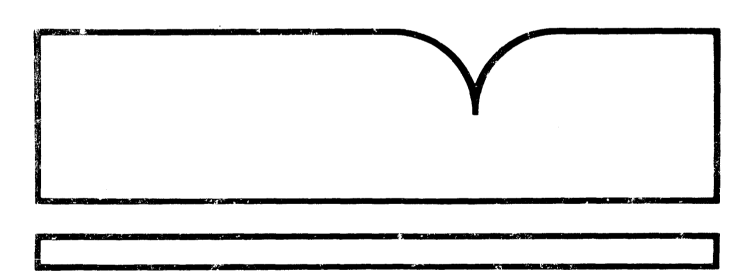
Proceedings: Workshop on Effects of Sewage Sludge Quality and Soil Properties on Plant Uptake of Sludge-Applied Trace Constituents

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PROCEEDINGS: WORKSHOP ON EFFECTS OF SEWAGE SLUDGE

QUALITY AND SOIL PROPERTIES ON PLANT UPTAKE

OF SLUDGE-APPLIED TRACE CONSTITUENTS

### Sponsored by:

U.S. Environmental Protection Agency Cincinnati, Ohio 45268

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Location: Las Vegas, Nevada Date: November 13-16, 1985 Cooperative Agreement No. CR-812673

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J. A. Ryan Water Engineering Research Laboratory U.S. Environmental Protection Agency Cincinnati, Ohio 45268

WATER ENGINEERING RESEARCH LABORATORY
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The workshop report summarizes the current research transfer of contaminants from sewage sludge to the human application. As such it addresses the important parametalter the rate and degree of movement of contaminants the The workshop group met in Las Vegas, Nevada, November participants were divided into five separate but related each workgroup are as follows: effects of soil properties elements by crops, effects of sludge properties on accurations, effects of long-term sludge applications on accurations, transfer of sludge-applied trace elements to the trace organics in sewage sludges on soil-plant systems a humans.  The report evaluates available data on effects of son plant uptake of municipal sewage sludge-applied trace transfer into the food chain. The summarized data and it to EPA in regulation and management of land application	n food chain via land ters in the system which can be rough the environment. The land two kgroups. The topics of es on accumulation of trace mulation of trace elements by food chain, and effects of and assessing their risk to sludge, soil, and plant factors e contaminants and their interpretation will be of value

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#### **FOREWORD**

The U.S. Environmental Protection Agency is charged by Congress with protecting the Nation's land, air, and water systems. Under a mandate of national environmental laws, the agency strives to formulate and implement actions leading to a compatible balance between human activities and the ability of natural systems to support and nurture life. The Clean Water Act, the Safe Drinking Water Act, and the Toxics Substances Control Act are three of the major congressional laws that provide the framework for restoring and maintaining the integrity of our Nation's water, for presering and enhancing the water we drink, and for protecting the environment from toxic substances. These laws direct the EPA to perform research to define our environmental problems, measure the impacts, and search for solutions.

The Water Engineering Research Laboratory is that component of EPA's Research and Development program concerned with preventing, treating, and managing municipal and industrial wastewater discharges; establishing practices to control and remove contaminants from drinking water and to prevent its deterioration during storage and distribution; and assessing the nature and controllability of releases of toxic substances to the air, water, and land from manufacturing processes and subsequent product uses. This publication is one of the products of that research and provides a vital communication link between the researcher and the user community.

This report evaluates available data on effects of sludge, soil, and plant factors on plant uptake of municipal sewage sludge applied trace contaminants and their transfer into the food chain. The summarized data and interpretation will be of value to EPA in regulation and management of land application of municipal sewage sludge.

Francis T. Mayo, Director Water Engineering Research Laboratory

#### **EXECUTIVE SUMMARY**

The workshop group met in Las Vegas, Nevada, November 13-16, 1985, to assess the state of our knowledge on potential problems of trace elements and trace organics associated with the land application of municipal sewage sludges. The participants were divided into five separate but related workgroups. The topics of each workgroup, participants, and their affiliations are as follows:

# I. <u>EFFECTS OF SOIL PROPERTIES ON ACCUMULATION OF TRACE ELEMENTS</u> BY CROPS

Lee E. Sommers, Chair; Colorado State University, Fort Collins, CO V. Van Volk, Oregon State University, Corvallis, OR Paul M. Giordano, Tennessee Valley Authority, Muscle Shoals, AL William E. Sopper, Pennsylvania State Univ., University Park, PA Robert Bastian, OMPC, U. S. EPA, Washington, D.C.

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### IV. TRANSFER OF SLUDGE-APPLIED TRACE ELEMENTS TO THE FOOD CHAIN

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Seattle, WA

# V. EFFECTS OF TRACE ORGANICS IN SEWAGE SLUDGES ON SOIL-PLANT SYSTEMS AND ASSESSING THEIR RISK TO HUMANS

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Each workgroup started out by reviewing the existing data base and prepared a working draft in their assigned subject matter areas. The salient features of each group's findings were presented at plenary sessions attended by the entire workshop. At this time each participant was afforded an opportunity to provide his/her input into workgroups other than the one he/she was assigned.

Following the workshop, chairs solicited participants for additional data or comments they wished to incorporate into the report. A revised draft was then prepared. These revised workgroup drafts were, in turn, reviewed by the workshop coordinators and the chairs, and recorders of other workgroups. Following this revision, the workshop coordinators, workgroup chairs and recorders met, and finalized the report. Findings of the workshop are summarized as follows:

#### I. Effects of Soil Properties on Accumulation of Trace Elements by Crops

- Although greenhouse pot studies may be useful to examine mechanisms and to establish relative response curves, the concentrations of trace elements in a particular crop are greater when the crop is grown in pots with studge treated soils than when it is grown under comparable conditions in the field.
- Experiments which employ either trace element salts or sludges spiked with trace element salts do not simulate trace element uptake by crops grown on sludge-amended soils. Therefore, results of such studies do not provide a reliable basis for establishment of

criteria, guidelines and regulations to control trace element concentrations of crops grown on sludge amended soils.

