+06	1.25E+06	1.11E+06	1.11E+06	8.72E+05	1.23E+06	1.09E+06	1.32E+06	1.30E+06	6.14E+05	1.61E+06
	EXT. GROUND CONTAM.	4.02E+04	3.81E+04	2.68E+04	2.39E+04	2.39E+04	1.87E+04	2.66E+04	2.35E+04	2.84E+04	2.81E+04	1.32E+04	3.45E+04
BI-214	INHALATION	5.38E+02	1.35E+02	6.97E+04	8.44E+01	7.17E+01	2.83E+01	7.93E+01	8.42E+03	2.05E+04	8.00E+01	7.69E+01	7.62E+01
	INGESTION	2.37E+02	2.83E+01	7.57E+00	1.63E+01	3.18E+03	2.57E+01	4.17E+00	3.85E+02	1.54E+02	1.05E+01	2.02E+01	4.88E+00
	EXT. AIR SUBMERSION	9.43E+06	8.72E+06	8.28E+06	7.54E+06	7.92E+06	6.81E+06	7.87E+06	7.04E+06	8.33E+06	8.74E+06	6.93E+06	6.74E+06
	EXT. GROUND CONTAM.	1.65E+05	1.52E+05	1.44E+05	1.31E+05	1.38E+05	1.17E+05	1.35E+05	1.23E+05	1.45E+05	1.52E+05	1.17E+05	1.20E+05
PB-210	INHALATION	6.52E+06	3.81E+05	1.08E+07	2.58E+06	1.18E+03	4.56E+04	1.48E+05	2.30E+06	3.58E+06	1.48E+05	1.48E+05	1.48E+05
	INGESTION	5.80E+06	2.91E+05	9.05E+01	2.16E+06	2.04E+02	1.77E+04	8.17E+04	1.00E+06	3.39E+05	8.17E+04	8.18E+04	8.17E+04
	EXT. AIR SUBMERSION	1.32E+04	1.19E+04	4.94E+03	3.93E+03	4.19E+03	2.38E+03	7.34E+03	4.36E+03	6.47E+03	6.29E+03	3.97E+03	6.72E+03
	EXT. GROUND CONTAM.	4.42E+02	3.97E+02	1.66E+02	1.28E+02	1.46E+02	8.07E+01	2.42E+02	1.42E+02	2.22E+02	2.28E+02	1.32E+02	2.62E+02
PO-210	INHALATION	3.78E+05	8.20E+05	7.84E+07	2.52E+06	2.22E+03	8.84E+04	8.10E+05	1.46E+07	2.37E+07	8.10E+05	8.10E+05	8.10E+05
	INGESTION	2.44E+05	5.32E+05	1.15E-02	1.63E+06	4.52E+03	1.80E+05	5.26E+05	9.44E+06	5.67E+05	5.26E+05	5.26E+05	5.26E+05
	EXT. AIR SUBMERSION	5.26E+01	4.87E+01	4.33E+01	3.93E+01	4.11E+01	3.08E+01	3.52E+01	4.25E+01	4.39E+01	4.62E+01	2.38E+01	4.49E+01
	EXT. GROUND CONTAM.	1.02E+00	9.45E-01	8.40E-01	7.62E-01	7.97E-01	5.97E-01	6.82E-01	8.24E-01	8.51E-01	8.95E-01	4.62E-01	8.70E-01

Table J.13 Health effects conversion factors for internal pathways

Nuclide	Somatic		Genetic	
	(fatal cancers/Ci intake)		(genetic defects/Ci intake)	
	Inhalation	Ingestion	Inhalation	Ingestion
U-238	3.67×10^3	4.79×10^1	8.07×10^1	7.20×10^1
U-234	4.14×10^3	5.38×10^1	9.57×10^1	8.55×10^1
Ra-226	4.21×10^3	1.01×10^2	8.57×10^2	7.65×10^2
Rn-222	4.51×10^{-2}	5.64	1.49×10^{-3}	1.49
Pb-214	4.15	6.47×10^{-2}	2.30×10^{-1}	4.24×10^{-2}
Bi-214	3.04	3.34×10^{-2}	1.13×10^{-1}	7.81×10^{-3}
Pb-210	5.31×10^2	3.94×10^1	1.69×10^2	7.82×10^1
Po-210	3.50×10^3	9.67×10^1	1.21×10^3	7.84×10^2

Table J.14 Health effects conversion factors for external pathways

Organ	Somatic	Genetic
	(fatal cancers/rem)	(genetic defects/rem)
Endosteal	4.35×10^{-6}	0
Red Marrow	4.59×10^{-5}	0
Lung	8.59×10^{-5}	0
Liver	2.20×10^{-5}	0
Stomach Wall	1.23×10^{-5}	0
LLI Wall	9.81×10^{-6}	0
Thyroid	1.20×10^{-5}	0
Kidney	4.90×10^{-6}	0
Muscle	9.31×10^{-5}	0
Ovaries	2.46×10^{-6}	3.00×10^{-4}
Testes	2.46×10^{-6}	3.00×10^{-4}

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APPENDIX K
AIRBORNE PATHWAY MODELING

K.1 Generic Sites

Two generic sites were selected to represent locations where various uranium mines are present. The characteristics of these sites, as shown in Table K.1, were used in the AIRDOS-EPA code (Mo79).

Table K.1 Characteristics of the generic sites

	New Mexico (Ambrosia Lake)	Wyoming (Gas Hills)
Meteorological data: Stability Categories	Grants/Gnt-Milan (WBAN=93057) A-F	Casper (WBAN-24089) A-G
Period of Record:	54/01-54/12	67/01-71/12
Annual Rainfall:	20 cm	29 cm
Average Mixing Height:	800 m	500 m
Mean Ambient Temperature:	13.8°C	7.4°C
Atmospheric dispersion factors (Chi/Q) for maximum individual:		
gases	5.5 E-6 sec/m ³	2.5 E-6 sec/m ³
particulates	1.4 E-6 sec/m ³	1.0 E-6 sec/m ³
Population: (0-80 km):	3.60E+4 persons	1.43E+4 persons
Dairy Cattle (0-80 km):	2.30E+3 head	1.17E+3 head
Meat Animals (0-80 km):	8.31E+4 head	1.03E+5 head
Vegetable Crop Area: (0-80 km)	2.78E+3 ha	3.20E+3 ha

The model active and inactive underground mines were assumed to be situated at the New Mexico site (see Section 2). The Wyoming site was used for both the model active and inactive surface mines and the model in situ leach mine (see Section 2).

K.2 Meteorological Data

Joint frequency distributions by stability category were obtained from the National Climatic Center (NOAA-Asheville, NC). These distributions are identified in Table K.1 by location, stability category range, and period of record.

The average mixing height (Table K.1) is the distance between the ground surface and a stable layer of air where no further mixing occurs. This average was computed by determining the harmonic mean of the annual morning mixing height and the annual afternoon mixing height for the location (Ho72). The rainfall rate determines the value used for the scavenging coefficient. No attempt was made to be any more accurate than one significant figure for both average mixing height and scavenging coefficient. Both sites are relatively dry locations, as reflected by a scavenging coefficient of $2.0 \times 10^{-6} \text{ sec}^{-1}$. A dry deposition velocity of 1 cm/s was assigned to particulates, while radon was assumed to be non-depositing.

K.3 Population

The population data for both generic sites were generated by a computer program (At74) that uses an edited and compressed version of the 1970 United States Census Bureau's "Master Enumeration District List with Coordinates" containing housing and population counts for each census enumeration district (CED) and the geographic coordinates of the population centroid for the district. In the Standard Metropolitan Statistical Areas, the CED is usually a "block group" that consists of a physical city block. In other areas, the district used is called an "enumeration district", and it may cover several square miles in a rural area.

There are approximately 250,000 CED's in the United States and the average population is about 800. The position of the population centroid for each CED was marked on the district maps by the individual census official responsible for each district and is based only on his judgment from inspection of the population distribution on the map. The CED entries are sorted in ascending order by longitude on the final data tape.

K.4 Dairy and Meat Animals

Dairy cattle and meat animal distributions are part of the AIRDOS-EPA

input. A constant animal density is assumed. The animal densities are provided by state in Table K.2. These densities were derived from information developed by NRC (NRC75). Milk production density in units of liter/day-square mile was converted to number of dairy cattle/square kilometer by assuming a milk production rate of 11.0 liters/day per dairy cow. Meat production density in units of kilograms/day-square mile was changed to an equivalent number of meat animals/square kilometer by assuming a slaughter rate of 0.00381 day^{-1} and 200 kilograms of meat/animal slaughtered.

Table K.2 Animal and vegetable crop distributions
for use with AIRDOS-EPA

State	Dairy Cattle Density (No./km ²)	Meat Animal Density (No./km ²)	Vegetable Crop Fraction (Km ² /km ²)
New Mexico	1.14E-1	4.13	1.38E-3
Wyoming	5.79E-2	5.12	1.59E-3

K.5 Vegetable Crop Area

A certain fraction of the land within 80 km of the source is used for vegetable crop production, which is assumed to be uniformly distributed throughout the entire assessment area. Information on the vegetable production density in terms of kilograms(fresh weight)/day-square mile was obtained from NRC data (NRC75). The vegetable crop fractions (Table K.2) by state were computed from the production densities by assuming a production rate of 2 kilograms (fresh weight)/year-square meter (NRC77).

K.6 Food Intake

Table K.3 summarizes the ingestion values used for both generic sites for the maximum individual. These values are based on a USDA report (USDA72) for a rural farm situation. F1 and F2 are the percentages produced at the

individual's home and within the 80 km assessment area, respectively. The balance of the diet, F3, is considered to be imported from outside the assessment area with negligible radionuclide concentrations due to the assessed source. The F1 values are obtained by dividing the home-produced quantity by the quantity from all sources. The meat values include a combination of beef and pork. The vegetable values only include fresh vegetables.

Table K.3 Sources of food for the maximum individual (percent)

	F1	F2	F3
Vegetables	70.0	0.0	30.0
Meat	44.2	0.0	55.8
Milk	39.9	0.0	60.1

For population exposure estimates, the AIRDOS-EPA code determines the imported fraction needed to supply the nutritional requirements of the entire population within 80 km. The quantity of food that is not imported is assumed to be grown or produced throughout the entire assessment area and consumed by the population within the assessment area as an average value for the entire assessment area. The surplus food grown at a given site we assumed was shipped outside the assessment area. We did not calculate dose for this exported fraction.

The ingestion pathway is handled by the terrestrial model (NRC77) portion of the AIRDOS-EPA code. The input values shown in Table K.4 were used and are independent of location and radionuclide. Selected terrestrial pathway parameters, which are radionuclide dependent, are given in Table K.5.

K.7 AIRDOS-EPA Output

An example output of AIRDOS-EPA can be found in the AIRDOS-EPA manual (Mo79). Doses calculated by AIRDOS-EPA were not used in this report. Another code, DARTAB, performed dose and risk estimates based on air and ground concentrations and ingestion and inhalation intakes and working levels calculated by AIRDOS-EPA. An explanation of the DARTAB code can be found in

Table K.4 Selected Input Parameters to AIRDOS-EPA

SYMBOLIC VARIABLE	DESCRIPTION	VALUE
PR	Plume rise	0.0 m
PH	Release height	1.0 m
BRTHRT	Human breathing rate	$9.17 \times 10^5 \text{ cm}^3/\text{hr}$
T	Buildup time for surface deposition	source dependent
DD1	Fraction of radioactivity retained on leafy vegetables and produce after washing	1.0
TSUBH1	Time delay-ingestion of pasture grass by animals	0.0 hr
TSUBH2	Time delay-ingestion of stored feed by animals	$2.16 \times 10^3 \text{ hr}$
TSUBH3	Time delay-ingestion of leafy vegetables by man	$3.36 \times 10^2 \text{ hr}$
TSUBH4	Time delay-ingestion of produce by man	$3.36 \times 10^2 \text{ hr}$
LAMW	Removal rate constant for physical loss by weathering	$2.1 \times 10^{-3} \text{ hr}^{-1}$
TSUBE1	Period of exposure during growing season-pasture grass	$7.2 \times 10^2 \text{ hr}$
TSUBE2	Period of exposure during growing season-crops or leafy vegetables	$1.44 \times 10^3 \text{ hr}$
YSUBV1	Agricultural productivity by unit area (grass-cow-milk- man pathway)	0.28 kg/m^2
YSUBV2	Agricultural productivity by unit area (produce or leafy vegetables ingested by man)	0.716 kg/m^2
FSUBP	Fraction of year animals graze or pasture	0.40
FSUBS	Fraction of daily feed that is pasture grass when animals graze on pasture	0.43
QSUBF	Consumption rate of contaminated feed or forage by an animal (dry weight)	15.6 kg/day
TSUBF	Transport time from animal feed-milk-man	2.0 days

Table K.4 (Continued)

SYMBOLIC VARIABLE	DESCRIPTION	VALUE
UV	Rate of ingestion of produce by man	1.76E+2 kg/yr ^(a)
UM	Rate of ingestion of milk by man	1.12E+2 l/yr
UF	Rate of ingestion of meat by man	85.0 kg/yr
UL	Rate of ingestion of leafy vegetables by man	18.0 kg/yr
TSUBS	Average time from slaughter of meat animal to consumption	20.0 days
FSUBG	Fraction of produce ingested grown in garden of interest	1.0
FSUBL	Fraction of leafy vegetables grown in garden of interest	1.0
TSUBB	Period of long-term buildup for activity in soil	Same as T
P	Effective surface density of soil (dry weight) (assumes 15 cm plow layer)	2.15E+2 kg/m ²
TAUBEF	Fraction of meat producing herd slaughtered per day	3.81E-3 day ⁻¹
MSUBB	Muscle mass of meat producing animal at slaughter	2.0E+2 kg
VSUBM	Milk production of cow	11.0 l/day
R1	Fallout interception fraction for pasture	0.57
R2	Fallout interception for vegetable crops	0.20

^(a) This value, which was used in our analysis, is conservative because it includes fruit consumption. Without fruit consumption, the ingestion rate is 122 kg/yr.

Table K.5 Selected terrestrial pathway parameters by radionuclide

Radionuclide	Environmental Removal Rate (day ⁻¹)	B _{iv1} ^(a)	B _{iv2} ^(b)	Milk-transfer Coefficient F _m (day/l)	Meat-transfer Coefficient F _f (day/kg)
Uranium-238	7.06E-7	2.1E-2	4.2E-3	1.4E-4	1.6E-6
Uranium-235	7.06E-7	2.1E-2	4.2E-3	1.4E-4	1.6E-6
Uranium-234	7.06E-7	2.1E-2	4.2E-3	1.4E-4	1.6E-6
Thorium-232	5.61E-8	6.3E-3	3.5E-4	5.0E-6	1.6E-6
Thorium-230	5.61E-8	6.3E-3	3.5E-4	5.0E-6	1.6E-6
Thorium-228	5.61E-8	6.3E-3	3.5E-4	5.0E-6	1.6E-6
Actinium-228	2.12E-6	1.0E-2	2.5E-3	2.0E-5	1.6E-6
Radium-228	2.12E-5	1.0E-1	7.2E-2 ^(c)	5.9E-4	5.0E-4
Radium-226	2.12E-5	1.0E-1	7.2E-2 ^(c)	5.9E-4	5.0E-4
Radium-224	2.12E-5	1.0E-1	7.2E-2 ^(c)	5.9E-4	5.0E-4
Radon-222	0.0	0.0	0.0	0.0	0.0
Bismuth-214	2.09E-4	6.0E-1	1.5E-1	5.0E-4	1.7E-2
Bismuth-212	2.09E-4	6.0E-1	1.5E-1	5.0E-4	1.3E-2
Lead-214	5.31E-7	1.4E-1	4.8E-3	8.7E-5	9.1E-4
Lead-212	5.31E-7	1.4E-1	4.8E-3	8.7E-5	9.1E-4
Lead-210	5.31E-7	1.1E-1	3.9E-3	9.9E-5	9.1E-4
Polonium-210	8.93E-6	4.2E-3	2.6E-4	1.2E-4	8.7E-3
Thallium-208	9.84E-4	1.0E+0	2.5E-1	2.3E-2	4.0E-2

(a) $B_{iv1} = \frac{\text{Radionuclide concentration in entire above-ground portion of pasture grasses at maturity per unit dry weight}}{\text{Radionuclide concentration in soil per unit dry weight}}$
 $= \frac{\text{pCi/kg dry weight}}{\text{pCi/kg dry soil}}$

(b) $B_{iv2} = \frac{\text{Radionuclide concentration in edible portion of leafy vegetables and fresh produce at maturity per unit fresh weight}}{\text{Radionuclide concentration in soil per unit dry weight}}$
 $= \frac{\text{pCi/kg wet weight}}{\text{pCi/kg dry soil}}$

(c) This value, which was used in our analysis is conservatively high. Using data from DeBortoli (De72) for vegetables and grain products, a more realistic value would be 2.6E-3.

Appendix L. Concentrations and intakes computed by AIRDOS-EPA are by radionuclide and receptor location. Table K.6 lists values for effective radioactive decay constants, assumed for the plume (air) and ground surface, which influence the concentration and intake estimates. The values chosen for the decay constants attempt to account for daughter buildup in the air and on the ground surface.

Table K.6 Effective radioactive decay constants

Radionuclide	Decay constant in plume (day^{-1})	Decay constant on ground surface (day^{-1})
Uranium-238	4.25E-13	4.25E-13
Uranium-235	2.68E-12	2.68E-12
Uranium-234	7.77E-9	7.77E-9
Thorium-232	1.35E-13	1.35E-13
Thorium-230	2.47E-8	2.47E-8
Thorium-228	9.92E-4	1.35E-13
Actinium-228	2.72E+0	1.35E-13
Radium-228	3.30E-4	1.35E-13
Radium-226	1.19E-6	1.19E-6
Radium-224	1.89E-1	1.35E-13
Radon-222	1.81E-1	1.81E-1
Bismuth-214	1.81E-1	1.19E-6
Bismuth-212	1.35E-13	1.35E-13
Lead-214	1.81E-1	1.19E-6
Lead-212	1.35E-13	1.35E-13
Lead-210	8.52E-5	1.19E-6
Polonium-210	5.02E-3	1.19E-6
Thallium-208	1.35E-13	1.35E-13

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APPENDIX L

HEALTH RISK ASSESSMENT METHODOLOGY

L.1 Airborne Emissions

We estimated the health impact risks of airborne radionuclide emissions with the DARTAB (Be80) computer code using external exposure input data (air concentration and ground surface concentration) and internal intake input data (inhalation and ingestion) from the AIRDOS-EPA (Mo79) computer code. For radon-222 daughter calculations, we performed a working level calculation using the AIRDOS-EPA code and the working level exposures for each location were used as input by the DARTAB code. We assumed the fraction of equilibrium for the working level calculations to be 0.700 (Ge78). Doses calculated by AIRDOS-EPA were not used in DARTAB.

Tables L.1 and L.2 contain the data used by DARTAB in the health impact assessment, and Table L.3 is an example input data file for a DARTAB run. The variables are described in the DARTAB manual (Be80). We calculated dose rates and somatic health risks with DARTAB using a data base developed using the RADRISK (Du80) computer code. The dose conversion factors for each radionuclide are shown in Table L.1.

The doses calculated using DARTAB were not used in the risk calculations. We used the risk conversion factors in Table L.1 for this purpose. We calculated genetically significant doses for a 30-year exposure period (the mean years of life where gonadal doses are genetically significant). In calculating external dose rates from the ground surface, external dose conversion factors for the ground surface (see Table L.1), which are for a perfect plane surface, are multiplied by 0.5 to correct for the roughness of the soil surface. A weighted mean dose equivalent rate is calculated instead of a total body dose equivalent rate. Weighting factors are chosen to have a sum of one (see Table L.2). These weighted mean dose equivalent rates are presented for perspective purposes and are not used in the risk calculations.

The somatic risk conversion factors for Rn-222 and particulate radionuclides (see Table L.1) are calculated based on external exposures and internal intakes existing for the cohort lifetime (70.7565 years average

lifetime expectancy). When the exposure time for the calculated risks was less than the expected lifetime for an individual, we calculated the risk by multiplying the risk calculated by DARTAB with the ratio of the actual exposure time to the average lifetime expectancy for an individual (e.g., the DARTAB calculated risk is multiplied by $1/70.7565$ for a one-year exposure time). The risk conversion factor for Rn-222 in Table L.1 is for Rn-222 only and does not include the risk due to radon daughters. The risk due to radon daughters was calculated using a working level calculation and the fatal lung cancer risk conversion factor for a lifetime exposure given in Table L.2. The somatic health impact for the regional population (fatal cancers per year) is calculated at equilibrium for continuous exposure and is equal to the additional fatal cancers committed over all time per year of exposure.

Genetic effect risks (effects/birth) to the descendants of the exposed parent are calculated based on a 30-year exposure period. When the exposure time for the calculated risks was less than 30 years, we calculated the risk by multiplying the risk calculated by DARTAB with the ratio of the actual exposure time to 30 years (e.g., the DARTAB calculated risk is multiplied by $1/30$ for a one year exposure time). Since the presented genetic effect risk is to descendants of the exposed individual or individuals, one cannot add the individual somatic and genetic effect risks presented in this report. The genetic effects per year in the regional population due to radionuclide releases from the mine are calculated for an equilibrium exposure situation. The calculated genetic effects per year at equilibrium is equal to the genetic effects committed over all time from one year exposure since the total genetic damage expressed over all generations is equal to the value in each generation reached after prolonged continuous exposure (UN77). The genetic effects committed to the regional population are calculated using risk coefficients (see Table L.2) that are based on a genetically significant dose (GSD). The fraction of the population gonadal dose that is genetically significant is $30/70.7565$ where 30 is the mean individual reproductive life in years and 70.7565 is the average individual lifetime expectancy in years.

For each model uranium mine site, calculations are done separately for each mine source as well as for the total source term for the evaluated mine. The additional runs for each source allows us to identify the percentage contribution of each source to the total risk. The tables in Chapter 6

reflect results for the total mine source term and tables in Appendix L present the risk by source term as well as the total risk for each model uranium mine type. Tables L.4-L.6 contain individual fatal cancer risks and Tables L.7-L.9 contain genetic effect risks.

L.2 Aqueous Emissions

The health effects conversion factors used in the aquatic pathways are based on information contained in the RADRISK data library (Du80). The RADRISK data library is the data base used by the DARTAB computer code in computing the health impact of airborne releases. Thus, most of the philosophy of health impact determination discussed in Section L.1 above applies to aquatic releases as well as to airborne releases. The numerical values for the health effects conversion factors used for the aquatic releases are given in Appendix J along with additional discussions of the use of these factors in the aquatic calculations.

TABLE I.1
RADIONUCLIDE DOSE RATE AND HEALTH EFFECT RISK CONVERSION
FACTORS USED IN URANIUM MINE ASSESSMENTS

FOR NUCLIDE : U-238, RESP CLEARANCE CLASS=Y, PARTICLE SIZE=1.0 AMAD, FI=0.200E-02

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	9.64E-07	1.53E-07	3.34E-05	3.89E-06	3.08E+05	232.
ENDOST	1.96E-06	5.63E-06	6.52E-05	1.43E-04	3.47E+05	271.
PUL	1.37E-09	2.43E-09	1.62E-02	4.13E-02	1.26E+05	85.9
MUSCLE	7.06E-09	2.76E-08	2.53E-06	7.08E-07	2.37E+05	329.
LIVER	6.07E-09	2.62E-08	5.38E-06	6.71E-07	8.53E+04	29.7
S WALL	4.31E-08	1.81E-07	6.41E-06	1.67E-07	1.07E+05	72.6
PANCREAS	6.74E-09	2.76E-08	3.29E-06	7.08E-07	7.63E+04	57.6
LLI WALL	9.76E-06	7.94E-06	2.05E-04	4.71E-06	8.18E+04	84.1
KIDNEYS	5.94E-07	2.92E-06	1.70E-05	7.47E-05	9.06E+04	30.4
BL WALL	4.57E-09	1.50E-08	7.15E-07	3.85E-07	6.58E+04	21.4
ULI WALL	1.73E-06	2.64E-06	6.94E-05	1.61E-06	6.92E+04	22.8
SI WALL	1.58E-07	4.48E-07	1.22E-05	3.23E-07	6.22E+04	20.8
OVARIES	1.45E-08	2.63E-08	1.42E-06	6.71E-07	9.60E+04	53.2
TESTES	6.04E-09	2.64E-08	1.19E-06	6.75E-07	2.62E+05	389.
SPLEEN	6.59E-09	2.76E-08	3.13E-06	7.08E-07	8.91E+04	45.8
UTERUS	7.76E-09	2.77E-08	1.38E-06	7.07E-07	2.36E+04	7.70
THYMUS	6.12E-09	2.76E-08	4.80E-06	7.08E-07	9.25E+04	30.4
THYROID	6.01E-09	2.76E-08	2.16E-06	7.07E-07	1.58E+05	70.5
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	1.55E-07	7.20E-07	1.73E-05	1.55E-05	7.86E+06	1.17E+04
OVARIES	4.03E-07	7.17E-07	2.27E-05	1.54E-05	2.88E+06	1.60E+03
AVERAGE	2.79E-07	7.19E-07	2.00E-05	1.55E-05	5.37E+06	6.63E+03
FATAL CANCER RISK CONVERSION FACTORS FOR LIFETIME EXPOSURE						
CANCER	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	2.27E-07	7.62E-07	7.15E-06	1.77E-05	.100	7.55E-05
ENDOST	4.22E-08	1.22E-06	1.26E-06	2.78E-05	1.07E-02	8.34E-06
PULMONARY	5.33E-10	1.48E-08	6.32E-03	.238	7.67E-02	5.23E-05
BREAST	2.42E-09	1.02E-08	5.92E-07	2.24E-07	9.46E-02	1.31E-04
LIVER	8.26E-10	3.78E-08	4.73E-07	8.33E-07	1.33E-02	4.64E-06
ST WALL	3.72E-09	1.57E-07	4.72E-07	1.36E-07	9.28E-03	6.30E-06
PANCREAS	7.07E-10	3.10E-08	2.38E-07	6.83E-07	9.27E-03	7.00E-06
LLI WALL	6.77E-07	5.51E-06	1.32E-05	3.22E-06	5.68E-03	5.84E-06
KIDNEYS	1.89E-08	9.29E-07	4.49E-07	2.04E-05	3.14E-03	1.05E-06
BL WALL	1.42E-10	4.85E-09	1.41E-08	1.07E-07	2.28E-03	7.43E-07
ULI WALL	6.00E-08	9.16E-07	2.23E-06	5.48E-07	2.40E-03	7.91E-07
SI WALL	2.73E-09	7.78E-08	1.94E-07	5.40E-08	1.08E-03	3.61E-07
OVARIES	2.36E-10	4.21E-09	1.44E-08	9.26E-08	1.67E-03	9.23E-07
TESTES	9.20E-11	4.23E-09	1.12E-08	9.30E-08	4.55E-03	6.75E-06
SPLEEN	9.96E-11	4.43E-09	3.23E-08	9.75E-08	1.55E-03	7.95E-07
UTERUS	1.21E-10	4.43E-09	1.34E-08	9.75E-08	4.09E-04	1.34E-07
THYMUS	9.22E-11	4.43E-09	5.02E-08	9.76E-08	1.61E-03	5.27E-07
THYROID	4.61E-10	2.22E-09	1.19E-07	5.13E-08	1.34E-02	5.97E-06
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	8.37E-14	2.16E-11	6.00E-12	4.65E-10	1.61E+00	2.00E-03

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
- (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
- (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
- (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
- (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
- (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
- (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
- (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
- (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
- (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
- (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
- (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : U-234, RESP CLEARANCE CLASS=Y, PARTICLE SIZE=1.0 AMAD, F1=0.200E-02

DOSE CONVERSION FACTORS

ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	4.40E-09	2.07E-07	1.21E-07	5.26E-06	9.80E+05	422.
ENDOST	1.84E-08	7.03E-06	4.83E-07	1.78E-04	1.08E+06	482.
PUL	2.77E-11	2.76E-09	2.73E-04	4.70E-02	4.68E+05	177.
MUSCLE	2.70E-10	3.13E-08	1.09E-07	8.01E-07	6.15E+05	471.
LIVER	1.48E-10	3.13E-08	4.24E-08	8.01E-07	3.76E+05	100.
S WALL	4.98E-08	2.06E-07	5.34E-08	1.89E-07	3.72E+05	145.
PANCREAS	2.02E-10	3.13E-08	1.92E-08	8.01E-07	2.87E+05	115.
LLI WALL	2.19E-06	9.01E-06	1.28E-06	5.34E-06	2.90E+05	146.
KIDNEYS	8.63E-09	3.32E-06	2.24E-07	8.49E-05	3.75E+05	99.2
BL WALL	2.66E-10	1.71E-08	1.42E-09	4.37E-07	3.20E+05	82.1
ULI WALL	7.37E-07	3.00E-06	4.33E-07	1.82E-06	3.14E+05	81.7
SI WALL	1.31E-07	5.10E-07	7.77E-08	3.68E-07	3.07E+05	79.4
OVARIES	5.67E-09	3.13E-08	5.92E-09	8.01E-07	2.96E+05	107.
TESTES	1.30E-10	3.13E-08	2.18E-09	8.01E-07	6.85E+05	553.
SPLEEN	1.62E-10	3.13E-08	3.19E-08	8.01E-07	3.50E+05	112.
UTERUS	4.13E-10	3.13E-08	2.72E-09	8.01E-07	1.77E+05	43.7
THYMUS	9.68E-11	3.13E-08	3.45E-08	8.01E-07	3.35E+05	89.2
THYROID	8.99E-11	3.13E-08	6.47E-09	8.01E-07	6.18E+05	185.
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	3.62E-09	8.55E-07	4.92E-08	1.84E-05	2.06E+07	1.66E+04
OVARIES	1.70E-07	8.55E-07	1.55E-07	1.84E-05	8.88E+06	3.21E+03
AVERAGE	8.68E-08	8.55E-07	1.02E-07	1.84E-05	1.47E+07	9.90E+03

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION	INGESTION	INHALATION	INHALATION	AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	1.10E-09	1.02E-06	2.70E-08	2.37E-05	.319	1.37E-04
ENDOST	3.99E-10	1.52E-06	9.50E-09	3.48E-05	3.32E-02	1.48E-05
PULMONARY	1.39E-11	1.68E-08	1.15E-04	.270	.285	1.08E-04
BREAST	1.02E-10	1.15E-08	2.96E-08	2.54E-07	.245	1.88E-04
LIVER	2.16E-11	4.52E-08	4.54E-09	9.94E-07	5.87E-02	1.56E-05
ST WALL	4.32E-09	1.78E-07	3.93E-09	1.54E-07	3.23E-02	1.26E-05
PANCREAS	2.29E-11	3.51E-08	1.61E-09	7.73E-07	3.49E-02	1.40E-05
LLI WALL	1.52E-07	6.25E-06	8.77E-08	3.65E-06	2.01E-02	1.01E-05
KIDNEYS	2.75E-10	1.06E-06	6.09E-09	2.32E-05	1.30E-02	3.44E-06
BL WALL	9.08E-12	5.51E-09	3.90E-11	1.21E-07	1.11E-02	2.85E-06
ULI WALL	2.56E-08	1.04E-06	1.48E-08	6.22E-07	1.09E-02	2.84E-06
SI WALL	2.27E-09	8.85E-08	1.32E-09	6.14E-08	5.33E-03	1.38E-06
OVARIES	9.83E-11	5.02E-09	9.11E-11	1.10E-07	5.14E-03	1.86E-06
TESTES	2.12E-12	5.02E-09	2.97E-11	1.10E-07	1.19E-02	9.60E-06
SPLEEN	2.53E-12	5.02E-09	3.80E-10	1.10E-07	6.07E-03	1.94E-06
UTERUS	7.03E-12	5.02E-09	3.71E-11	1.10E-07	3.07E-03	7.58E-07
THYMUS	1.49E-12	5.02E-09	4.12E-10	1.10E-07	5.81E-03	1.55E-06
THYROID	7.02E-12	2.51E-09	4.24E-10	5.81E-08	5.23E-02	1.57E-05
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	2.61E-14	2.57E-11	3.06E-14	5.52E-10	4.41E+00	2.97E-03