- Concentrations of trace elements in crops grown on sludge-amended soils vary with soil conditions such as the content of iron and aluminum oxides and soil pH. Iron and aluminum oxides in soils, sludges, and sludge-amended soils may reduce solubilities of trace elements and, in turn, their plant availabilities. In general, trace element uptake by crops (except Mo and Se) decreases with increasing soil pH.
- The pH measurement of a soil depends upon the method used to prepare the soil suspension. Suspensions of 1:1 soil:water or soil:0.01 M CaCl<sub>2</sub> have been used for measuring the pH of soils and/or sludge amended soils. However, the 0.01 M CaCl<sub>2</sub> method is preferred because it compensates for soluble salt contents in the soil-sludge mixture. Soil pH's in 0.01 M CaCl<sub>2</sub> are generally lower than those measured in water and regulations based on soil pri should specify the method to be used.
- Sewage sludge additions have been effective in correcting trace element deficiencies (e.g., iron, copper and zinc) of crops, particularly those grown on calcareous soils.
- Trace metal uptake by crops grown on sludge-amended soil is not directly related to the soil's cation exchange capacity or texture. Available research data do not support the continued use of cation exchange capacity or soil texture alone to determine maximum available trace metal loadings.

## II. Effects of Sludge Properties on Accumulation of Trace Elements by Crops

- Trace elements in raw sewage are associated primarily with suspended solids, and they remain as suspended solids in the sludge following wastewater treatment.
- Over the past decade, concentrations of trace elements in many publicly-operated treatment works (POTW) sludges have decreased markedly as a result of implementing industrial waste pretreatment, and this trend is expected to continue.
- During sewage treatment, addition of materials containing Fe, Al or lime reduces solubilities of metals in sludges.
- A variety of factors determine equilibrium trace element solubility in sludges, particularly the presence of trace-element precipitates (relatively pure compounds or coprecipitated with Fe, Al, or Ca precipitates), the strength of bonding to organic and mineral adsorption sites, the proportion of potential adsorbing sites

filled, and the presence of dissolved ligands capable of complexing the trace elements.

- If, within the pH range normally found in soils of a given region, a sludge maintains the availability of a trace element below the level that causes phytotoxicity or potentially harmful accumulation of that element in plants, there is no need to limit land application of that sludge because of that element.
- If, within the pH range normally found in soils of a given region, a sludge maintains the availability of a trace element above the level that causes phytotoxicity or potentially harmful accumulation of that element in plants, loading limits should be established based on characteristics of the sludge and of the soil to which it is applied that interact to control the availability of that element.
- Development of methods for measuring trace-element desorption characteristics of sludges and adsorption characteristics of soils (particularly for Cd, Zn, Ni and Cu) should be given high priority.
- Immediately following land application all sludges will undergo changes which will affect trace element solubility and plant uptake. This effect is a function of sludge treatment prior to land application. Most research indicates that plant availability of sludgederived metals stays the same or decreases with time following their land application.

### III. <u>Effects of Long-Term Sludge Applications on Accumulation of Trace</u> Elements by Crops

- Application of Cd and Zn to soils from municipal sewage sludge will cause the Cd and Zn concentrations of crops grown on these soils to exceed those of untreated controls. When the sludge is applied at rates to satisfy the N requirement of the crop grown, the Cd and Zn contents of plant tissue remain low and at nearly constant levels with successive sludge applications.
- In sludge-treated soils maintained at pH >6.0, Cu and Ni contents of vegetative tissue may become slightly elevated. Phytotoxicity from sludge-applied Cu and Ni, however, has rarely been reported.
- Available data suggest that four or more years following sludge application the trace element concentration of the affected vegetative tissue would be determined by the total amounts of trace elements in the soil and would not be affected by the frequency of sludge application (e.g., single addition vs. multiple applications).
- Plant availability of sludge-borne metals is highest during the first year sludge is applied. Using the first year response curve generated by a large single sludge addition will overestimate metal

- accumulation in vegetative tissue from plants grown in well stabilized sludge/soil systems.
- Field data indicate that trace element concentration in vegetative tissue will not rise after the termination of sludge applications if chemical conditions of the soil remain constant. Cadmium and In levels of plants grown in soils which were no longer receiving sludges either remained at the pretermination level or decreased with time.

## IV. Transfer of Sludge-Applied Trace Elements to the Food Chain

- Contents of some trace elements in edible crop tissues can be increased when sewage sludges rich in these elements are applied to soils, especially to soils that are highly acidic (Cd, Zn, Ni) or alkaline (Mo). Under conditions which allow the concentration of a trace element in crops to increase substantially (responsive conditions), the relative increases in element concentration among crop species are sufficiently consistent to be used to generate input data for modeling the dietary exposure of the element. The relative increase of trace element concentration among crops may vary when the results are extrapolated from soils with average to soils with high organic matter contents, or from acidic to calcareous soils. High organic matter and high soil pH (except for Mo and Se) both reduce element uptake and would not increase risk above that determined from the risk assessment based on conditions of maximum intake.
- Relatively high and low Cd-accumulating crop types (lettuce vs. cabbage; carrot vs. beet) within a food group should be accounted for when using the FDA food groups to model the dietary intake of Cd.
- Representative food intake from birth to age 50 should be used to calculate daily Cd ingestion and not the maximum daily intake.
- Increased Cd ingestion from consumption of crops grown on sludgeamended soils can be expressed in terms of their Cd uptake relative to a reference crop (e.g., lettuce).
- Models developed to predict Cd retention by humans should consider not only Cd content of the diet but also other constituents in the diet (e.g., Fe, Zn) that affect Cd retention.
- The highest exposure to sludge-applied Cd would result from ingestion of a substantial fraction of the daily diet of foods grown in a strongly acidic vegetable garden for many years.
- Crop cultivars differ in their Cd uptake. However, in determining dietary Cd intake, these differences are less important than differences caused by crop species and soil and sludge characteristics.

• Surface application of sludge without soil-incorporation presents a greater potential risk to humans, livestock and wildlife due to possible direct ingestion of sludge-borne trace elements. The bioavailability of a trace element in ingested sludge is strongly influenced by the concentration of the element, the presence of other interacting elements, and the sludge redox potential. Livestock showed no harmful effects when grazing on pastures treated with sewage sludge containing median trace element concentrations.