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : TH-230, RESP CLEARANCE CLASS=Y, PARTICLE SIZE=1.0 AMAD, FI=0.200E-03

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	2.21E-08	5.79E-05	5.23E-06	1.42E-02	3.24E+06	884.
ENDOST	1.19E-07	9.09E-04	2.91E-05	.223	3.54E+06	977.
PUL	1.65E-10	5.01E-11	3.09E-04	4.60E-02	1.66E+06	438.
MUSCLE	1.11E-09	2.28E-07	3.01E-07	5.70E-05	1.83E+06	633.
LIVER	3.49E-09	1.09E-06	9.01E-07	2.73E-04	1.40E+06	339.
S WALL	5.76E-08	1.99E-07	1.00E-07	1.29E-07	1.30E+06	353.
PANCREAS	1.10E-09	2.28E-07	2.64E-07	5.70E-05	1.03E+06	274.
LLI WALL	2.52E-06	8.83E-06	1.52E-06	5.18E-06	9.88E+05	275.
KIDNEYS	1.04E-09	2.28E-07	2.17E-07	5.70E-05	1.38E+06	335.
BL WALL	1.09E-09	1.14E-07	9.59E-08	2.85E-05	1.17E+06	284.
ULI WALL	8.45E-07	2.94E-06	5.24E-07	1.73E-06	1.20E+06	289.
SI WALL	1.49E-07	4.97E-07	1.16E-07	3.03E-07	1.16E+06	280.
OVARIES	6.69E-09	2.28E-07	1.98E-07	5.70E-05	1.04E+06	270.
TESTES	8.91E-10	2.28E-07	1.83E-07	5.70E-05	2.14E+06	769.
SPLEEN	9.95E-10	2.28E-07	2.62E-07	5.70E-05	1.32E+06	329.
UTERUS	1.78E-09	2.28E-07	1.88E-07	5.70E-05	7.27E+05	171.
THYMUS	7.83E-10	2.28E-07	2.88E-07	5.70E-05	1.29E+06	310.
THYROID	7.74E-10	2.28E-07	2.08E-07	5.70E-05	2.16E+06	544.
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	2.38E-08	6.20E-06	3.96E-06	1.30E-03	6.42E+07	2.31E+04
OVARIES	1.97E-07	6.20E-06	4.24E-06	1.30E-03	3.12E+07	8.10E+03
AVERAGE	1.11E-07	6.20E-06	4.10E-06	1.30E-03	4.77E+07	1.56E+04

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	4.76E-09	2.39E-04	1.00E-06	5.36E-02	1.05	2.88E-04
ENDOST	2.22E-09	1.70E-04	4.91E-07	3.76E-02	.109	3.01E-05
PULMONARY	6.15E-11	3.05E-10	1.30E-04	.265	1.01	2.66E-04
BREAST	3.91E-10	8.35E-08	8.53E-08	1.80E-05	.730	2.53E-04
LIVER	4.98E-10	1.57E-06	1.09E-07	3.39E-04	.219	5.29E-05
ST WALL	5.00E-09	1.72E-07	6.47E-09	1.09E-07	.113	3.06E-05
PANCREAS	1.18E-10	2.54E-07	2.33E-08	5.49E-05	.125	3.33E-05
LLI WALL	1.75E-07	6.13E-06	1.02E-07	3.55E-06	6.86E-02	1.91E-05
KIDNEYS	3.21E-11	7.26E-08	5.52E-09	1.57E-05	4.79E-02	1.16E-05
BL WALL	3.59E-11	3.63E-08	2.48E-09	7.85E-06	4.06E-02	9.86E-06
ULI WALL	2.93E-08	1.02E-06	1.75E-08	5.92E-07	4.16E-02	1.00E-05
SI WALL	2.58E-09	8.62E-08	1.73E-09	5.17E-08	2.01E-02	4.86E-06
OVARIES	1.14E-10	3.63E-08	2.59E-09	7.85E-06	1.80E-02	4.68E-06
TESTES	1.40E-11	3.63E-08	2.42E-09	7.85E-06	3.71E-02	1.33E-05
SPLEEN	1.51E-11	3.63E-08	3.28E-09	7.85E-06	2.29E-02	5.71E-06
UTERUS	2.94E-11	3.63E-08	2.50E-09	7.85E-06	1.26E-02	2.97E-06
THYMUS	1.19E-11	3.63E-08	3.65E-09	7.85E-06	2.24E-02	5.38E-06
THYROID	5.91E-11	1.82E-08	1.43E-08	4.14E-06	.183	4.61E-05
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	3.31E-14	1.86E-10	1.23E-12	3.90E-08	1.43E+01	4.68E-03

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : RA-226, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, F1=0.200E+00

DOSE CONVERSION FACTORS

ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	2.54E-05	6.40E-05	2.84E-05	7.18E-05	5.49E+07	1.21E+04
ENDOST	5.07E-05	1.13E-03	5.69E-05	1.27E-03	5.87E+07	1.30E+04
PUL	1.91E-06	.0	6.33E-06	4.77E-03	3.37E+07	7.43E+03
MUSCLE	2.27E-06	2.83E-05	2.50E-06	3.17E-05	3.42E+07	7.57E+03
LIVER	1.51E-06	2.65E-05	1.70E-06	2.98E-05	2.91E+07	6.41E+03
S WALL	1.30E-06	2.08E-07	1.49E-06	1.21E-07	2.49E+07	5.49E+03
PANCREAS	2.14E-06	2.83E-05	2.41E-06	3.17E-05	2.19E+07	4.84E+03
LLI WALL	4.15E-05	1.47E-05	2.38E-05	7.84E-06	2.09E+07	4.61E+03
KIDNEYS	2.01E-06	2.65E-05	2.21E-06	2.98E-05	2.76E+07	6.09E+03
BL WALL	1.58E-06	1.41E-05	1.56E-06	1.59E-05	2.42E+07	5.35E+03
ULI WALL	8.07E-06	3.58E-06	5.57E-06	1.94E-06	2.69E+07	5.94E+03
SI WALL	2.58E-06	4.63E-07	2.42E-06	2.59E-07	2.60E+07	5.74E+03
OVARIES	2.96E-06	2.83E-05	2.68E-06	3.17E-05	2.00E+07	4.41E+03
TESTES	1.66E-06	2.83E-05	1.80E-06	3.17E-05	4.68E+07	1.03E+04
SPLEEN	1.56E-06	2.65E-05	1.75E-06	2.98E-05	2.95E+07	6.49E+03
UTERUS	1.75E-06	2.83E-05	1.73E-06	3.17E-05	2.13E+07	4.68E+03
THYMUS	1.35E-06	2.83E-05	1.58E-06	3.17E-05	2.89E+07	6.36E+03
THYROID	1.49E-06	2.83E-05	1.69E-06	3.17E-05	3.81E+07	8.41E+03
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	2.69E-05	7.65E-04	2.82E-05	8.57E-04	1.40E+09	3.09E+05
OVARIES	6.17E-05	7.65E-04	5.00E-05	8.57E-04	6.00E+08	1.32E+05
AVERAGE	4.43E-05	7.65E-04	3.91E-05	8.57E-04	1.00E+09	2.21E+05

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	5.77E-06	3.37E-04	6.43E-06	3.78E-04	17.9	3.94E-03
ENDOST	1.05E-06	2.34E-04	1.18E-06	2.63E-04	1.81	4.00E-04
PULMONARY	6.95E-07	.0	3.32E-06	2.90E-02	20.5	4.52E-03
BREAST	5.48E-07	1.04E-05	5.95E-07	1.16E-05	13.6	3.02E-03
LIVER	1.42E-07	3.80E-05	1.60E-07	4.25E-05	4.54	1.00E-03
ST WALL	7.19E-08	1.80E-07	8.39E-08	1.04E-07	2.16	4.76E-04
PANCREAS	1.56E-07	3.15E-05	1.75E-07	3.53E-05	2.66	5.88E-04
LLI WALL	2.82E-06	1.02E-05	1.59E-06	5.43E-06	1.45	3.20E-04
KIDNEYS	4.20E-08	8.45E-06	4.57E-08	9.46E-06	.958	2.11E-04
BL WALL	3.75E-08	4.50E-06	3.46E-08	5.04E-06	.840	1.86E-04
ULI WALL	2.59E-07	1.24E-06	1.69E-07	6.72E-07	.934	2.06E-04
SI WALL	3.36E-08	8.03E-08	2.93E-08	4.48E-08	.451	9.96E-05
OVARIES	3.81E-08	4.50E-06	3.16E-08	5.04E-06	.347	7.65E-05
TESTES	1.77E-08	4.50E-06	1.86E-08	5.04E-06	.812	1.79E-04
SPLEEN	1.64E-08	4.22E-06	1.83E-08	4.73E-06	.512	1.13E-04
UTERUS	2.09E-08	4.50E-06	1.94E-08	5.04E-06	.370	8.12E-05
THYMUS	1.38E-08	4.50E-06	1.67E-08	5.04E-06	.501	1.10E-04
THYROID	8.46E-08	2.26E-06	9.60E-08	2.53E-06	3.23	7.12E-04
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	1.33E-11	2.29E-08	1.17E-11	2.56E-08	3.00E+02	6.62E-02

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : RN-222, RESP CLEARANCE CLASS=Y, PARTICLE SIZE=1.0 AMAD, FI=0.0

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	.0	.0	4.29E-11	2.71E-10	2.48E+06	512.
ENDOST	.0	.0	4.84E-11	1.66E-09	2.59E+06	533.
PUL	.0	.0	2.67E-09	5.14E-08	2.00E+06	412.
MUSCLE	.0	.0	3.29E-11	4.96E-11	2.14E+06	440.
LIVER	.0	.0	9.80E-11	3.61E-10	1.85E+06	380.
S WALL	.0	.0	2.08E-10	1.40E-11	2.12E+06	436.
PANCREAS	.0	.0	6.38E-11	4.96E-11	1.19E+06	246.
LLI WALL	.0	.0	1.02E-11	2.05E-13	1.56E+06	321.
KIDNEYS	.0	.0	2.37E-10	2.22E-09	1.86E+06	383.
BL WALL	.0	.0	7.87E-12	2.48E-11	1.71E+06	352.
ULI WALL	.0	.0	4.60E-11	2.22E-12	1.74E+06	359.
SI WALL	.0	.0	8.80E-11	6.38E-12	1.56E+06	322.
OVARIES	.0	.0	1.51E-11	4.96E-11	7.96E+05	164.
TESTES	.0	.0	7.59E-12	4.96E-11	2.40E+06	494.
SPLEEN	.0	.0	9.53E-11	6.08E-10	2.27E+06	468.
UTERUS	.0	.0	1.60E-11	4.96E-11	1.44E+06	296.
THYMUS	.0	.0	8.27E-11	4.96E-11	1.31E+06	270.
THYROID	.0	.0	2.67E-11	4.96E-11	1.77E+06	365.
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	.0	.0	.0	.0	7.20E+07	1.48E+04
OVARIES	.0	.0	.0	.0	2.39E+07	4.92E+03
AVERAGE	.0	.0	.0	.0	4.79E+07	9.87E+03

FATAL CANCER RISK CONVERSION FACTORS FOR LIFETIME EXPOSURE						
CANCER	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	.0	.0	1.39E-11	1.75E-09	.807	1.67E-04
ENDOST	.0	.0	1.48E-12	5.05E-10	7.97E-02	1.64E-05
PULMONARY	.0	.0	1.63E-09	3.13E-07	1.22	2.51E-04
BREAST	.0	.0	1.31E-11	1.98E-11	.854	1.76E-04
LIVER	.0	.0	1.53E-11	5.63E-10	.289	5.93E-05
ST WALL	.0	.0	1.81E-11	1.22E-11	.184	3.78E-05
PANCREAS	.0	.0	7.74E-12	6.02E-11	.145	2.99E-05
LLI WALL	.0	.0	7.07E-13	1.42E-13	.108	2.23E-05
KIDNEYS	.0	.0	8.23E-12	7.70E-10	6.45E-02	1.33E-05
BL WALL	.0	.0	2.73E-13	8.60E-12	5.93E-02	1.22E-05
ULI WALL	.0	.0	1.60E-12	7.71E-13	6.04E-02	1.25E-05
SI WALL	.0	.0	1.53E-12	1.11E-12	2.71E-02	5.59E-06
OVARIES	.0	.0	2.62E-13	8.60E-12	1.38E-02	2.85E-06
TESTES	.0	.0	1.32E-13	8.60E-12	4.16E-02	8.57E-06
SPLEEN	.0	.0	1.65E-12	1.06E-10	3.94E-02	8.12E-06
UTERUS	.0	.0	2.77E-13	8.60E-12	2.50E-02	5.14E-06
THYMUS	.0	.0	1.43E-12	8.60E-12	2.27E-02	4.68E-06
THYROID	.0	.0	2.26E-12	4.20E-12	.150	3.09E-05
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	0.00E-01	0.00E-01	0.00E-01	0.00E-01	1.44E+01	2.96E-03

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICO CURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICRO CURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICRO CURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICO CURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICRO CURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICRO CURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICO CURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICO CURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICO CURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICO CURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICRO CURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICRO CURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : PB-214, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, FI=0.200E+00

DOSE CONVERSION FACTORS

ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	1.46E-08	1.52E-08	9.59E-09	5.70E-08	1.76E+09	3.81E+05
ENDOST	7.30E-09	1.08E-07	1.10E-08	3.68E-07	1.86E+09	4.02E+05
PUL	7.70E-09	.0	6.07E-07	.0	1.25E+09	2.68E+05
MUSCLE	1.15E-08	1.20E-09	7.21E-09	7.45E-09	1.30E+09	2.81E+05
LIVER	1.82E-08	2.00E-08	2.19E-08	7.32E-08	1.11E+09	2.39E+05
S WALL	1.40E-06	8.68E-08	4.60E-08	2.84E-09	1.11E+09	2.39E+05
PANCREAS	6.50E-08	1.20E-09	1.40E-08	7.45E-09	7.90E+08	1.69E+05
LLI WALL	5.57E-08	2.36E-09	2.16E-09	4.31E-11	8.72E+08	1.87E+05
KIDNEYS	2.86E-08	4.89E-08	5.05E-08	4.53E-07	1.09E+09	2.35E+05
BL WALL	1.32E-08	6.00E-10	1.55E-09	3.72E-09	9.80E+08	2.10E+05
ULI WALL	3.69E-07	2.44E-08	9.86E-09	4.70E-10	1.05E+09	2.25E+05
SI WALL	8.01E-07	6.12E-08	1.89E-08	1.36E-09	9.69E+08	2.08E+05
OVARIES	4.04E-08	1.20E-09	3.04E-09	7.45E-09	6.14E+08	1.32E+05
TESTES	3.02E-09	1.20E-09	1.40E-09	7.45E-09	1.61E+09	3.45E+05
SPLEEN	3.84E-08	1.13E-09	1.16E-08	7.21E-09	1.26E+09	2.70E+05
UTERUS	3.52E-08	1.20E-09	3.14E-09	7.45E-09	8.45E+08	1.81E+05
THYMUS	2.88E-09	1.20E-09	1.86E-08	7.45E-09	9.40E+08	2.02E+05
THYROID	1.06E-09	1.20E-09	5.81E-09	7.45E-09	1.23E+09	2.66E+05
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	9.05E-08	3.59E-08	4.20E-08	2.23E-07	4.83E+10	1.04E+07
OVARIES	1.21E-06	3.59E-08	9.11E-08	2.23E-07	1.84E+10	3.96E+06
AVERAGE	6.52E-07	3.59E-08	6.65E-08	2.23E-07	3.34E+10	7.16E+06

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION	INHALATION	AIR	GROUND		
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	4.73E-09	9.72E-08	3.11E-09	3.69E-07	573.	.124
ENDOST	2.22E-10	3.20E-08	3.34E-10	1.12E-07	57.3	1.24E-02
PULMONARY	4.68E-09	.0	3.69E-07	2.82E-05	760.	.163
BREAST	4.58E-09	4.78E-10	2.88E-09	2.97E-09	519.	.112
LIVER	2.83E-09	3.11E-08	3.41E-09	1.14E-07	173.	3.73E-02
ST WALL	1.22E-07	7.53E-08	3.99E-09	2.46E-09	96.3	2.07E-02
PANCREAS	7.90E-09	1.45E-09	1.70E-09	9.04E-09	95.9	2.05E-02
LLI WALL	3.87E-09	1.64E-09	1.50E-10	2.99E-11	60.5	1.30E-02
KIDNEYS	9.94E-10	1.70E-08	1.75E-09	1.57E-07	37.8	8.16E-03
BL WALL	4.59E-10	2.08E-10	5.39E-11	1.29E-09	34.0	7.29E-03
ULI WALL	1.28E-08	8.47E-09	3.42E-10	1.63E-10	36.4	7.81E-03
SI WALL	1.39E-08	1.06E-08	3.28E-10	2.36E-10	16.8	3.61E-03
OVARIES	7.01E-10	2.08E-10	5.27E-11	1.29E-09	10.7	2.29E-03
TESTES	5.23E-11	2.08E-10	2.43E-11	1.29E-09	27.9	5.99E-03
SPLEEN	6.67E-10	1.96E-10	2.02E-10	1.25E-09	21.9	4.68E-03
UTERUS	6.11E-10	2.08E-10	5.45E-11	1.29E-09	14.7	3.14E-03
THYMUS	5.00E-11	2.08E-10	3.23E-10	1.29E-09	16.3	3.51E-03
THYROID	8.95E-11	1.01E-10	4.92E-10	6.30E-10	104.	2.25E-02
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	1.95E-13	1.08E-12	2.00E-14	6.70E-12	1.00E+04	2.14E+00

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : BI-214, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, FI=0.500E-01

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	6.93E-09	1.07E-09	4.37E-09	6.52E-09	8.72E+09	1.52E+06
ENDOST	3.38E-09	1.17E-08	3.65E-09	2.67E-08	9.43E+09	1.65E+06
PUL	6.37E-09	6.00E-11	3.18E-07	.0	8.28E+09	1.44E+06
MUSCLE	7.02E-09	1.72E-10	4.41E-09	3.78E-09	8.74E+09	1.52E+06
LIVER	9.67E-09	3.33E-10	7.59E-09	3.84E-09	7.54E+09	1.31E+06
S WALL	1.06E-06	1.06E-07	1.89E-08	2.64E-09	7.92E+09	1.38E+06
PANCREAS	5.60E-08	1.72E-10	8.35E-09	3.78E-09	7.94E+09	1.36E+06
LLI WALL	1.76E-08	4.07E-10	9.08E-10	1.37E-09	6.81E+09	1.17E+06
KIDNEYS	1.69E-08	1.84E-08	4.36E-08	4.19E-07	7.04E+09	1.23E+06
BL WALL	5.89E-09	1.16E-10	8.13E-10	2.57E-09	7.40E+09	1.29E+06
ULI WALL	9.75E-08	5.95E-09	2.96E-09	1.44E-09	8.83E+09	1.51E+06
SI WALL	2.89E-07	2.81E-08	4.95E-09	1.71E-09	6.77E+09	1.17E+06
OVARIES	1.68E-08	1.72E-10	1.35E-09	3.78E-09	6.93E+09	1.17E+06
TESTES	1.44E-09	1.72E-10	6.48E-10	3.78E-09	6.74E+09	1.20E+06
SPLEEN	3.22E-08	1.66E-10	7.27E-09	3.79E-09	8.18E+09	1.44E+06
UTERUS	1.52E-08	1.72E-10	1.43E-09	3.78E-09	6.24E+09	1.08E+06
THYMUS	2.37E-09	1.72E-10	1.22E-08	3.78E-09	7.48E+09	1.27E+06
THYROID	7.26E-10	1.72E-10	3.70E-09	3.78E-09	7.87E+09	1.35E+06
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	4.31E-08	5.08E-09	1.94E-08	1.13E-07	2.02E+11	3.60E+07
OVARIES	5.03E-07	5.08E-09	4.06E-08	1.13E-07	2.08E+11	3.51E+07
AVERAGE	2.73E-07	5.08E-09	3.00E-08	1.13E-07	2.05E+11	3.56E+07

FATAL CANCER RISK CONVERSION FACTORS FOR LIFETIME EXPOSURE						
CANCER	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	2.25E-09	5.53E-09	1.41E-09	4.09E-08	2.84E+03	.495
ENDOST	1.02E-10	2.60E-09	1.10E-10	7.14E-09	290.	5.08E-02
PULMINARY	3.87E-09	3.65E-10	1.93E-07	2.11E-05	5.04E+03	.876
BREAST	2.80E-09	6.77E-11	1.76E-09	1.51E-09	3.49E+03	.607
LIVER	1.51E-09	4.41E-10	1.18E-09	5.91E-09	1.18E+03	.205
ST WALL	9.21E-08	9.19E-08	1.64E-09	2.29E-09	687.	.120
PANCREAS	6.80E-09	2.06E-10	1.01E-09	4.58E-09	964.	.165
LLI WALL	1.22E-09	2.82E-10	6.30E-11	9.54E-10	473.	8.12E-02
KIDNEYS	5.85E-10	6.39E-09	1.51E-09	1.45E-07	244.	4.27E-02
BL WALL	2.04E-10	3.99E-11	2.82E-11	8.93E-10	257.	4.48E-02
ULI WALL	3.38E-09	2.07E-09	1.03E-10	5.00E-10	306.	5.24E-02
SI WALL	5.01E-09	4.88E-09	8.59E-11	2.96E-10	117.	2.03E-02
OVARIES	2.91E-10	2.94E-11	2.35E-11	6.55E-10	120.	2.03E-02
TESTES	2.49E-11	2.94E-11	1.12E-11	6.55E-10	117.	2.08E-02
SPLEEN	5.59E-10	2.85E-11	1.26E-10	6.57E-10	142.	2.50E-02
UTERUS	2.64E-10	2.94E-11	2.47E-11	6.55E-10	108.	1.87E-02
THYMUS	4.11E-11	2.94E-11	2.12E-10	6.55E-10	130.	2.20E-02
THYROID	6.14E-11	1.44E-11	3.13E-10	3.20E-10	666.	.114
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	8.19E-14	1.53E-13	9.00E-15	3.39E-12	6.15E+04	1.07E+01

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

L-11

FOR NUCLIDE : PB-210, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, F1=0.200

DOSE CONVERSION FACTORS

ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	5.28E-05	1.19E-05	5.92E-05	1.61E-05	1.19E+07	3.97E+03
ENDOST	1.42E-04	2.83E-04	1.59E-04	3.18E-04	1.32E+07	4.42E+03
PUL	9.05E-08	.0	3.01E-04	5.25E-04	4.94E+06	1.66E+03
MUSCLE	1.54E-06	4.01E-06	1.78E-06	7.29E-06	6.29E+06	2.28E+03
LIVER	3.90E-05	1.06E-04	4.37E-05	1.27E-04	3.93E+06	1.28E+03
S WALL	2.04E-07	2.71E-13	1.07E-06	5.42E-09	4.19E+06	1.46E+03
PANCREAS	1.58E-06	4.01E-06	1.82E-06	7.29E-06	2.60E+06	884.
LLI WALL	1.75E-05	1.06E-08	4.04E-05	2.60E-07	2.38E+06	807.
KIDNEYS	1.82E-05	4.93E-05	2.35E-05	1.14E-04	4.36E+06	1.42E+03
BL WALL	7.61E-07	2.00E-06	8.68E-07	3.65E-06	3.09E+06	1.01E+03
ULI WALL	3.73E-06	7.55E-10	1.38E-05	7.79E-08	3.11E+06	1.01E+03
SI WALL	4.73E-07	1.02E-11	2.43E-06	1.24E-08	2.71E+06	883.
OVARIES	1.56E-06	4.01E-06	1.77E-06	7.29E-06	3.97E+06	1.32E+03
TESTES	1.50E-06	4.01E-06	1.72E-06	7.29E-06	6.72E+06	2.62E+03
SPLEEN	1.47E-06	3.81E-06	1.69E-06	8.72E-05	3.74E+06	1.23E+03
UTERUS	1.52E-06	4.01E-06	1.74E-06	7.29E-06	1.12E+06	366.
THYMUS	1.52E-06	4.01E-06	1.77E-06	7.29E-06	4.41E+06	1.44E+03
THYROID	1.53E-06	4.01E-06	1.75E-06	7.29E-06	7.34E+06	2.42E+03
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	2.93E-05	7.79E-05	3.37E-05	1.69E-04	2.02E+08	7.86E+04
OVARIES	3.05E-05	7.79E-05	3.48E-05	1.69E-04	1.19E+08	3.96E+04
AVERAGE	2.99E-05	7.79E-05	3.43E-05	1.69E-04	1.60E+08	5.91E+04

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	1.26E-05	5.81E-05	1.41E-05	8.32E-05	3.87	1.29E-03
ENDOST	3.11E-06	6.15E-05	3.47E-06	6.91E-05	.406	1.36E-04
PULMONARY	3.59E-08	.0	1.83E-04	3.19E-03	3.01	1.01E-03
BREAST	4.36E-07	1.13E-06	5.06E-07	2.36E-06	2.51	9.10E-04
LIVER	4.33E-06	1.17E-04	4.84E-06	1.44E-04	.614	2.00E-04
ST WALL	1.64E-08	2.35E-13	9.02E-08	4.63E-09	.364	1.27E-04
PANCREAS	1.36E-07	3.45E-06	1.57E-07	7.17E-06	.316	1.07E-04
LLI WALL	1.22E-06	7.34E-09	2.77E-06	1.78E-07	.165	5.60E-05
KIDNEYS	4.51E-07	1.21E-05	6.08E-07	3.36E-05	.151	4.93E-05
BL WALL	1.88E-08	4.93E-07	2.15E-08	1.02E-06	.107	3.50E-05
ULI WALL	1.29E-07	2.62E-10	4.73E-07	2.66E-08	.108	3.50E-05
SI WALL	7.83E-09	1.76E-12	4.14E-08	2.12E-09	4.70E-02	1.53E-05
OVARIES	1.93E-08	4.93E-07	2.19E-08	1.02E-06	6.89E-02	2.29E-05
TESTES	1.85E-08	4.93E-07	2.13E-08	1.02E-06	.117	4.55E-05
SPLEEN	1.81E-08	4.70E-07	2.10E-08	1.46E-05	6.49E-02	2.13E-05
UTERUS	1.88E-08	4.93E-07	2.15E-08	1.02E-06	1.94E-02	6.35E-06
THYMUS	1.88E-08	4.93E-07	2.20E-08	1.02E-06	7.65E-02	2.50E-05
THYROID	9.99E-08	2.63E-07	1.15E-07	5.27E-07	.621	2.05E-04
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	8.97E-12	2.34E-09	1.03E-11	5.08E-09	4.81E+01	1.77E-02

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

L-12

FOR NUCLIDE : PO-210, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, F1=0.100E+00

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET	HIGH LET	LOW LET	HIGH LET	IMMERSION	SURFACE
	(A)	(A)	(A)	(A)	(B)	(C)
R MAR	1.91E-11	2.66E-05	3.90E-11	4.10E-05	4.87E+04	9.45
ENDOST	1.21E-11	1.22E-05	2.91E-11	1.89E-05	5.26E+04	10.2
PUL	1.15E-11	.0	3.26E-10	3.92E-03	4.33E+04	8.40
MUSCLE	1.43E-11	2.63E-05	3.51E-11	4.05E-05	4.62E+04	8.95
LIVER	3.14E-11	8.15E-05	7.57E-11	1.26E-04	3.93E+04	7.62
S WALL	2.64E-11	2.26E-07	5.59E-11	1.11E-07	4.11E+04	7.97
PANCREAS	3.76E-11	2.63E-05	8.49E-11	4.05E-05	3.85E+04	7.46
LLI WALL	7.89E-11	9.01E-06	5.02E-11	4.42E-06	3.08E+04	5.97
KIDNEYS	7.50E-11	4.72E-04	1.24E-10	7.28E-04	4.25E+04	8.24
BL WALL	1.92E-11	1.31E-05	2.15E-11	2.03E-05	4.00E+04	7.75
ULI WALL	5.31E-11	3.00E-06	4.47E-11	1.47E-06	4.02E+04	7.80
SI WALL	4.11E-11	5.08E-07	3.67E-11	2.49E-07	3.50E+04	6.79
OVARIES	4.27E-11	2.63E-05	3.64E-11	4.05E-05	2.38E+04	4.62
TESTES	1.08E-11	2.63E-05	1.42E-11	4.05E-05	4.49E+04	8.70
SPLEEN	1.15E-10	8.15E-04	2.02E-10	1.26E-03	4.69E+04	9.09
UTERUS	3.51E-11	2.63E-05	4.26E-11	4.05E-05	3.18E+04	6.17
THYMUS	1.54E-11	2.63E-05	7.40E-11	4.05E-05	3.03E+04	5.87
THYROID	8.16E-12	2.63E-05	2.95E-11	4.05E-05	3.52E+04	6.82
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	3.23E-10	7.84E-04	4.24E-10	1.21E-03	1.35E+06	261.
OVARIES	1.28E-09	7.84E-04	1.09E-09	1.21E-03	7.14E+05	139.
AVERAGE	8.01E-10	7.84E-04	7.55E-10	1.21E-03	1.03E+06	200.

CANCER	FATAL CANCER RISK		CONVERSION FACTORS FOR LIFETIME EXPOSURE			
	INGESTION		INHALATION		AIR	GROUND
	LOW LET	HIGH LET	LOW LET	HIGH LET	IMMERSION	SURFACE
	(G)	(G)	(G)	(G)	(H)	(I)
R MARROW	6.23E-12	1.73E-04	1.27E-11	2.67E-04	1.59E-02	3.08E-06
ENDOST	3.73E-13	3.77E-06	8.95E-13	5.80E-06	1.62E-03	3.14E-07
PULMONARY	6.97E-12	.0	1.98E-10	2.38E-02	2.63E-02	5.11E-06
BREAST	5.70E-12	1.05E-05	1.40E-11	1.61E-05	1.84E-02	3.57E-06
LIVER	4.90E-12	1.27E-04	1.18E-11	1.96E-04	6.14E-03	1.19E-06
ST WALL	2.29E-12	1.96E-07	4.84E-12	9.60E-08	3.57E-03	6.91E-07
PANCREAS	4.57E-12	3.19E-05	1.03E-11	4.91E-05	4.68E-03	9.06E-07
LLI WALL	5.47E-12	6.25E-06	3.48E-12	3.06E-06	2.14E-03	4.14E-07
KIDNEYS	2.60E-12	1.64E-04	4.30E-12	2.52E-04	1.47E-03	2.86E-07
BL WALL	6.67E-13	4.56E-06	7.43E-13	7.01E-06	1.39E-03	2.69E-07
ULI WALL	1.84E-12	1.04E-06	1.55E-12	5.09E-07	1.40E-03	2.71E-07
SI WALL	7.14E-13	8.82E-08	6.36E-13	4.31E-08	6.07E-04	1.18E-07
OVARIES	7.41E-13	4.56E-06	6.30E-13	7.01E-06	4.13E-04	8.02E-08
TESTES	1.88E-13	4.56E-06	2.46E-13	7.01E-06	7.79E-04	1.51E-07
SPLEEN	2.00E-12	1.41E-04	3.50E-12	2.17E-04	8.14E-04	1.58E-07
UTERUS	6.09E-13	4.56E-06	7.38E-13	7.01E-06	5.52E-04	1.07E-07
THYMUS	2.67E-13	4.56E-06	1.28E-12	7.01E-06	5.26E-04	1.02E-07
THYROID	6.91E-13	2.22E-06	2.50E-12	3.42E-06	2.98E-03	5.77E-07
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	2.40E-16	2.35E-08	2.26E-16	3.63E-08	3.09E-01	6.00E-05

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
- (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
- (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
- (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
- (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
- (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
- (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
- (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
- (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
- (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
- (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
- (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : TH-232, RESP CLEARANCE CLASS=Y, PARTICLE SIZE=1.0 AMAD, F1=0.200E-03