# V. Effects of Trace Organics in Sewage Sludges on Soil-Plant Systems and Assessing Their Risk to Humans

- Sewage sludges could contain thousands of trace organics.
   Organics discharged by major contributors to wastewater treatment plants should be identified to help select compounds for analysis in sewage sludge.
- Although some industrially derived organic compounds can be present in sewage sludge at relatively high concentrations (i.e., a few percent dry weight) most detected compounds are present at concentrations less than 10 mg/kg, dry weight.
- Results of bioassays of sludges for their mutagenic activity are difficult to interpret. Information obtained from these tests is not presently adequate to predict adverse environmental impacts associated with land application of sludge.
- Organic chemicals applied to soil may undergo adsorption, volatilization, degradation, leaching, and plant uptake. Many organics are strongly adsorbed to organic matter and/or undergo degradation, thus reducing the potential for plant uptake or leaching.
- Because experimental data are not always available for organics found in sludges, use of mathematical models based on physical/ chemical properties of representative organic compounds is a logical approach to predict the fate of similar sludge-derived organics in soils. Field research with selected sludge organics, which are representative of organic chemical groups, is needed to calibrate and validate these models.
- No adverse effects on the growth of crops have been observed when sludges containing these organics are applied to soil at fertilizer rates for nitrogen or lower.

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#### SECTION 1

#### INTRODUCTION

In the last two decades, this nation has experienced a dramatic increase in the construction of publicly-owned treatment works with a corresponding increase in residual solids from treating the waste water. Because the common methods of sludge disposal, such as landfill, incineration and ocean dumping may not be adequate or suitable to accommodate the ever-increasing quantities of POTW sludge, interest in applying sludges to agricultural, forest and disturbed land has increased.

In addition to valuable plant nutrients, sewage sludge contains variable concentrations of trace elements and synthetic organic compounds. Concern for trace element contamination of the food chain from land application of sewage sludge stems from extensive prior experience with phytotoxicity of elements such as Cu, Ni and Zn from smelters and other sources (Page, 1974) and from human and livestock toxicities associated with environmental contamination by Pb, Hg, Cd, Cu, F, Mo, As and other trace elements (Logan and Chaney, 1983).

The Joint Conference on Recycling Municipal Sludges and Effluents on Land (1973) raised the issue of trace element contamination from sewage sludge but the available data base on actual land application research was for the most part limited to pot studies with metal salts or sludge and a few field experiments of no more than a few years duration (Logan and Chaney, 1983). Subsequent conferences in 1980 (CAST, 1980) and 1983 (Page et al., 1983) reexamined these issues in light of the increasing body of research. By 1983 Logan and Chaney had concluded that the environmental threat from sludges applied to land at agronomic rates was minimal when existing federal regulations and guidelines (EPA, 1979a) were followed. Phytotoxicity from sludge-applied metals was no longer believed to be of concern except for high-metal content sludges applied at high loading rates on acid soils.

Inputs of sludge-borne trace elements to agricutural land in the U.S. has been governed since 1979 by EPA regulations and guidelines (EPA, 1979a). Under various provisions of existing federal statutes, cadmium was the only trace element addressed and the regulatory approach was to limit annual and cumulative applications of Cd to land, based on soil pH and soil cation exchange capacity (CEC). In addition to federal regulations, many states imposed limitations on cumulative applications of elements such as Cu, Ni, Zn (to protect against phototoxicity) and Pb (to protect the human food chain) (Logan and Chaney, 1983).

Implicit in this regulatory approach was the belief that bioavailability of sludge-applied trace elements was controlled by soil processes such as adsorption, chelation, and precipitation and that these processes were reflected by soil properties such as pH and CEC. By 1980, however, data from the increasing number of long-term field studies were beginning to indicate that sludge properties could also influence trace element bioavailability. The sludge's effect on bioavailability of trace elements was postulated by Corey et al. (1981) and later reiterated by Logan and Chaney (1983) as being due to binding of trace elements by the sludge itself. A corollary to this hypothesis is the prediction that, at high enough sludge application rates, the solubility of trace elements in soil would be controlled by the sludge and not by the soil. The implication of this theory, if true, is that the present regulation of Cd application to land with sludge on the basis of Cd loading and soil properties alone ignores what may be the equally or more important sludge properties, and may overestimate crop Cd uptake particularly from low-Cd sludges.

Parallel to, but more recent than, the evolution of our knowledge of trace element chemistry and bioavailability is the growing concern over contamination of the environment by synthetic organic compounds. This concern has led to recent but limited studies of synthetic organic compounds in sewage sludges (Naylor and Loehr, 1982; Overcash, 1983; Overcash et al., 1986) and proposals for their regulation. Under the 1979 regulations, only polychlorinated biphenyls (PCBs) were specifically controlled (EPA, 1979a). The research data base on the fate of sludge-borne organics is extremely limited, as is the information on the content of various synthetic organic compounds in sludges. As a result, uncertainties as to the health effects and threshold exposures of any of these compounds has made the evaluation of risk from sludge organics difficult.

In 1984, EPA began a process to reevaluate the existing regulations and criteria by which land application of municipal sewage sludge is controlled in the U.S. The Office of Water Regulations and Standards working with several technical advisory committees and with the Environmental Criteria and Assessment Office (ECAO) screened those pollutants found in sludge that had the potential to adversely affect the food chain, thus possibly requiring regulation. Based on this evaluation, and using a risk assessment approach developed by ECAO, hazard indices were developed for a number of trace elements and synthetic organic compounds, and subsequently were used to evaluate the potential risk from land application of sludge (EPA, 1985). Presently, a comprehensive risk assessment methodology which will be used to evaluate potential risk from land application of municipal sludge is under development by ECAO Cincinnati. The result of this and related efforts will be development of revised or, if necessary, new regulations governing land application and other means of sludge use and disposal.

The development of hazard indices and their use in risk assessment is limited by the availability of valid data for the pollutants of concern. A critical review of the data bases used to develop the hazard indices revealed that they often included studies involving metal salts addition

rather than sludge only sources and did not include many of the long-term field studies with sludges which were conducted in the late 1970's and early 1980's and which are now beginning to enter the literature.