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	9.24E-08	5.22E-05	5.54E-05	1.41E-02	1.40E+06	455.
ENDOST	2.50E-07	9.30E-04	9.91E-05	.243	1.54E+06	511.
PUL	6.82E-09	4.29E-11	5.84E-03	4.22E-02	6.72E+05	209.
MUSCLE	8.80E-09	1.99E-07	2.81E-05	9.18E-05	7.91E+05	389.
LIVER	9.44E-09	9.49E-07	4.81E-05	3.56E-04	5.55E+05	145.
S WALL	5.18E-08	1.70E-07	4.01E-05	2.67E-07	5.39E+05	178.
PANCREAS	8.21E-09	1.99E-07	4.68E-05	9.18E-05	4.09E+05	131.
LLI WALL	2.15E-06	7.57E-06	3.40E-05	9.48E-06	3.97E+05	138.
KIDNEYS	8.46E-09	1.88E-07	2.19E-05	8.82E-05	5.56E+05	145.
BL WALL	5.00E-09	9.95E-08	4.24E-06	4.59E-05	4.69E+05	121.
ULI WALL	7.14E-07	2.52E-06	2.90E-05	3.35E-06	4.63E+05	120.
SI WALL	1.31E-07	4.27E-07	1.52E-05	5.84E-07	4.50E+05	116.
OVARIES	1.33E-08	1.99E-07	8.14E-06	9.18E-05	4.31E+05	131.
TESTES	6.08E-09	1.99E-07	3.85E-06	9.18E-05	8.78E+05	477.
SPLEEN	6.72E-09	1.88E-07	4.22E-05	8.70E-05	5.10E+05	144.
UTERUS	7.41E-09	1.99E-07	7.07E-06	9.18E-05	2.51E+05	62.5
THYMUS	6.31E-09	1.99E-07	7.63E-05	9.18E-05	4.97E+05	130.
THYROID	5.79E-09	1.99E-07	2.43E-05	9.18E-05	9.11E+05	253.
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	7.86E-08	5.41E-06	4.35E-05	1.58E-03	2.63E+07	1.43E+04
OVARIES	2.64E-07	5.41E-06	1.13E-04	1.58E-03	1.29E+07	3.93E+03
AVERAGE	1.72E-07	5.41E-06	7.83E-05	1.58E-03	1.96E+07	9.12E+03

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	1.77E-08	2.14E-04	1.05E-05	5.26E-02	.456	1.48E-04
ENDOST	4.33E-09	1.73E-04	1.63E-06	4.06E-02	4.74E-02	1.57E-05
PULMONARY	1.88E-09	2.61E-10	1.85E-03	.242	.409	1.27E-04
BREAST	1.74E-09	7.29E-08	5.52E-06	2.31E-05	.316	1.55E-04
LIVER	9.62E-10	1.36E-06	3.76E-06	3.75E-04	8.67E-02	2.26E-05
ST WALL	4.31E-09	1.48E-07	1.85E-06	2.09E-07	4.68E-02	1.54E-05
PANCREAS	5.04E-10	2.22E-07	2.82E-06	7.04E-05	4.97E-02	1.59E-05
LLI WALL	1.49E-07	5.25E-06	1.91E-06	6.00E-06	2.76E-02	9.58E-06
KIDNEYS	1.46E-10	5.98E-08	3.77E-07	1.93E-05	1.93E-02	5.03E-06
BL WALL	9.78E-11	3.17E-08	7.39E-08	1.01E-05	1.63E-02	4.20E-06
ULI WALL	2.47E-08	8.74E-07	7.44E-07	1.06E-06	1.61E-02	4.16E-06
SI WALL	2.21E-09	7.40E-08	1.74E-07	9.19E-08	7.81E-03	2.01E-06
OVARIES	1.65E-10	3.17E-08	7.56E-08	1.01E-05	7.48E-03	2.27E-06
TESTES	5.49E-11	3.17E-08	3.11E-08	1.01E-05	1.52E-02	8.28E-06
SPLEEN	5.97E-11	3.00E-08	3.64E-07	9.52E-06	8.85E-03	2.50E-06
UTERUS	7.45E-11	3.17E-08	6.27E-08	1.01E-05	4.36E-03	1.08E-06
THYMUS	5.63E-11	3.17E-08	6.59E-07	1.01E-05	8.62E-03	2.26E-06
THYROID	2.95E-10	1.59E-08	1.21E-06	5.54E-06	7.71E-02	2.14E-05
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	5.16E-14	1.62E-10	2.36E-11	4.74E-08	5.89E+00	2.73E-03

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : RA-228, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, F1=0.200E+00

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	1.76E-05	3.01E-05	2.26E-05	4.08E-05	2.49E-02	5.10E-05
ENDOST	3.38E-05	3.02E-04	4.12E-05	4.16E-04	3.08E-02	6.31E-05
PUL	2.03E-06	.0	4.75E-04	3.82E-04	1.68E-02	3.45E-05
MUSCLE	4.68E-06	2.04E-05	7.64E-06	2.75E-05	6.97E-02	1.43E-04
LIVER	4.41E-06	1.94E-05	9.64E-06	3.74E-05	1.80E-04	3.68E-07
S WALL	2.15E-06	9.94E-13	8.82E-06	2.51E-08	4.23E-02	8.68E-05
PANCREAS	5.03E-06	2.04E-05	1.00E-05	2.75E-05	1.21E-02	2.47E-05
LLI WALL	7.07E-05	2.08E-08	5.07E-05	9.31E-07	6.97E-03	1.43E-05
KIDNEYS	4.87E-06	1.94E-05	7.41E-06	2.62E-05	2.45E-05	5.01E-08
BL WALL	3.56E-06	1.02E-05	4.00E-06	1.37E-05	2.94E-09	6.03E-12
ULI WALL	2.16E-05	1.81E-09	2.68E-05	3.18E-07	6.00E-05	1.23E-07
SI WALL	5.49E-06	3.58E-11	1.07E-05	5.31E-08	3.28E-05	6.73E-08
OVARIES	6.98E-06	2.04E-05	7.64E-06	2.75E-05	8.99E-03	1.84E-05
TESTES	4.15E-06	2.04E-05	4.74E-06	2.75E-05	.150	3.07E-04
SPLEEN	4.43E-06	1.94E-05	8.85E-06	2.60E-05	4.23E-03	8.68E-06
UTERUS	6.40E-06	2.04E-05	7.43E-06	2.75E-05	1.68E-06	3.43E-09
THYMUS	4.80E-06	2.04E-05	1.27E-05	2.75E-05	2.49E-06	5.10E-09
THYROID	3.89E-06	2.04E-05	6.63E-06	2.75E-05	8.47E-03	1.74E-05
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	1.14E-04	5.53E-04	1.30E-04	7.46E-04	4.50	9.21E-03
OVARIES	1.97E-04	5.53E-04	2.15E-04	7.46E-04	.270	5.52E-04
AVERAGE	1.56E-04	5.53E-04	1.72E-04	7.46E-04	2.38	4.88E-03

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	5.27E-06	1.82E-04	6.82E-06	2.48E-04	8.11E-09	1.66E-11
ENDOST	9.41E-07	8.02E-05	1.15E-06	1.12E-04	9.48E-10	1.94E-12
PULMINARY	1.10E-06	.0	2.89E-04	2.32E-03	1.02E-08	2.10E-11
BREAST	1.72E-06	7.45E-06	2.88E-06	1.00E-05	2.78E-08	5.71E-11
LIVER	6.41E-07	2.76E-05	1.45E-06	5.40E-05	2.81E-11	5.75E-14
ST WALL	1.74E-07	8.63E-13	7.47E-07	2.13E-08	3.67E-09	7.53E-12
PANCREAS	5.66E-07	2.27E-05	1.17E-06	3.05E-05	1.47E-09	3.00E-12
LLI WALL	4.89E-06	1.44E-08	3.48E-06	6.32E-07	4.84E-10	9.92E-13
KIDNEYS	1.57E-07	6.14E-06	2.42E-07	8.32E-06	8.50E-13	1.74E-15
BL WALL	1.16E-07	3.24E-06	1.30E-07	4.37E-06	1.02E-16	2.09E-19
ULI WALL	7.42E-07	6.29E-10	9.16E-07	1.08E-07	2.08E-12	4.27E-15
SI WALL	9.19E-08	6.21E-12	1.81E-07	9.00E-09	5.69E-13	1.17E-15
OVARIES	1.15E-07	3.24E-06	1.25E-07	4.36E-06	1.56E-10	3.19E-13
TESTES	6.70E-08	3.24E-06	7.62E-08	4.36E-06	2.60E-09	5.33E-12
SPLEEN	7.14E-08	3.07E-06	1.47E-07	4.14E-06	7.34E-11	1.51E-13
UTERUS	1.05E-07	3.24E-06	1.21E-07	4.36E-06	2.92E-14	5.95E-17
THYMUS	7.75E-08	3.24E-06	2.13E-07	4.36E-06	4.32E-14	8.85E-17
THYROID	3.14E-07	1.63E-06	5.43E-07	2.20E-06	7.17E-10	1.47E-12
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	4.68E-11	1.67E-08	5.16E-11	2.24E-08	7.16E-07	1.46E-09

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : AC-228, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, F1=0.100E-02

DOSE CONVERSION FACTORS

ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	1.22E-07	9.47E-09	8.38E-08	2.26E-06	5.41E+09	1.02E+06
ENDOST	4.26E-08	1.10E-07	1.01E-07	2.60E-05	5.85E+09	1.11E+06
PUL	2.59E-08	1.95E-14	3.62E-06	3.28E-06	4.82E+09	9.04E+05
MUSCLE	8.14E-08	1.70E-10	5.19E-08	5.72E-08	5.11E+09	9.59E+05
LIVER	9.92E-08	1.35E-08	2.79E-07	1.61E-06	4.34E+09	8.14E+05
S WALL	2.08E-06	8.80E-12	3.50E-07	2.37E-10	4.48E+09	8.39E+05
PANCREAS	1.78E-07	1.70E-10	9.70E-08	5.72E-08	4.49E+09	8.37E+05
LLI WALL	6.16E-06	7.00E-09	9.12E-07	1.02E-08	3.54E+09	6.61E+05
KIDNEYS	1.37E-07	1.65E-10	5.71E-08	5.51E-08	4.49E+09	8.45E+05
BL WALL	1.44E-07	8.52E-11	2.68E-08	2.86E-08	4.42E+09	8.28E+05
ULI WALL	8.05E-06	1.33E-09	1.17E-06	3.24E-09	4.64E+09	8.65E+05
SI WALL	3.74E-06	8.78E-11	5.49E-07	5.15E-10	3.90E+09	7.32E+05
OVARIES	5.15E-07	1.70E-10	8.49E-08	5.72E-08	3.20E+09	5.90E+05
TESTES	3.75E-08	1.70E-10	1.13E-08	5.72E-08	4.73E+09	9.00E+05
SPLEEN	1.19E-07	1.61E-10	7.62E-08	5.42E-08	5.12E+09	9.59E+05
UTERUS	2.98E-07	1.70E-10	5.28E-08	5.72E-08	3.51E+09	6.58E+05
THYMUS	1.08E-08	1.70E-10	1.09E-07	5.72E-08	3.71E+09	6.90E+05
THYROID	3.32E-09	1.70E-10	3.61E-08	5.72E-08	4.19E+09	7.86E+05
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	1.12E-06	4.89E-09	3.35E-07	1.63E-06	1.42E+11	2.70E+07
OVARIES	1.54E-05	4.89E-09	2.54E-06	1.63E-06	9.60E+10	1.77E+07
AVERAGE	8.28E-06	4.89E-09	1.44E-06	1.63E-06	1.19E+11	2.24E+07

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION	INHALATION	AIR	GROUND		
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	3.97E-08	5.92E-08	2.71E-08	1.41E-05	1.76E+03	.332
ENDOST	1.31E-09	3.23E-08	3.08E-09	7.62E-06	180.	3.42E-02
PULMONARY	1.57E-08	1.19E-13	2.20E-06	1.99E-05	2.93E+03	.550
BREAST	3.25E-08	6.55E-11	2.07E-08	2.18E-08	2.04E+03	.383
LIVER	1.55E-08	2.03E-08	4.35E-08	2.42E-06	678.	.127
ST WALL	1.80E-07	7.63E-12	3.04E-08	2.04E-10	389.	7.28E-02
PANCREAS	2.17E-08	1.99E-10	1.18E-08	6.64E-08	545.	.102
LLI WALL	4.28E-07	4.86E-09	6.33E-08	7.05E-09	246.	4.59E-02
KIDNEYS	4.77E-09	5.52E-11	1.98E-09	1.83E-08	156.	2.93E-02
BL WALL	5.00E-09	2.85E-11	9.27E-10	9.50E-09	153.	2.87E-02
ULI WALL	2.79E-07	4.62E-10	4.07E-08	1.12E-09	161.	3.00E-02
SI WALL	6.49E-08	1.52E-11	9.52E-09	8.85E-11	67.7	1.27E-02
OVARIES	8.93E-09	2.85E-11	1.47E-09	9.49E-09	55.5	1.02E-02
TESTES	6.50E-10	2.85E-11	1.94E-10	9.49E-09	82.1	1.56E-02
SPLEEN	2.07E-09	2.70E-11	1.32E-09	9.00E-09	88.8	1.66E-02
UTERUS	5.16E-09	2.85E-11	9.14E-10	9.49E-09	60.9	1.14E-02
THYMUS	1.88E-10	2.85E-11	1.89E-09	9.49E-09	64.4	1.20E-02
THYROID	2.81E-10	1.41E-11	3.05E-09	4.71E-09	355.	6.65E-02
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	2.49E-12	1.47E-13	4.32E-13	4.89E-11	3.57E+04	6.70E+00

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : TH-228, RESP CLEARANCE CLASS=Y, PARTICLE SIZE=1.0 AMAD, FI=0.200E-03

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	2.27E-07	1.17E-05	1.52E-05	1.13E-03	1.68E+07	3.98E+03
ENDOST	2.05E-07	1.33E-04	2.04E-05	1.29E-02	1.81E+07	4.32E+03
PUL	1.75E-08	7.32E-11	1.44E-03	6.21E-02	9.28E+06	2.17E+03
MUSCLE	9.50E-08	3.33E-07	7.94E-06	3.23E-05	9.69E+06	2.44E+03
LIVER	6.37E-08	1.20E-06	1.45E-05	1.26E-04	8.02E+06	1.84E+03
S WALL	1.93E-07	2.36E-07	1.77E-05	3.55E-06	7.10E+06	1.67E+03
PANCREAS	6.26E-08	3.33E-07	1.22E-05	3.23E-05	5.81E+06	1.36E+03
LLI WALL	3.44E-05	2.20E-05	3.91E-04	1.19E-04	5.70E+06	1.35E+03
KIDNEYS	7.92E-08	3.43E-07	7.68E-06	3.99E-05	7.75E+06	1.78E+03
BL WALL	2.62E-07	1.67E-07	4.91E-06	1.61E-05	6.93E+06	1.59E+03
ULI WALL	4.62E-06	5.08E-06	1.31E-04	4.30E-05	7.02E+06	1.61E+03
SI WALL	7.45E-07	6.39E-07	2.73E-05	7.39E-06	6.92E+06	1.59E+03
OVARIES	7.91E-07	3.33E-07	1.41E-05	3.23E-05	5.28E+06	1.23E+03
TESTES	9.75E-08	3.33E-07	2.55E-06	3.23E-05	1.18E+07	2.98E+03
SPLEEN	6.31E-08	3.18E-07	1.09E-05	3.08E-05	7.57E+06	1.74E+03
UTERUS	2.88E-07	3.33E-07	6.79E-06	3.23E-05	4.91E+06	1.12E+03
THYMUS	1.71E-08	3.33E-07	1.70E-05	3.23E-05	7.04E+06	1.61E+03
THYROID	1.11E-08	3.33E-07	5.28E-06	3.23E-05	1.18E+07	2.74E+03
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	2.91E-06	9.68E-06	7.19E-05	8.92E-04	3.54E+08	8.94E+04
OVARIES	2.37E-05	9.68E-06	4.05E-04	8.92E-04	1.58E+08	3.69E+04
AVERAGE	1.33E-05	9.68E-06	2.38E-04	8.92E-04	2.56E+08	6.32E+04

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	7.29E-08	7.31E-05	4.70E-06	6.90E-03	5.47	1.30E-03
ENDOST	6.12E-09	3.91E-05	5.91E-07	3.67E-03	.557	1.33E-04
PULMINARY	1.04E-08	4.45E-10	8.16E-04	.367	5.65	1.32E-03
BREAST	3.77E-08	1.29E-07	2.94E-06	1.19E-05	3.87	9.74E-04
LIVER	9.82E-09	1.82E-06	2.10E-06	1.81E-04	1.25	2.87E-04
ST WALL	1.68E-08	2.05E-07	1.44E-06	2.93E-06	.616	1.45E-04
PANCREAS	7.55E-09	3.94E-07	1.37E-06	3.62E-05	.706	1.65E-04
LLI WALL	2.38E-06	1.53E-05	2.58E-05	7.90E-05	.396	9.37E-05
KIDNEYS	2.73E-09	1.16E-07	2.49E-07	1.29E-05	.269	6.18E-05
BL WALL	9.09E-09	5.63E-08	1.61E-07	5.17E-06	.240	5.52E-05
ULI WALL	1.60E-07	1.76E-06	4.33E-06	1.42E-05	.244	5.59E-05
SI WALL	1.29E-08	1.11E-07	4.49E-07	1.22E-06	.120	2.76E-05
OVARIES	1.37E-08	5.63E-08	2.32E-07	5.17E-06	9.16E-02	2.13E-05
TESTES	1.68E-09	5.63E-08	4.14E-08	5.17E-06	.205	5.17E-05
SPLEEN	1.09E-09	5.38E-08	1.76E-07	4.94E-06	.131	3.02E-05
UTERUS	4.98E-09	5.63E-08	1.11E-07	5.17E-06	8.52E-02	1.94E-05
THYMUS	2.90E-10	5.63E-08	2.73E-07	5.17E-06	.122	2.79E-05
THYROID	9.26E-10	2.77E-08	4.26E-07	2.59E-06	.999	2.32E-04
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	3.99E-12	2.91E-10	7.16E-11	2.67E-08	7.68E+01	1.89E-02

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
- (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
- (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
- (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
- (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
- (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
- (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
- (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
- (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
- (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
- (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
- (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : RA-224, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, FI=0.200E+00

DOSE CONVERSION FACTORS						
ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	8.41E-07	2.06E-05	4.77E-07	1.54E-05	7.61E+07	1.66E+04
ENDOST	7.41E-07	1.78E-04	5.10E-07	1.33E-04	8.06E+07	1.76E+04
PUL	8.77E-08	.0	7.35E-06	6.14E-04	5.04E+07	1.10E+04
MUSCLE	4.06E-07	2.89E-06	2.36E-07	2.16E-06	5.17E+07	1.13E+04
LIVER	4.91E-07	4.16E-06	3.47E-07	2.96E-06	4.40E+07	9.58E+03
S WALL	5.97E-07	7.89E-07	4.94E-07	3.08E-07	3.99E+07	8.69E+03
PANCREAS	3.59E-07	2.89E-06	2.59E-07	2.16E-06	3.24E+07	7.06E+03
LLI WALL	9.03E-05	2.86E-05	3.47E-05	1.07E-05	3.28E+07	7.16E+03
KIDNEYS	5.07E-07	3.59E-06	3.31E-07	2.93E-06	4.21E+07	9.18E+03
BL WALL	8.91E-07	1.44E-06	3.78E-07	1.08E-06	3.69E+07	8.05E+03
ULI WALL	2.40E-05	1.02E-05	1.01E-05	3.86E-06	4.15E+07	9.04E+03
SI WALL	3.25E-06	1.66E-06	1.61E-06	6.42E-07	3.92E+07	8.53E+03
OVARIES	2.73E-06	2.89E-06	1.12E-06	2.16E-06	2.80E+07	6.11E+03
TESTES	3.81E-07	2.89E-06	1.88E-07	2.16E-06	6.99E+07	1.52E+04
SPLEEN	3.16E-07	2.76E-06	2.27E-07	2.07E-06	4.75E+07	1.03E+04
UTERUS	1.12E-06	2.89E-06	5.02E-07	2.16E-06	3.37E+07	7.34E+03
THYMUS	1.58E-07	2.89E-06	2.13E-07	2.16E-06	4.25E+07	9.25E+03
THYROID	1.10E-07	2.89E-06	1.07E-07	2.16E-06	5.26E+07	1.15E+04
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	1.14E-05	8.66E-05	5.64E-06	6.48E-05	2.10E+09	4.56E+05
OVARIES	8.19E-05	8.66E-05	3.37E-05	6.48E-05	8.40E+08	1.83E+05
AVERAGE	4.67E-05	8.66E-05	1.97E-05	6.48E-05	1.47E+09	3.20E+05

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION		INHALATION		AIR	GROUND
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	2.74E-07	1.34E-04	1.55E-07	1.00E-04	24.8	5.40E-03
ENDOST	2.28E-08	5.49E-05	1.57E-08	4.09E-05	2.48	5.42E-04
PULMONARY	5.34E-08	.0	4.47E-06	3.73E-03	30.7	6.69E-03
BREAST	1.62E-07	1.15E-06	9.40E-08	8.63E-07	20.6	4.51E-03
LIVER	7.67E-08	6.50E-06	5.42E-08	4.63E-06	6.87	1.50E-03
ST WALL	5.18E-08	6.84E-07	4.28E-08	2.67E-07	3.46	7.54E-04
PANCREAS	4.36E-08	3.51E-06	3.15E-08	2.63E-06	3.94	8.58E-04
LLI WALL	6.26E-06	1.98E-05	2.41E-06	7.41E-06	2.28	4.97E-04
KIDNEYS	1.76E-08	1.25E-06	1.15E-08	1.02E-06	1.46	3.19E-04
BL WALL	3.09E-08	5.01E-07	1.31E-08	3.75E-07	1.28	2.79E-04
ULI WALL	8.33E-07	3.54E-06	3.49E-07	1.34E-06	1.44	3.14E-04
SI WALL	5.65E-08	2.87E-07	2.79E-08	1.11E-07	.680	1.48E-04
OVARIES	4.74E-08	5.01E-07	1.95E-08	3.75E-07	.486	1.06E-04
TESTES	6.61E-09	5.01E-07	3.26E-09	3.75E-07	1.21	2.64E-04
SPLEEN	5.49E-09	4.79E-07	3.94E-09	3.58E-07	.824	1.79E-04
UTERUS	1.94E-08	5.01E-07	8.71E-09	3.75E-07	.585	1.27E-04
THYMUS	2.74E-09	5.01E-07	3.70E-09	3.75E-07	.737	1.61E-04
THYROID	9.31E-09	2.45E-07	9.05E-09	1.83E-07	4.45	9.74E-04
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	1.40E-11	2.59E-09	5.91E-12	1.95E-09	4.41E+02	9.59E-02

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : PB-212, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, FI=0.200E+00

DOSE CONVERSION FACTORS

ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET (A)	HIGH LET (A)	LOW LET (A)	HIGH LET (A)	IMMERSION (B)	SURFACE (C)
R MAR	4.76E-07	3.64E-06	2.67E-07	2.45E-06	1.15E+09	2.57E+05
ENDOST	4.80E-07	2.73E-05	3.29E-07	1.79E-05	1.23E+09	2.75E+05
PUL	7.63E-08	.0	1.14E-05	9.73E-05	7.25E+08	1.61E+05
MUSCLE	2.21E-07	2.08E-07	1.46E-07	1.90E-07	7.49E+08	1.66E+05
LIVER	9.06E-07	5.23E-06	6.39E-07	3.42E-06	6.32E+08	1.40E+05
S WALL	2.31E-06	1.27E-07	8.89E-07	5.57E-08	5.75E+08	1.27E+05
PANCREAS	3.25E-07	2.08E-07	2.47E-07	1.90E-07	4.62E+08	1.02E+05
LLI WALL	2.49E-05	2.12E-06	4.93E-06	4.18E-07	4.67E+08	1.03E+05
KIDNEYS	7.05E-07	3.30E-06	8.59E-07	6.96E-06	6.08E+08	1.35E+05
BL WALL	3.80E-07	1.04E-07	1.02E-07	9.47E-08	5.34E+08	1.18E+05
ULI WALL	2.16E-05	1.76E-06	4.40E-06	3.56E-07	5.85E+08	1.29E+05
SI WALL	6.40E-06	4.57E-07	1.48E-06	1.10E-07	5.55E+08	1.23E+05
OVARIES	1.27E-06	2.08E-07	3.00E-07	1.90E-07	4.06E+08	9.02E+04
TESTES	1.56E-07	2.08E-07	6.06E-08	1.90E-07	9.78E+08	2.16E+05
SPLEEN	2.27E-07	1.96E-07	2.04E-07	1.78E-07	6.64E+08	1.47E+05
UTERUS	6.35E-07	2.08E-07	1.77E-07	1.90E-07	4.55E+08	1.00E+05
THYMUS	8.88E-08	2.08E-07	2.58E-07	1.90E-07	5.95E+08	1.31E+05
THYROID	5.47E-08	2.08E-07	9.20E-08	1.90E-07	7.98E+08	1.78E+05
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	4.68E-06	6.24E-06	1.82E-06	5.69E-06	2.93E+10	6.48E+06
OVARIES	3.82E-05	6.24E-06	9.01E-06	5.69E-06	1.22E+10	2.71E+06
AVERAGE	2.15E-05	6.24E-06	5.41E-06	5.69E-06	2.08E+10	4.59E+06

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION	INHALATION	AIR	GROUND		
	LOW LET (G)	HIGH LET (G)	LOW LET (G)	HIGH LET (G)	IMMERSION (H)	SURFACE (I)
R MARROW	1.55E-07	2.37E-05	8.69E-08	1.59E-05	374.	8.37E-02
ENDOST	1.48E-08	8.40E-06	1.01E-08	5.52E-06	37.9	8.46E-03
PULMINARY	4.64E-08	.0	6.92E-06	5.92E-04	440.	9.79E-02
BREAST	8.83E-08	8.30E-08	5.84E-08	7.56E-08	298.	6.62E-02
LIVER	1.42E-07	8.16E-06	9.98E-08	5.35E-06	98.6	2.19E-02
ST WALL	2.01E-07	1.10E-07	7.72E-08	4.84E-08	49.9	1.10E-02
PANCREAS	3.95E-08	2.53E-07	3.00E-08	2.30E-07	56.1	1.24E-02
LLI WALL	1.73E-06	1.47E-06	3.42E-07	2.90E-07	32.4	7.15E-03
KIDNEYS	2.45E-08	1.15E-06	2.98E-08	2.42E-06	21.1	4.68E-03
BL WALL	1.32E-08	3.61E-08	3.54E-09	3.29E-08	18.5	4.09E-03
ULI WALL	7.48E-07	6.09E-07	1.53E-07	1.24E-07	20.3	4.48E-03
SI WALL	1.11E-07	7.93E-08	2.57E-08	1.91E-08	9.63	2.13E-03
OVARIES	2.21E-08	3.61E-08	5.21E-09	3.29E-08	7.04	1.57E-03
TESTES	2.70E-09	3.61E-08	1.05E-09	3.29E-08	17.0	3.75E-03
SPLEEN	3.93E-09	3.40E-08	3.53E-09	3.10E-08	11.5	2.55E-03
UTERUS	1.10E-08	3.61E-08	3.08E-09	3.29E-08	7.89	1.74E-03
THYMUS	1.54E-09	3.61E-08	4.48E-09	3.29E-08	10.3	2.27E-03
THYROID	4.63E-09	1.76E-08	7.79E-09	1.60E-08	67.5	1.51E-02
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	6.44E-12	1.88E-10	1.62E-12	1.71E-10	6.23E+03	1.38E+00

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : BI-212, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, FI=0.500E-01

DOSE CONVERSION FACTORS

ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET	HIGH LET	LOW LET	HIGH LET	IMMERSSION	SURFACE
	(A)	(A)	(A)	(A)	(B)	(C)
R MAR	2.43E-08	3.34E-09	1.25E-08	2.35E-08	1.08E+09	2.00E+05
ENDOST	1.24E-08	9.66E-09	1.00E-08	6.80E-08	1.15E+09	2.14E+05
PUL	1.29E-08	7.98E-10	9.87E-07	1.06E-05	9.78E+08	1.80E+05
MUSCLE	2.08E-08	2.23E-09	1.24E-08	1.57E-08	1.04E+09	1.92E+05
LIVER	2.79E-08	2.15E-09	2.04E-08	1.51E-08	8.85E+08	1.63E+05
S WALL	1.95E-06	1.98E-07	8.49E-08	1.23E-08	9.43E+08	1.74E+05
PANCREAS	9.57E-08	2.23E-09	2.37E-08	1.57E-08	8.64E+08	1.58E+05
LLI WALL	1.72E-07	1.40E-08	8.14E-09	6.06E-09	7.66E+08	1.40E+05
KIDNEYS	7.68E-08	2.51E-07	1.81E-07	1.76E-06	8.75E+08	1.63E+05
BL WALL	2.29E-08	1.51E-09	3.54E-09	1.06E-08	8.89E+08	1.64E+05
ULI WALL	8.74E-07	7.77E-08	3.53E-08	8.24E-09	9.71E+08	1.78E+05
SI WALL	1.23E-06	1.30E-07	4.68E-08	1.00E-08	7.91E+08	1.46E+05
OVARIES	7.08E-08	2.23E-09	6.52E-09	1.57E-08	6.87E+08	1.23E+05
TESTES	8.85E-09	2.23E-09	2.96E-09	1.57E-08	9.14E+08	1.71E+05
SPLEEN	5.77E-08	2.15E-09	2.15E-08	1.51E-08	1.04E+09	1.93E+05
UTERUS	5.72E-08	2.23E-09	6.59E-09	1.57E-08	7.18E+08	1.32E+05
THYMUS	9.85E-09	2.23E-09	3.01E-08	1.57E-08	7.86E+08	1.43E+05
THYROID	5.17E-09	2.23E-09	1.01E-08	1.57E-08	8.81E+08	1.61E+05
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	2.65E-07	6.68E-08	8.88E-08	4.70E-07	2.74E+10	5.13E+06
OVARIES	2.12E-06	6.68E-08	1.95E-07	4.70E-07	2.06E+10	3.69E+06
AVERAGE	1.19E-06	6.68E-08	1.42E-07	4.70E-07	2.40E+10	4.41E+06

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		FOR LIFETIME EXPOSURE	
	INGESTION		INHALATION		AIR	GROUND
	LOW LET	HIGH LET	LOW LET	HIGH LET	IMMERSSION	SURFACE
	(G)	(G)	(G)	(G)	(H)	(I)
R MARROW	7.92E-09	2.18E-08	4.08E-09	1.53E-07	352.	6.51E-02
ENDOST	3.82E-10	2.97E-09	3.09E-10	2.09E-08	35.4	6.59E-03
PULMINARY	7.88E-09	4.86E-09	6.00E-07	6.44E-05	593.	.110
BREAST	8.30E-09	8.89E-10	4.96E-09	6.25E-09	415.	7.66E-02
LIVER	4.36E-09	3.36E-09	3.19E-09	2.36E-08	138.	2.55E-02
ST WALL	1.70E-07	1.72E-07	7.36E-09	1.07E-08	81.8	1.51E-02
PANCREAS	1.16E-08	2.71E-09	2.88E-09	1.90E-08	105.	1.92E-02
LLI WALL	1.19E-08	9.72E-09	5.65E-10	4.21E-09	53.1	9.72E-03
KIDNEYS	2.67E-09	8.69E-08	6.29E-09	6.11E-07	30.4	5.66E-03
BL WALL	7.95E-10	5.25E-10	1.23E-10	3.69E-09	30.8	5.69E-03
ULI WALL	3.03E-08	2.70E-08	1.23E-09	2.86E-09	33.7	6.18E-03
SI WALL	2.13E-08	2.26E-08	8.12E-10	1.74E-09	13.7	2.53E-03
OVARIES	1.23E-09	3.87E-10	1.13E-10	2.72E-09	11.9	2.13E-03
TESTES	1.54E-10	3.87E-10	5.13E-11	2.72E-09	15.9	2.97E-03
SPLEEN	1.00E-09	3.73E-10	3.73E-10	2.62E-09	18.0	3.35E-03
UTERUS	9.93E-10	3.87E-10	1.14E-10	2.72E-09	12.5	2.29E-03
THYMUS	1.71E-10	3.87E-10	5.21E-10	2.72E-09	13.6	2.48E-03
THYROID	4.37E-10	1.89E-10	8.53E-10	1.33E-09	74.6	1.36E-02
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	3.59E-13	2.01E-12	4.26E-14	1.41E-11	7.20E+03	1.32E+00