The purpose of this workshop was to bring together researchers involved in land application of sewage sludge to evaluate their most recent data and, in light of this information, to assess the validity of assumptions made in the risk assessment process on fate of sludge contaminants. In this report, pertinent unpublished data from experiments in progress and from papers submitted for publication were supplied by the participants and incorporation in the report where appropriate. Specifically, the workshop was organized into five groups and the topics and questions addressed by group are given below:

# WORKGROUP I. Effects of Soil Properties on Accumulation of Trace Elements by Crops

- 1. What information is most relevant for defining background levels of trace elements in soils and crops? Are background levels in soils related to soil properties?
- 2. What soil properties have a direct role in regulating plant uptake of trace elements and can these be quantified?

# WORKGROUP II. Effects of Sludge Properties on Accumulation of Trace Elements by Crops

- Is there evidence that concentration and chemical form have a significant effect on plant uptake of sludge-applied trace elements?
- Are there synergistic or antagonistic effects of certain trace elements? If so, are they sufficiently well quantified to be considered in setting criteria?

# WORKGROUP III: Effects of Long-Term Sludge Applications on Accumulation of Trace Elements by Crops

- Do the long-term field plot data show any significant differences in plant uptake of annual vs. cumulative sludge-applied trace elements?
- 2. What is the evidence for increases vs. decreases of plant uptake of sludge-applied trace elements with time after sludge application?

# WORKGROUP IV: Transfer of Sludge-Applied Trace Elements to the Food Chain

- 1. How can plant uptake rates be used to provide a basis for "good practice" and "worst case" risk scenarios?
- 2. Are there sludges so low in trace constituents that no limits should be applied?
- 3. What effect may routes of exposure from land application other than crop uptake have upon "good practice" or "worst case" risk scenarios (e.g., direct ingestion of soil from sludge-treated land)?

# WORKGROUP V: Effects of Trace Organics in Sewage Sludges on Soil-Plant Systems and Assessing Their Risk to Humans

- 1. How do soil and sludge properties influence rates at which sludge organic matter is decomposed?
- 2. What synthetic organic compounds are absorbed by plants and what is known concerning the machanism of absorption?

Although the complete list of trace elements for which hazard indices were developed was addressed in the workshop, the focus of Workgroups I-IV was limited to Cd, Zn, Mo, Fe, Pb and Se, as the other trace elements usually present in sludges were considered to present little potential risk to the human or animal food chain.

#### SECTION 2

#### EFFECTS OF SOIL PROPERTIES ON

#### ACCUMULATION OF TRACE ELEMENTS BY CROPS

#### INTRODUCTION

The fate and effects of sewage sludge constituents in a soil-plant system are influenced by factors such as climate (rainfall and temperature), management (irrigation, drainage, liming, fertilization, addition of amendments), and composition of the sewage sludge. In addition, soil properties affect the chemical reactions and processes which occur after application of sewage sludge to a soil. Soil properties that affect the reactions and resultant plant uptake of sewage sludge constituents include pH, organic matter, cation exchange capacity, iron and aluminum oxides, texture, aeration, specific sorption sites and water availability. Mean values for selected soil properties are shown in Table 1. Many of these factors are interrelated and thus create a rather complex medium involving chemical and microbial reactions. The factors which tend to be stable are texture, CEC, organic matter, and iron and aluminum oxides. Factors such as pH, water content, and aeration (relates to water content) vary frequently or are easier to adjust. For example, soil pH can be increased by lime additions while ammoniacal fertilizers acidify soils.

Soil cation exchange capacity (CEC) is dependent on soil properties such as organic matter, pH, and type and percentage of clay. Thus it serves as an easily measured, integrating parameter to characterize a soil. Soil pH, like CEC, is an easily measured soil property which provides background information relevant to assessing elemental availability to plants. The soil pH measured in the laboratory is a representation of that which occurs under field conditions. The pH at any individual site in the soil may be significantly different from the pH of other sites. For example, the pH at the root-soil interface may be lower because of exuded organic acids. Due to differential uptake of cations and anions, the pH in the root cylinder of active root hairs may be lower than in older parts of the root system (Römheld and Marschner, 1986). Also, pH reductions with time in sludge-treated soils are due to protons generated during oxidation of reduced forms of N and S mineralized from sludge organic matter. Similar pH reductions occur after addition of fertilizers, particularly those containing ammonium.

Plant uptake of elements from the soil solution initially requires positional availability to the plant root. Either the element must be moved to the root through diffusion or mass flow processes, or the root must grow to the element. The element must then occur in a form which can move into the plant via the uptake mechanism. This transfer requires that the element move through a solution phase; thus, water solubility and a variety of complexation, chelation, and other chemical reactions become important.

Considerable research on microelement uptake by plants has been done with metal salts. However, metals applied to a soil as a salt, commonly a sulfate, chloride, or nitrate salt, accumulate in plants more readily than the same quantity of metal added in sewage sludge (Logan and Chaney, 1983; Dijkshoorn et al., 1981). Metal salt additions to soils can cause formation of metal chloride complexes and ion pairs which may increase metal diffusion and plant uptake (Bingham, 1980). Metals in sludges are often associated with the insoluble inorganic components (such as phosphates, sulfides, and carbonates) and are not readily plant available (Soon, 1981; Page, 1974). Elemental uptake by plants grown in soils treated with metal salts or sewage sludge amended with metal salts will be higher than actually exists for equal amounts of metal contained in sewage sludges. If results from sludge-treated soils are available, human or animal exposure models should be based on these observations and not on extrapolation of data from additions of soluble metal salts to soils.

To predict the impact of sludge use on elemental content in the human diet, plant uptake of trace elements from sewage sludge should be measured in field experiments. Greenhouse or pot study experiments generally create a root environment which increases the magnitude of trace element uptake (deVries and Tiller, 1978; Davis, 1981). The enhanced uptake of trace elements generally results from four factors: 1) the use of acid forming fertilizers; 2) increased soluble salt content from fertilizers in a smaller soil volume than in the field; 3) root confinement; and 4) unnatural watering patterns. However, greenhouse pot experiments can have value if plants are harvested in an early growth stage or if pots are sufficiently large to allow unrestricted root growth and natural water drainage. In addition, pot experiments are valuable for evaluating factors affecting plant uptake of trace elements, realizing that plant concentrations may differ from those found in a field study.