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

TABLE L.1
(CONTINUED)

FOR NUCLIDE : TL-208, RESP CLEARANCE CLASS=W, PARTICLE SIZE=1.0 AMAD, F1=0.950E+0

ORGAN	INGESTION		INHALATION		AIR	GROUND
	LOW LET	HIGH LET	LOW LET	HIGH LET	IMMERSION	SURFACE
	(A)	(A)	(A)	(A)	(B)	(C)
R MAR	1.72E-09	.0	1.28E-09	.0	1.90E+10	2.98E+06
ENDOST	1.03E-09	.0	1.03E-09	.0	2.13E+10	3.32E+06
PUL	2.57E-09	.0	5.01E-08	.0	1.91E+10	2.94E+06
MUSCLE	2.23E-09	.0	1.33E-09	.0	2.01E+10	3.10E+06
LIVER	3.22E-09	.0	2.30E-09	.0	1.82E+10	2.79E+06
S WALL	2.45E-07	.0	2.35E-09	.0	1.67E+10	2.60E+06
PANCREAS	2.25E-08	.0	2.32E-09	.0	2.33E+10	3.46E+06
LLI WALL	2.45E-09	.0	1.17E-10	.0	1.66E+10	2.52E+06
KIDNEYS	5.12E-09	.0	1.19E-09	.0	1.61E+10	2.50E+06
BL WALL	8.99E-10	.0	1.55E-10	.0	1.96E+10	2.98E+06
ULI WALL	6.36E-09	.0	5.22E-10	.0	2.20E+10	3.32E+06
SI WALL	1.20E-08	.0	4.18E-10	.0	1.67E+10	2.55E+06
OVARIES	2.10E-09	.0	2.57E-10	.0	1.85E+10	2.72E+06
TESTES	3.71E-10	.0	1.05E-10	.0	1.31E+10	2.13E+06
SPLEEN	1.26E-08	.0	2.18E-09	.0	1.60E+10	2.53E+06
UTERUS	2.24E-09	.0	2.98E-10	.0	1.55E+10	2.36E+06
THYMUS	8.80E-10	.0	3.81E-09	.0	2.26E+10	3.36E+06
THYROID	3.65E-10	.0	1.11E-09	.0	2.25E+10	3.39E+06
GENETICALLY SIGNIFICANT DOSE CONVERSION FACTORS FOR 30 YEAR EXPOSURE PERIOD						
	(D)	(D)	(D)	(D)	(E)	(F)
TESTES	1.11E-08	.0	3.16E-09	.0	3.93E+11	6.39E+07
OVARIES	6.29E-08	.0	7.70E-09	.0	5.55E+11	8.16E+07
AVERAGE	3.70E-08	.0	5.43E-09	.0	4.74E+11	7.28E+07

CANCER	FATAL CANCER RISK		CONVERSION FACTORS		AIR	GROUND
	INGESTION	INHALATION	INGESTION	INHALATION	IMMERSION	SURFACE
	LOW LET	HIGH LET	LOW LET	HIGH LET	(H)	(I)
	(G)	(G)	(G)	(G)		
R MARROW	5.59E-10	.0	4.18E-10	.0	6.18E+03	.970
ENDOST	3.16E-11	.0	3.17E-11	.0	656.	.102
PULMONARY	1.56E-09	.0	3.05E-08	.0	1.16E+04	1.79
BREAST	8.89E-10	.0	5.30E-10	.0	8.02E+03	1.24
LIVER	5.02E-10	.0	3.59E-10	.0	2.84E+03	.436
ST WALL	2.13E-08	.0	2.04E-10	.0	1.45E+03	.226
PANCREAS	2.73E-09	.0	2.82E-10	.0	2.83E+03	.420
LLI WALL	1.70E-10	.0	8.15E-12	.0	1.15E+03	.175
KIDNEYS	1.78E-10	.0	4.12E-11	.0	559.	8.68E-02
BL WALL	3.12E-11	.0	5.39E-12	.0	680.	.103
ULI WALL	2.21E-10	.0	1.81E-11	.0	763.	.115
SI WALL	2.07E-10	.0	7.25E-12	.0	290.	4.42E-02
OVARIES	3.64E-11	.0	4.46E-12	.0	321.	4.72E-02
TESTES	6.44E-12	.0	1.82E-12	.0	227.	3.70E-02
SPLEEN	2.18E-10	.0	3.79E-11	.0	278.	4.39E-02
UTERUS	3.89E-11	.0	5.17E-12	.0	269.	4.09E-02
THYMUS	1.53E-11	.0	6.62E-11	.0	392.	5.83E-02
THYROID	3.09E-11	.0	9.40E-11	.0	1.90E+03	.287
GENETIC EFFECT RISK CONVERSION FACTORS						
	(J)	(J)	(J)	(J)	(K)	(L)
AVERAGE	1.11E-14	0.00E-01	1.64E-15	0.00E-01	1.42E+05	2.18E+01

TABLE OF UNITS

- (A) - 70 YEAR COMMITTED DOSE (MILLIRAD)/(PERSON PICOCURIE)
 (B) - (MILLIRAD/YR)/(PERSON MICROCURIE/CC)
 (C) - (MILLIRAD/YR)/(PERSON MICROCURIE/CM**2)
 (D) - (MILLIRAD)/(PERSON PICOCURIE/YR)
 (E) - (MILLIRAD)/(PERSON MICROCURIE/CC)
 (F) - (MILLIRAD)/(PERSON MICROCURIE/CM**2)
 (G) - (DEATHS)/(1E+5 PERSONS PICOCURIE/YR)
 (H) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CC)
 (I) - (DEATHS)/(1E+5 PERSONS PICOCURIE/CM**2)
 (J) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON PICOCURIE/YR)
 (K) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CC)
 (L) - (GENETIC EFFECTS/LIVE BIRTH)/(PERSON MICROCURIE/CM**2)

Table L.2

ADDITIONAL INPUT DATA USED BY DARTAB IN THE
HEALTH IMPACT ASSESSMENT OF AIRBORNE EMISSIONS

Ground surface correction factor for external dose = 0.50
 Quality factor = 1.0 (low LET) and 20.0 (high LET)
 Genetic effect risk coefficient for genetically significant dose (GSD) =
 300 effects/ 10^6 live birth rad(GSD) (low LET) and
 30000 effects/ 10^6 live birth rad (GSD) (high LET)
 Regional population birth rate = 0.14133E-01 births/yr
 Average lifetime expectancy = 70.7565 years
 Rn-222 decay products risk conversion factor for a lifetime exposure =
 1.69 cancer deaths/person-WL
 Organ dose weighting factors used to determine weighted mean dose equivalent

<u>target organ</u>	<u>weighting factor</u>
red marrow	0.15590
endosteal cells	0.01470
pulmonary	0.29080
muscle	0.19080
liver	0.07460
stomach wall	0.04150
pancreas	0.05810
LLI wall	0.03320
kidneys	0.01660
bladder wall	0.01660
ULI wall	0.01660
SI wall	0.00830
ovaries	0.00830
testes	0.00830
spleen	0.00830
uterus	0.00830
thymus	0.00830
thyroid	0.04050

TABLE L.3. EXAMPLE INPUT DATA FILE FOR DARTAB

```

ACT. AV. SURFACE MINE(ORE SOURCE)-MAX. INDIV. DOSE AND RISK PARAMETERS
&INPUT
  ILOC=0, JLOC=0, PLOC=100., ILET=2*1,
  DTABLE=0, 0, 1, 1, 0, 0, 1,
  RTABLE=0, 0, 1, 1, 0, 0, 1,
  FTABLE=0, 0, 1, 1, 0, 0, 1,
  OUTPUT=F, GSCFAC=0.5,
&END
&ORGAN
  NORGN=18,
  ORGN='R MAR', 'ENDOST', '*PUL*', 'MUSCLE', 'LIVER', 'S WALL',
  'PANCREAS', 'LLI WALL', 'KIDNEYS', 'BL WALL', 'ULI WALL', 'SI WALL',
  'OVARIES', 'TESTES', 'SPLEEN', 'UTERUS', 'THYMUS', 'THYROID',
  TIME=20*70.,
&END
&QFACTR
  LLET=20*1., HLET=20*20.,
&CANCER
  NCANC=18,
  CANC='R MARROW', 'ENDOST', 'PULMINARY', 'BREAST', 'LIVER', 'ST WALL',
  'PANCREAS', 'LLI WALL', 'KIDNEYS', 'BL WALL', 'ULI WALL', 'SI WALL',
  'OVARIES', 'TESTES', 'SPLEEN', 'UTERUS', 'THYMUS', 'THYROID',
  RELABS=20*1.,
&END
&GENETIC
  GENEFF=T, NGEN=3, GEN='TESTES', 'OVARIES', 'AVERAGE',
  GRFAC=200., 20000., REPPER=.014133,
  GLLET=3*1., GHLET=3*20.,
&END
&LOCTBL
  NTLOC=1,
  RNLOC='SUM',
  OGLOC='SUM',
  PTLOC=7,
  FALOC=2,
  HLLOC=1,
  LTABLE=1,
&END
&ORGANF
  NORGB=18,
  ORGB='R MAR', 'ENDOST', '*PUL*', 'MUSCLE', 'LIVER', 'S WALL',
  'PANCREAS', 'LLI WALL', 'KIDNEYS', 'BL WALL', 'ULI WALL', 'SI WALL',
  'OVARIES', 'TESTES', 'SPLEEN', 'UTERUS', 'THYMUS', 'THYROID',
  IPATH=18*5,
  ORGDAT=.1559, .0147, .2908, .1908, .0746, .0415, .0581, .0332, .0166,
  .0166, .0166, .0083, .0083, .0083, .0083, .0083, .0083, .0405,
&END

```

Table L.4 Maximum individual fatal cancer risk for one year
of exposure to atmospheric radioactive emissions
from model uranium mines

Lifetime fatal cancer risk						
Source	Mining Activities	Ore	Sub-ore	Overburden	Vehicular Dust	Total
Average Surface Mine						
Particulates & Rn-222	1.6E-7	3.6E-7	1.7E-8	8.4E-8	4.8E-8	6.7E-7
Rn-222 daughters	3.3E-6	7.0E-7	8.3E-7	6.6E-7	0	5.5E-6
Total	3.5E-6	1.1E-6	8.5E-7	7.5E-7	4.8E-8	6.2E-6
Average Large Surface Mine						
Particulates & Rn-222	9.6E-7	1.9E-6	5.9E-8	4.7E-7	2.9E-7	3.7E-6
Rn-222 daughters	1.3E-5	1.3E-6	2.8E-6	2.9E-6	0	1.9E-5
Total	1.4E-5	3.2E-6	2.8E-6	3.4E-6	2.9E-7	2.3E-5
Average Underground Mine						
Particulates & Rn-222	3.1E-8	5.8E-8	6.5E-8	6.3E-10	4.0E-9	1.6E-7
Rn-222 daughters	9.0E-6	2.2E-7	1.8E-6	1.4E-8	0	1.1E-5
Total	9.0E-6	2.8E-7	1.9E-6	1.5E-8	4.0E-9	1.1E-5
Average Large Underground Mine						
Particulates & Rn-222	3.4E-7	6.4E-7	3.7E-7	3.3E-9	8.0E-9	1.4E-6
Rn-222 daughters	1.0E-4	2.0E-6	9.8E-6	7.6E-8	0	1.1E-4
Total	1.0E-4	2.6E-6	1.0E-5	7.9E-8	8.0E-9	1.1E-4
Inactive Surface Mine						
Particulates & Rn-222	0	N.A. (a)	N.A.	5.5E-8 (b)	N.A.	5.5E-8
Rn-222 daughters	1.3E-7	N.A.	N.A.	2.9E-7 (b)	N.A.	4.2E-7
Total	1.3E-7	N.A.	N.A.	3.4E-7 (b)	N.A.	4.7E-7
Inactive Underground Mine						
Particulates & Rn-222	0	N.A.	N.A.	1.5E-8 (b)	N.A.	1.5E-8
Rn-222 daughters	2.2E-7	N.A.	N.A.	5.0E-8 (b)	N.A.	2.7E-7
Total	2.2E-7	N.A.	N.A.	6.5E-8 (b)	N.A.	2.8E-7
In situ Leach Mine						
Particulates & Rn-222	1.6E-6	N.A.	N.A.	N.A.	N.A.	1.6E-6
Rn-222 daughters	1.1E-5	N.A.	N.A.	N.A.	N.A.	1.1E-5
Total	1.3E-5	N.A.	N.A.	N.A.	N.A.	1.3E-5

(a) N.A. - Not applicable.

(b) Consists of waste rock covered over by sub-ore (see Sections 3.7.1.1 and 3.7.2.1).

Table L.5 Fatal cancer risk to an average individual in the regional population for one year of exposure to atmospheric radioactive emissions from model uranium mines

Source	Lifetime fatal cancer risk					Total
	Mining Activities	Ore	Sub-ore	Overburden	Vehicular Dust	
Average Surface Mine						
Particulates & Rn-222	1.9E-10	4.0E-10	2.0E-11	9.1E-11	4.7E-11	7.5E-10
Rn-222 daughters	6.5E-9	1.4E-9	1.6E-9	1.3E-9	0	1.1E-8
Total	6.7E-9	1.8E-9	1.6E-9	1.4E-9	4.7E-11	1.2E-8
Average Large Surface Mine						
Particulates & Rn-222	1.1E-9	1.7E-9	7.0E-11	5.5E-10	2.8E-10	3.7E-9
Rn-222 daughters	2.6E-8	3.1E-9	5.4E-9	6.6E-9	0	4.1E-8
Total	2.7E-8	4.8E-9	5.5E-9	7.1E-9	2.8E-10	4.5E-8
Average Underground Mine						
Particulates & Rn-222	9.7E-11	7.8E-11	9.8E-11	9.2E-13	6.5E-12	2.8E-10
Rn-222 daughters	4.0E-8	1.0E-9	8.0E-9	6.6E-11	0	4.9E-8
Total	4.0E-8	1.1E-9	8.1E-9	6.7E-11	6.5E-12	4.9E-8
Average Large Underground Mine						
Particulates & Rn-222	1.1E-9	8.6E-10	5.6E-10	4.9E-12	1.3E-11	2.5E-9
Rn-222 daughters	4.5E-7	9.0E-9	4.4E-8	3.4E-10	0	5.0E-7
Total	4.5E-7	9.9E-9	4.5E-8	3.4E-10	1.3E-11	5.0E-7
Inactive Surface Mine						
Particulates & Rn-222	0	N.A. (a)	N.A.	6.4E-11 (b)	N.A.	6.4E-11
Rn-222 daughters	2.6E-10	N.A.	N.A.	5.7E-10 (b)	N.A.	8.3E-10
Total	2.6E-10	N.A.	N.A.	6.3E-10 (b)	N.A.	8.9E-10
Inactive Underground Mine						
Particulates & Rn-222	0	N.A.	N.A.	2.0E-11 (b)	N.A.	2.0E-11
Rn-222 daughters	9.9E-10	N.A.	N.A.	2.2E-10 (b)	N.A.	1.2E-9
Total	9.9E-10	N.A.	N.A.	2.4E-10 (b)	N.A.	1.2E-9
In situ Leach Mine						
Particulates & Rn-222	8.7E-10	N.A.	N.A.	N.A.	N.A.	8.7E-10
Rn-222 daughters	2.1E-8	N.A.	N.A.	N.A.	N.A.	2.1E-8
Total	2.2E-8	N.A.	N.A.	N.A.	N.A.	2.2E-8

(a) N.A. - not applicable.

(b) Consists of waste rock covered over by sub-ore (see Sections 3.7.1.1 and 3.7.2.1).