#### BACKGROUND LEVELS OF TRACE ELEMENTS

The trace element content of crops is a function of the plant available level in the soil and the modifying influences of soil chemical and physical properties. Trace element levels of soil vary with the parent material. Except for a few special cases (Lund et al., 1981), plant tissue concentrations are not positively correlated with the total trace element content in untreated soils.

Background levels of metals have been summarized for soils from Ohio (Logan and Miller, 1983), Minnesota (Pierce et al., 1982), and from 3,305 sites across the U.S. (Table 1; Holmgren et al., 1986). Although quite extensive, the data contained in Table 1 were selected on the basis of being agricultural soils removed from mobile and point sources of contamination and means representative of all agricultural soils in the U.S. may differ from those presented in Table 1. For most elements, the minimum and maximum values differ by 2 to 3 orders of magnitude. The unusually high mean values for Pb in soils from Virginia and West Virginia are due to metalliferous deposits near a few sites. One statistical approach to evaluating whether a soil has been impacted by industrial sources of metals is to compare trace element concentrations with the concentrations at the 95th percentile. As shown, the 95th percentile for soil metals (Table 2) is appreciably smaller than the maximum value. Current U.S. EPA Cd limits imposed on sludge applications (U.S. EPA, 1979a) are 5 to 10 fold greater than a liberal estimate of natural background levels. Further, a soil's total trace element content enables a preliminary evaluation of metal contamination from prior waste disposal activities. The total soil metal data also allows identification of sites where parent material contains unusually high levels of a given element.

The total metal concentration reported for soils may be influenced by the analytical methods employed, especially if a dissolution procedure is used. The total metal content in a soil requires either a non-destructive analytical method such as neutron activation analysis or a total dissolution of the soil matrix with strong acids plus HF, or partial dissolution with boiling 4  $_{\rm M}$  HNO $_{\rm 3}$ , or refluxing HNO $_{\rm 3}$ -HClO $_{\rm 4}$  (Lund et al., 1981). Once the soil matrix is dissolved, standard atomic absorption or equivalent methods can be used to analyze metals in the digest.

A need still exists for a standard extractant to assess the level of plant available metals in soils. Logan and Chaney (1983) summarize recent research on common soil metal extractants. The DTPA-TEA reagent used to detect trace metal deficiencies in calcareous soils (Lindsay and Norvell, 1978) has been used to monitor soils. Other extractants which have been used include double acid (HCl +  $H_2SO_4$ ), dilute HCl,  $Ca(NO_3)_2$ , and water saturation extracts. One approach is the "agnostic soil test used on a routine basis for soils treated with sludges in Pennsylvania (Baker and Amacher, 1981). This method involves equilibrating soils with a test solution containing cations (Ca. Mg. K, and H) at the activities and ratios determined to be near the minimum for optimum growth of plants. The solution also contains  $4 \times 10^{-4}$  M DTPA (diethylene triamine pentaacetic acid) to render a small exchange of trace metals. The extracted metal provides a measure of the labile pool and the metal-DTPA formation constants are used to calculate activities of trace metals (Baker and Amacher, 1981). However, no method used has been proven acceptable to predict plant uptake of metals from a wide range of soils (Logan and Chaney, 1983). Ideally, extraction of a soil, sewage sludge, or a sludge amended soil could be used to predict the eventual plant uptake of trace

elements. With this approach, bioavailable elements of both the soil and sewage sludge could be assessed. A procedure is also needed for quantitative measurement of specific sorption sites in soils.

Information on the background levels of metals in crops is also needed to evaluate the impact of metals entering animal or human diets. A recent survey has been conducted by the USDA-EPA-FDA for a variety of crops grown in major regions of the U.S. (Wolnick et al., 1983a, 1983b, 1985). As previously discussed for soils from the same survey, the data presented in Table 3 were for crops grown on soils removed from mobile and point sources of contamination. The data, therefore, may not be representative of the crops for U.S. agricultural soils in general. The Cd. In and Pb content of 12 common crops varies by 1 to 3 orders of magnitude (Table 3). Median concentrations of Cd in leafy vegetables were highest (spinach, 800 µg/kg); median concentrations of Cd in root crops ranged from 68 µg/kg (peanuts) to 160 µg/kg (carrots); and for grains the median Cd concentrations varied from 4 μg/kg (field corn) to 45 μg/kg (soybeans). Somewhat similar median concentrations of Pb in crops were observed (Table 3). Median concentrations of Zn across the 12 crops tested, however, were more uniform. They varied from a low of 15 µg/kg (rice) to a high of 46 µg/kg (lettuce).

### REGIONAL STUDY OF SLUDGE USE

The regional project W-124 (Optimum utilization of Sewage Sludge on Cropland) has collected data on the uptake of metals by barley grown at 15 locations in the U.S. (Table 4). At each location, the same sewage sludge sample from Chicago was applied either at 100 mt/ha in the initial year or at 20 mt/ha each year for 5 consecutive years. A 100 mt/ha application of this sludge resulted in addition of 20 kg Cd/ha, the upper limit allowed by current federal regulations (U.S. EPA, 1979a). Barley was also grown on soils fertilized according to soil test recommendations. Barley leaf and grain and soil samples were collected and analyzed for Cd, Zn, Cu and Ni each year (Tables 5 to 8).

The major conclusions from this experiment are summarized as:

- (1) The metal content of barley grain and tissue were similar for untreated and NPK fertilized soils.
- (2) Yearly variations in plant metal composition were observed for sludge-treated and untreated plots at most locations.
- (3) Metal levels in a plant grown on untreated soil could exceed those found at another location where sewage sludge was added to the soil.
- (4) Cadmium concentrations in barley leaf tissue were greater than those in grain.

- (5) Sewage sludge application increased metal concentrations for soil and plant tissues with a single 100 mt/ha or annual applications of 20 mt/ha.
- (6) The initial application of 20 mt/ha caused a greater increase in plant metal levels than the subsequent 4 annual sludge treatments.
- (7) The increase in 2n and Cd in plants grown on sewage sludge treated soils was greater than for Pb, Cu, and Ni.
- (8) After the fifth year, the concentrations of Cd and Zn in the barley tissue depended only on the total amount of sludge applied and not upon the frequency of application (i.e., 100 mt/ha in year 1 vs. 20 mt/ha in years 1 through 5).

# SOIL PROPERTIES INFLUENCING THE ACCUMULATION OF TRACE ELEMENTS BY PLANTS

The accumulation of trace elements by plants is a reflection of the influence of soil physical properties on plant growth, soil hydraulic properties and chemical properties such as pH, CEC, and clay mineral sorption reactions.

### Physical Properties

Soil particle size distribution (i.e., texture), structure, and depth are important in determining soil hydraulic properties such as porosity, permeability and drainage rates; these properties in turn influence soil moisture content and aeration/respiration which impact the type and rates of both soil microbial activity and chemical reactions, as well as plant root development and growth rates.

Although soil texture, hard pans, and other physical features can be observed in the field or identified from soil maps and likely influence soil chemical reactions, clear identification of these effects in relation to plant uptake of metals has been difficult. Soil texture, however, has been recommended as a quantifiable soil property to limit metal loadings to soils in the Northeast (Baker et al., 1985), although experimental data are not available to support this concept. Presumably as the texture of the soil becomes finer (i.e., greater clay content), the limiting application rates may also increase.

### Soil pH

The impact of pH on metal accumulation by plants has been extensively reviewed (Logan and Chaney, 1983) and little additional pertinent data has been reported which would refute their conclusions. Basically, metal availability (except for Mo and Se) tends to decrease with liming.

Solubility of solid phase minerals including metal carbonates, phosphates, tes, and sulfides is enhanced at low soil pH (Lindsay, 1979); however, the importance of this phenomenon in sewage sludge-treated soils has not been adequately defined by solid phase equilibria studies.

Soil pH is one of the easier characteristics to measure but care must be exercised in interpreting results. The pH measured in a soil-neutral salt suspension will be lower than the pH measured in a soil-water system, although some exceptions exist. When pH was measured at 19 sites over a growing season, a maximum variation of 1.6 pH units was reported for measurements in a 1:2 soil:water suspension or in 0.01 N CaCl $_2$  (Collins et al., 1970). Bates et al. (1982) found similar variations in Ontario, Canada, soils, whether pH was measured in 0.01 N CaCl $_2$  or in a soil-water saturated paste. These authors found greater variation in the pH for soils cropped to corn which received large applications of N fertilizer than for soils cropped to alfalfa.

The water content of the soil and its electrolyte content affects soil pH readings significantly (Thomas and Hargrove, 1984). Bates et al. (1982) measured the pH in 245 soils and obtained the following pH values: pH 5.77  $\pm$  0.91 for a soil:water saturated paste; pH 5.26  $\pm$  0.91 for a 1:2 soil:solution of 0.01 N CaCl<sub>2</sub>; and pH 5.00  $\pm$  0.95 for a 1:2 soil:solution of 1 N KCl. The differences between methods varied with pH, such that: pHH<sub>2</sub>0 = 0.82  $\pm$  0.94 pHCaCl<sub>2</sub>, R<sup>2</sup> = 0.904. This variability emphasizes the need for use of a standardized method to measure soil pH.

Soil organic matter and the resultant impact on pH buffering can influence the effect of liming on trace element uptake. Liming acid soils to pH 6.5 as measured in water paste (1:1 soil-water ratio), often is costly and can require considerable amounts of lime. Some trace element deficiencies (e.g., Fe, and Mn) may occur as pH approaches neutrality. In addition, the increase in soil pH may not reduce markedly the uptake of trace elements from sludge treated soils, especially for crops not accumulating metals. Pepper et al. (1983) attributed the ineffectiveness of liming on reducing Cd uptake by corn to a drop in soil pH during the growing season. Others (Hemphill et al., 1982; Giordano and Mays, 1981) have observed similar effects on metal content of corn grain. In general, lime applications reduce uptake of Zn and Ni more than Cd (Singh and Narwal, 1984).

Metal uptake by plant species may vary in response to liming. Giordano (CAST, 1980) observed that liming reduced In concentrations in soybean seed to a greater extent than in corn grain or cotton seed. Whereas the Ni content of soybean and cotton seed was depressed by lime, the content of Cu in all crops and plant parts was relatively unaffected.

Soil pH influences uptake of most metals at least to some extent, but the current recommendation of pH 6.5 should be reconsidered for food-chain agricultural soils since some reports indicate adequate control of metal uptake at pH 6. For example, Hajjar (1985) conducted a greenhouse study using soils treated with sludge at rates from 0 to

27 mt/ha. Replicate soils were adjusted to pH <5.6, pH 5.7-6.0, and pH >6.4 with either  $\rm H_2SO_4$  or  $\rm Ca(OH)_2$ . For both tobacco and peanut, plant tissue concentrations of Ni, Zn, and Cd decreased with increasing soil pH; however, in general, metal uptake by peanuts was similar at pH >6.4 and pH 5.7-6.0, suggesting that attainment of pH 6.5 is not essential for minimizing plant availability of Ni, Zn and Cd. A soil pH  $\approx$ 6.0 may be equally acceptable to regulate metal uptake. This conclusion is supported by field data with barley (Vlamis et al., 1985). More specific pH values where accumulator crops such as tobacco are grown on highly buffered acid soils may be required.

It has not been demonstrated that pH control to prevent transfer of metals into the food chain is necessary on forested sites. If a forest site is shifted to agricultural use or residential development, the soil pH should be adjusted by limestone addition at that time to meet existing standards.

### Iron

Iron deficiency chlorosis on calcareous soils is a unique soil fertility problem worldwide. Application of sewage sludges can correct Fe chlorosis problems (McCaslin and O'Connor, 1982; McCaslin et al., 1985). In New Mexico, where Fe chlorosis affects large acreages of farmland, sludges applied at 34 to 90 mt/ha increased the levels of plant available Fe, Zn, and P in a severely Fe-deficient calcareous soil. Sorghum grain yields from sludge-treated soils were significantly higher than those receiving dairy manure or chemical fertilizers. Uptake of Zn and Cd by barley was minimal after sludge application to calcareous soils in the regional W-124 study (Tables 5 to 8).