Table L.6 Fatal cancer risk to the population for one year of exposure
to atmospheric radioactive emissions from model uranium mines

Source	Estimated fatal cancers					Total
	Mining Activities	Ore	Sub-ore	Overburden	Vehicular Dust	
Average						
Surface Mine						
Particulates & Rn-222	2.6E-6	5.7E-6	2.8E-7	1.3E-6	6.8E-7	1.1E-5
Rn-222 daughters	9.3E-5	2.0E-5	2.3E-5	1.9E-5	0	1.6E-4
Total	9.5E-5	2.6E-5	2.4E-5	2.0E-5	6.8E-7	1.7E-4
Average Large						
Surface Mine						
Particulates & Rn-222	1.6E-5	2.5E-5	1.0E-6	7.9E-6	4.0E-6	5.4E-5
Rn-222 daughters	3.7E-4	4.5E-5	7.7E-5	9.4E-5	0	5.9E-4
Total	3.9E-4	7.0E-5	7.8E-5	1.0E-4	4.0E-6	6.4E-4
Average						
Underground Mine						
Particulates & Rn-222	3.5E-6	2.8E-6	3.5E-6	3.3E-8	2.3E-7	1.0E-5
Rn-222 daughters	1.4E-3	3.6E-5	2.9E-4	2.4E-6	0	1.7E-3
Total	1.4E-3	3.9E-5	2.9E-4	2.4E-6	2.3E-7	1.7E-3
Average Large						
Underground Mine						
Particulates & Rn-222	3.8E-5	3.1E-5	2.0E-5	1.8E-7	4.6E-7	9.0E-5
Rn-222 daughters	1.6E-2	3.2E-4	1.6E-3	1.2E-5	0	1.8E-2
Total	1.6E-2	3.5E-4	1.6E-3	1.2E-5	4.6E-7	1.8E-2
Inactive						
Surface Mine						
Particulates & Rn-222	0	N.A. (a)	N.A.	9.1E-7 (b)	N.A.	9.1E-7
Rn-222 daughters	3.8E-6	N.A.	N.A.	8.1E-6 (b)	N.A.	1.2E-5
Total	3.8E-6	N.A.	N.A.	9.0E-6 (b)	N.A.	1.3E-5
Inactive						
Underground Mine						
Particulates & Rn-222	0	N.A.	N.A.	7.4E-7 (b)	N.A.	7.4E-7
Rn-222 daughters	3.6E-5	N.A.	N.A.	8.0E-6 (b)	N.A.	4.4E-5
Total	3.6E-5	N.A.	N.A.	8.7E-6 (b)	N.A.	4.5E-5
In situ						
Leach Mine						
Particulates & Rn-222	1.2E-5	N.A.	N.A.	N.A.	N.A.	1.2E-5
Rn-222 daughters	3.0E-4	N.A.	N.A.	N.A.	N.A.	3.0E-4
Total	3.1E-4	N.A.	N.A.	N.A.	N.A.	3.1E-4

(a) N.A. - Not applicable.

(b) Consists of waste rock covered over by sub-ore (see Sections 3.7.1.1 and 3.7.2.1).

Table L.7 Genetic effect risk to descendants of maximum exposed individual for one year of parental exposure to atmospheric radioactive particulate and Rn-222 emissions from model uranium mines

Source	Genetic risk (effects/birth)					Total
	Mining Activities	Ore	Sub-ore	Overburden	Vehicular Dust	
Average Surface Mine	1.5E-7	3.4E-7	1.5E-8	8.0E-8	4.4E-8	6.3E-7
Average Large Surface Mine	9.2E-7	2.0E-6	5.2E-8	4.5E-7	2.6E-7	3.7E-6
Average Underground Mine	1.3E-8	5.7E-8	6.2E-8	5.8E-10	3.6E-9	1.4E-7
Average Large Underground Mine	1.4E-7	6.3E-7	3.4E-7	3.2E-9	7.2E-9	1.1E-6
Inactive Surface Mine	3.9E-13	N.A. (a)	N.A.	6.0E-8 ^(b)	N.A.	6.0E-8
Inactive Underground Mine	6.3E-13	N.A.	N.A.	1.6E-8 ^(b)	N.A.	1.6E-8
In situ Leach Mine	8.0E-9	N.A.	N.A.	N.A.	N.A.	8.0E-9

(a) N.A. - Not applicable.

(b) Consists of waste rock covered over by sub-ore (see Sections 3.7.1.1 and 3.7.2.1).

Table L.8 Genetic effect risk to descendants of average individual of the population for one year of parental exposure to atmospheric radioactive particulate and Rn-222 emissions from model uranium mines

Source	Genetic risk (effects/birth)					Total
	Mining Activities	Ore	Sub-ore	Overburden	Vehicular Dust	
Average Surface Mine	6.2E-10	1.4E-9	6.0E-11	3.3E-10	1.6E-10	2.6E-9
Average Large Surface Mine	3.8E-9	6.3E-9	2.1E-10	2.0E-9	9.6E-10	1.3E-8
Average Underground Mine	2.7E-11	1.2E-10	1.3E-10	1.2E-12	9.3E-12	2.9E-10
Average Large Underground Mine	3.0E-10	1.3E-9	7.4E-10	6.4E-12	1.8E-11	2.4E-9
Inactive Surface Mine	7.5E-16	N.A. ^(a)	N.A.	2.4E-10 ^(b)	N.A.	2.4E-10
Inactive Underground Mine	2.8E-15	N.A.	N.A.	3.4E-11 ^(b)	N.A.	3.4E-11
In situ Leach Mine	2.7E-11	N.A.	N.A.	N.A.	N.A.	2.7E-11

(a) N.A. - Not applicable.

(b) Consists of waste rock covered over by sub-ore (see Sections 3.7.1.1 and 3.7.2.1).

Table L.9 Genetic effect risk to descendants of the regional population for one year of parental exposure to atmospheric radioactive particulates and Rn-222 emissions from model uranium mines

Source	Genetic risk (effects/yr)					Total
	Mining Activities	Ore	Sub-ore	Overburden	Vehicular Dust	
Average Surface Mine	3.8E-6	8.6E-6	3.6E-7	2.0E-6	9.9E-7	1.6E-5
Average Large Surface Mine	2.2E-5	3.8E-5	1.3E-6	1.2E-5	5.8E-6	7.9E-5
Average Underground Mine	4.2E-7	1.8E-6	2.0E-6	1.8E-8	1.4E-7	4.4E-6
Average Large Underground Mine	4.5E-6	2.0E-5	1.1E-5	9.9E-8	2.8E-7	3.6E-5
Inactive Surface Mine	4.5E-12	N.A. (a)	N.A.	1.4E-6 (b)	N.A.	1.4E-6
Inactive Underground Mine	4.3E-11	N.A.	N.A.	5.0E-7 (b)	N.A.	5.0E-7
In situ Leach Mine	1.6E-7	N.A.	N.A.	N.A.	N.A.	1.6E-7

(a) N.A. - Not applicable.

(b) Consists of waste rock covered over by sub-ore (see Sections 3.7.1.1 and 3.7.2.1).

L.3 References

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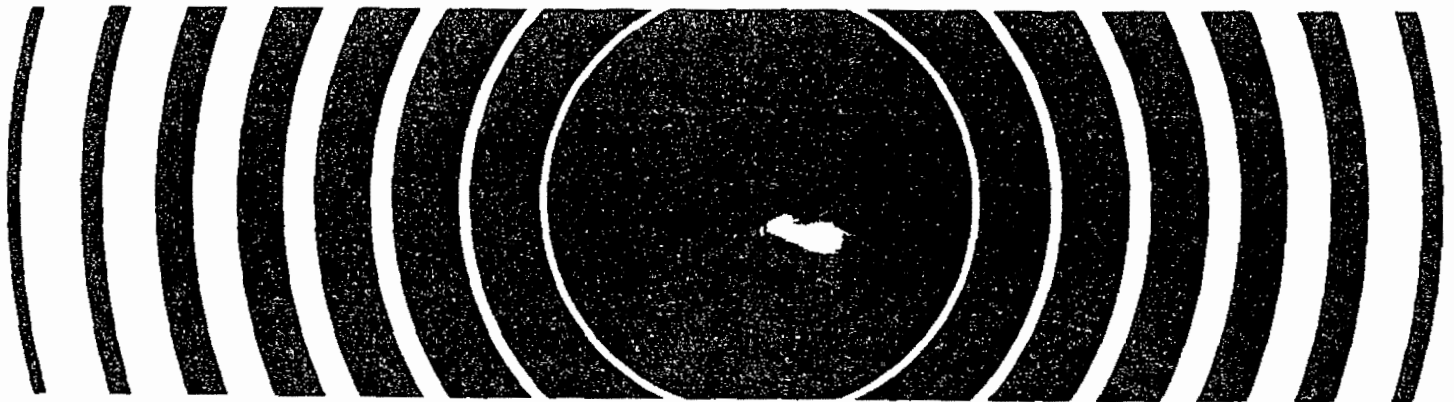
Radiation



Potential Health and Environmental Hazards of Uranium Mine Wastes

Appendixes

Report To The Congress Of The United States



Volume 3 of 3 Volumes

POTENTIAL HEALTH AND ENVIRONMENTAL
HAZARDS OF URANIUM MINE WASTES

Appendixes

A Report to the Congress of the United States
in Response to Public Law 95-604

June 10, 1983

U.S. Environmental Protection Agency
Office of Radiation Programs
Washington, D.C. 20460

CONTENTS

Appendixes

- A. Summary of Federal Laws Potentially Affecting Uranium Mining
- B. Federal Water Programs and Rights Activities
- C. Congressionally Approved Compacts that Apportion Water
- D. State Laws, Regulations, and Guides for Uranium Mining
- E. Active Uranium Mines in the United States
- F. Inactive Uranium Mines in the United States
- G. General Observations of Uranium Mine Sites in Colorado, New Mexico, Texas, and Wyoming
- H. Influence of Mine Drainage on Seepage to Groundwater and Surface Water Outflow
- I. Computation of Mass Emission Factors for Wind Erosion
- J. Aquatic Dosimetry and Health Effects Models and Parameter Values
- K. Airborne Pathway Modeling
- L. Health Risk Assessment Methodology

FIGURES

<u>Appendix G</u>	Page
G.1 Plan view of inactive underground uranium mine No. 1, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-3
G.2 Sectional view of inactive underground uranium mine No. 2, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-4
G.3 Plan view of inactive underground uranium mine No. 3, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-6
G.4 Sectional view of inactive underground uranium mine No. 4, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-7
G.5 Plan view of inactive underground uranium mine No. 5, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-8
G.6 Plan view of inactive underground uranium mine No. 6, related waste rock piles, and surface gamma exposure rates, Uravan Mineral Belt, Colorado	G-9
G.7 Plan view of inactive underground uranium mine No. 7, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado	G-11
G.8 Plan view of inactive underground uranium mine No. 8, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado	G-12
G.9 Plan view of inactive underground fluorspar uranium mine No. 9, related waste rock piles, and surface gamma exposure rates, near Jamestown, Colorado	G-13
G.10 Plan view of inactive underground uranium mine No. 10, related waste rock piles, and surface gamma exposure rates, Central City District, Colorado	G-15

	Page
G.11 Typical mine waste pile associated with a small- to medium-sized inactive underground uranium mine, Uravan Mineral Belt, Colorado	G-16
G.12 Side view of typical underground uranium mine located on the rim of a sandstone mesa, Uravan Mineral Belt, Colorado	G-16
G.13 Mine waste accumulations near the portal of a typical underground rim-type uranium mine, western Colorado .	G-17
G.14 Mine waste dump associated with a typical rim-type underground uranium mine, western Colorado . .	G-17
G.15 Movement of fluorspar-uranium mine wastes from a tailings pile into a stream, Jamestown area, Colorado . .	G-19
G.16 1972 aerial photograph of the Galen and Pavelek open pit mines, Karnes County, Texas	G-27
G.17 1978 aerial photograph of the Galen and Pavelek open pit mines, Karnes County, Texas	G-27
G.18 Results of gamma exposure rate survey at the 1601 pit and environs, Morton Ranch uranium mine, Converse County, Wyoming	G-31
G.19 Location of sampling stations at the Morton Ranch mine, South Powder River Basin, Wyoming	G-33
G.20 Sample locations for radionuclides and select trace metals in sediments, San Mateo mine, New Mexico	G-34

Appendix H

H.1 Wyoming model area sub-basin drainage system . .	H-3
H.2 Model area stream cross section	H-3
H.3 New Mexico model area sub-basin drainage system . .	H-9

Appendix J

J.1 Surface stream flow pattern within drainage area .	J-3
J.2 Conservation of mass relationship for resuspension model	J-13

TABLES

<u>Appendix A</u>	Page
A.1 Federal laws, regulations, and guides for uranium mining	A-1
<u>Appendix D</u>	
D.1 State laws, regulations, and guides for uranium mining	D-1
<u>Appendix E</u>	
E.1 Active uranium mines in the United States	E-1
<u>Appendix F</u>	
F.1 Inactive uranium mines in the United States	F-1
<u>Appendix G</u>	
G.1 Uravan and Jamestown areas	G-14
G.2 Inactive uranium mine sites surveyed in New Mexico	G-21
G.3 Status and location of uranium mines in Texas	G-25
G.4 Trace elements and radionuclides in water in the south fork of Box Creek drainage at UNC Morton Ranch lease	G-35
G.5 Radionuclides and trace metals in sediments in the south fork of Box Creek at UNC Morton Ranch lease	G-36
G.6 Radionuclides and trace metals in soils near the 1601 open pit mine, UNC Morton Ranch lease, Wyoming	G-37
G.7 Radionuclides and trace metals in soil profiles at the open pit mines, UNC Morton Ranch lease, Wyoming	G-38
G.8 Radionuclides and trace metals in sediments from the drainage of the San Mateo mine and from San Mateo Creek, New Mexico	G-40
G.9 Radium-226 and trace elements in water from San Mateo Creek near San Mateo mine discharge point	G-40

Appendix H

Page

H.1	Characteristics of the sub-basin containing the model mines	H-2
H.2	Seepage and outflow calculations for the Wyoming model mine drainage system	H-6
H.3	Characteristics of the sub-basin hydrographic unit in the model underground uranium mine area	H-8
H.4	Seepage and outflow calculations for the New Mexico model mine area drainage system	H-11

Appendix J

J.1	Aquatic environmental transport pathways examined	J-6
J.2	Characteristics of the generic sites	J-20
J.3	Stream data for Valencia County	J-23
J.4	Estimation of meat production in Valencia County for 1977	J-25
J.5	Estimates of meat production in Converse County, Wyoming for 1976	J-26
J.6	Annual radionuclide release rates to streams for active uranium mines	J-29
J.7	Freshwater fish concentration factors	J-29
J.8	Normalized human intake rate factors for radionuclide uptake via plant root systems	J-31
J.9	Irrigated land usage	J-31
J.10	Soil removal rate constants and radioactive decay constants	J-33
J.11	Milk and beef concentration factors	J-33
J.12	Dose equivalent conversion factors	J-36
J.13	Health effects conversion factors for internal pathways	J-37
J.14	Health effects conversion factors for external pathways	J-37

Appendix K

K.1	Characteristics of the generic sites	K-1
-----	--	-----

K.2	Animal and vegetable crop distribution for use with AIRDOS-EPA	K-3
K.3	Sources of food for the maximum individual (percent) .	K-4
K.4	Selected input parameters to AIRDOS-EPA	K-5
K.5	Selected terrestrial pathway parameters by radionuclide .	K-7
K.6	Effective radioactive decay constants	K-8

Appendix L

L.1	Radionuclide dose rate and health effect risk conversion factors used in uranium mine assessments	L-4
L.2	Additional input data used by DARTAB in the health impact assessment of airborne emissions	L-21
L.3	Example input data file for DARTAB	L-22
L.4	Maximum individual fatal cancer risk for one year of exposure to atmospheric radioactive emissions from model uranium mines	L-23
L.5	Fatal cancer risk to an average individual in the regional population for one year of exposure to atmospheric radioactive emissions from model uranium mines	L-24
L.6	Fatal cancer risk to the population for one year of exposure to atmospheric radioactive emissions from model uranium mines	L-25
L.7	Genetic effect risk to descendants of maximum exposed individual for one year of parental exposure to atmospheric radioactive particulate and Rn-222 emissions from model uranium mines	L-26
L.8	Genetic effect risk to descendants of average individual of the population for one year of parental exposure to atmospheric radioactive particulate and Rn-222 emissions from model uranium mines	L-27

L.9	Genetic effect risk to descendants of the regional population for one year of parental exposure to atmospheric radioactive particulates and Rn-222 emissions from model uranium mines	L-28
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