## Molybdenum

Soon and Bates (1985) present data which show that the application of a lime-treated sewage sludge supplying 0.21 kg Mo per hectare per year raised the Mo concentration in both bromegrass and corn stover above the control by significant margins (0.29 vs. 1.16 mg/kg for bromegrass, and 0.20 vs. 0.47 mg/kg for corn stover). A non-significant increase occurred in the Mo content of corn grain even though the lime-treated sludge increased soil pH from 7.4 to 8.1. In the same experiment, application of Al- and Fe-treated sludges raised the Mo concentration in bromegrass from 0.29 to 0.69 and 0.46 mg/kg, respectively. The amount of Mo added with the Al- and Fe-treated sludges averaged 2.18 and 1.66 kg per hectare per year, respectively.

Pierzynski and Jacobs (1986b) applied a sludge containing 1500 mg Mo/kg  $\alpha$  rates of 42 and 94 mt/ha (equivalent to 63 and 141 kg Mo/ha). During the three year study the Mo content of corn seedlings (25-31 cm height) ranged from 47 to 724 mg/kg for those grown with the higher sludge application as compared to a range of 1.9-6.0 mg/kg in those from control plots.

Similar increases were observed for soybean seedling tissue (18-23 cm height) and diagnostic leaf tissue of both crops. At the end of the study, soil pH had increased from 4.6 in the control to 6.9 in the 94 mt/ha sludge treatment. This change in soil pH may explain the increase of Mo concentration in plant tissue. The Mo content of corn grain also increased with time (0.2 to 0.6 mg Mo/kg for control and 2.0 to 6.9 mg Mo/kg for the high sludge rate) but the effect on the Mo concentration in soybean seeds was greater than on corn grain (8.9 to 19.9 mg Mo/kg for control and 122 to 242 mg Mo/kg for the high sludge rate). Elevated Mo levels did not affect growth of either crop.

A greenhouse study has shown that uptake of Mo by ryegrass and white clover was enhanced more by addition of up to 0.41 kg Mo/ha from sludge than from Na molybdate (Williams and Gogna, 1983). However. Mo additions from sludge did not always enhance Mo uptake by corn, soybeans, and alfalfa compared to Na molybdate, when rates of Mo from 60 to 400 kg Mo/ha were used (Pierzynski and Jacobs, 1986a). Results from another study indicated that sludge applications, especially to high pH soils, tended to reduce the Cu/Mo ratio of the affected vegetation (Soon and Bates, 1985). Under these conditions the likelihood of Mo induced Cu deficiency in grazing ruminant animals consuming the forage is enhanced.

### Selenium

Sludge application to agricultural soils did not increase Se uptake by crops (Dowdy et al., 1984; Logan et al., 1987). In these studies, Se inputs ranging from 0.024 to 66 kg/ha did not result in significant accumulation of Se in plant tissue. Unless additional data become available to indicate otherwise, Se should not be a limiting factor in land application of municipal sludges.

## Cation Exchange Capacity

Cation exchange capacity has been used as the primary soil property to govern metal loadings for the past 10-15 years. The basic concept originated in England and was adopted to prevent metal toxicities to crops; however, its use was mainly intended for soils where organic matter contributes a significant fraction to the CEC.

## Method of Analysis

One problem when using CEC to regulate sewage sludge addition to soil is that no single method of determining CEC is universally accepted for its determination. The two most widely used methods are: 1) summation of exchangeable cations, and 2) saturation with either a buffered or unbuffered index cation. The above mentioned methods can give vastly different CEC values for the same soil. Hence, the recommended total metal loading rate and subsequent metal uptake by plants can vary depending upon the method used to determine CEC.

Criteria developed under 40 CFR part 257 of the Resource Conservation and Recovery Act (RCRA) state that the method to be used for CEC analysis should depend upon the type of soil (US EPA, 1979b). For distinctly acid soils the summation method should be used and for neutral, calcareous, or saline soils the sodium acetate method should be used (see Rhoades, 1982). If CEC is to be used as an index for metal loadings, then the method of analysis must be standardized.

### Correlation of CEC and Plant Uptake of Metals

Research on the relationship between CEC and plant uptake of metals has been minimal and results have been conflicting (CAST, 1980; Logan and Chaney, 1983). Hinesly et al. (1982) conducted a study to determine the effect of CEC on Cd uptake by corn. Soil samples of the Bl horizon of the Ava series and the Ap horizon of the Maumee series were separately mixed with samples of the Plainfield series to obtain soil mixtures having a CEC from 5.3 to 15.9 cmol(+)/kg(meq/100 g). Additions of CdCl $_2$  or 100 mt/ha of dried, digested sewage sludge were used to provide a soil-Cd concentration of 10 mg/kg. Corn was grown in pots of each mixture and harvested at 3- and 7-week intervals for tissue analyses. The soil CEC inversely affected the uptake of Cd by corn when Cd was supplied as a soluble salt, but not when it was supplied as a constituent of municipal sewage sludge. This conclusion was confirmed in greenhouse studies conducted by Korcak and Fanning (1985).

In terms of phytotoxicity, research data available indicates that the maximum metal loadings allowed in the CEC-metal limit approach are conservative. Furthermore, a large degree of safety is provided by the CEC-metal limit approach. For example, no phytotoxicities have occurred in studies where the total metal loading equals or greatly exceeds those recommended in the CEC table at pH 6.5 (Chang et al., 1983; Ellis et al., 1981; Hinesly et al., 1984a; Vlamis et al., 1985). These observations indicate that the present practice of using CEC as a basis for establishing metal-loading limits should be abandoned.

#### CONCLUSIONS

The following conclusions are supported by previous literature or by new information reported in this total report.

 Conclusions on the impact of sewage sludge on trace element uptake by plants should be based on field studies rather than greenhouse or pot studies. Plant tissue concentrations obtained in greenhouse or pot studies may not be representative of those obtained in the field unless root growth is not restricted and accumulation of soluble salts is avoided.

- 2. Concentrations of trace elements in crops grown in soils treated with metal salts exceed those of crops grown on sewage sludge-amended soils and should not be used to predict dietary intake.
- 3. Soil physical properties are related to trace element uptake by plants but current data does not allow quantification of these relationships.
- 4. Due to natural variations in soil and crop characteristics, trace element content of crops grown on untreated soils differ.
- 5. The soil pH value used in conjunction with application of sewage sludge should be measured on the sewage sludge-soil mixture or on the untreated soil using a 1:1 soil water ratio, realizing that 0.01 M CaCl<sub>2</sub> is a preferred matrix to compensate for soluble salt levels in sewage sludge or soils.
- 6. The impact of the pH reduction on increased metal uptake is more marked with high metal sludges and crops responsive to metal additions.
- 7. The relationship of either CEC or texture to metal uptake in sewage sludge-amended soils has not been conclusively demonstrated under field conditions. The current guidelines that are based on the use of CEC to limit metal additions to soils are not supported by current long-term field experimentation.
- 8. Trace element deficiencies rather than toxicities are a major concern in soils containing free  $CaCO_3$ . Sewage sludge additions have been used to correct Fe deficiency in calcareous soils. Based on plant uptake, Mo is the principal metal of concern in calcareous soils treated with sewage sludge.

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TABLE 1. METAL AND ORGANIC CARBON CONTENTS, CEC, AND pH FOR SOILS FROM SELECTED SITES IN THE CONTINENTAL UNITED STATES (FROM HOLGREN ET AL., 1985).\*

State	Nt	Mean	d SD≢	P Mean	2D	Meai	Zn n SD		u SD	Mean	Ni SD	Mear	H SD	Orga Mean	nic C SD	Mear	EC SD
						<del></del>									%)		(+)/kg)
Arizona	14	0.24	0.06	14	4	72	18	39.4	11.3	28.7	6.5	7.7	0.5	0.37	0.14	143	36
Iowa	85	0.24	0.07	14	4	62	16	21.3	5.7	28.2	8.1	5.9	0.7	2.53	0.94	282	77
Missouri	33	0.27	0.08	20	5	60	9	18.8	3.8	24.8	3.5	6.5	0.7	1.80	0.61	200	46
Minnesota	89	0.30	0.09	12	2	71	20	22.3	4.5	30.0	5.5	5.9	0.8	3.01	0.90	342	89
California	283	0.31	0.28	12	12	93	41	46.6	30.2	74.3	63.9	7.2	0.8	1.00	0.88	214	140
Kansas	38	0.32	0.07	15	2	53	10	15.6	2.5	20.4	3.4	5.6	0.9	1.14	0.18	192	25
W.Virginia	40	0.32	0.34	646	1127	84	37	96.9	143	23.3	13.1	5.3	0.7	2.99	2.19	141	78
Wisconsin	164	0.35	0.20	12	7	59	29	37.7	36.8	16.4	9.4	5.9	0.8	14.64	16.45	556	552
Montana	33	0.37	0.07	11	2	75	13	21.0	4.6	26.1	5.5	6.8	0.9	1.41	0.38	171	35
N.Dakota	30	0.37	0.21	10	5	69	42	22.0	12.9	31.1	16.9	7.1	0.8.		0.76	260	151
Idaho	54	0.38	0.16	11	2	68	24	22.0	' 8.0	25.2	7.9	7.4	1.0	1.16	0.54	173	39
Ohio	81	0.38	0.15	19	4	89	41	28.1	11.7	28.2	9.7	6.4	0.6	1.83	0.54	189	58
Colorado	89	0.39	0.30	16	13	85	42	19.4	8.0	15.9	7.2	7.7	0.5	0.80	0.29	137	49
Nebraska	72	0.39	0.18	14	3	58	25	17.3	7.7	21.7	12.3	6.4	0.8	1.49	0.48	199	51
Florida	89	0.44	0.28	10	10	88	67	103.7	88.2	10.3	7.7	5.8	0.9	26.80	18.11	970	657
New York	173	0.45	0.36	17	5	64	37	74.8	77.5	19.5	10.1	5.4	0.8	16.71	17.32	767	769
Oregon	106	0.49	0.45	11	7	71	30	33.4	16.5	27.1	6.0	6.1	1.0	3.36	5.03	338	279
S.Dakota	44	0.56	0.12	14	3	96	25	30.3	8.8	42.3	18.8	6.5	0.9	2.62	0.49	300	49
Michigan	86	0.94	0.30	16	6	80	37	111.5	75.6	14.7	8.6	5.7	0.7	28.43	5.13	1358	315
S.Carolina	10	0.03	0.01	10	3	12	5	4.1	1.7	4.1	1.8	4.2	0.3	2.27	1.00	81	22
Georgia	146	0.05	0.05	8	5	18	19	7.0	5.3	9.0	7.9	5.9	0.5	0.74	0.26	35	15
Alahama	92	0.06	0.07	7	4	16	7	8.1	5.9	11.3	6.8	5.8	0.7	0.65	0.30	31	19
Maryland	57	0.08	0.02	ıí	4	31	16	8.1	2.6	12.4	4.4	5.7	0.7	0.75	0.17	44	21
N.Carolina	242	0.09	0.07	10	6	15	14	8.9	12.9	8.6	13.1	5.1	0.5	0.69	1.74	74	84

(continued)

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TABLE 1 (continued)

			<u>d</u>	P			Zn		u		Ni		Н		nic C		CEC
State	Nt	Mean	SD#	Mean	SD		n SD	Mean	SD	Mear	SD	Mean	SD	Mean		Mear	
						- (mg/k	g) <b>-</b> -							(	(1)	(mmol(	(+)/kg
Oklahoma	94	0.10	0.08	7	3	33	47	4.0	13.3	14.9	14.4	6.4	0.6	0.65	0.23	94	54
Virginia	46	0.14	0.07	98	118	59	29	9.4	28.4	22.3	12.0	5.6	0.8	2.07	0.97	92	34
Texas	362	0.16	0.11	9	5	40	27	2.2	6.9	15.9	10.2	7.1	1.0	0.78	0.41	153	105
Delaware	4	0.17	0.06	10	2	- 25	9	5.0	2.2	6.6	4.4	6.3	1.1	0.55	0.22	4	1
Maine	31	0.17	0.03	13	. 2	74	13	0.7	24.0	41.5	6.4	4.5	0.5	2.25	0.43	134	18
Arkansas	62	0.18	0.15	15	8	45	33	5.5	8.8	17.2	9.8	5.7	0.3	1.07	0.32	145	97
Illinois	135	0.20	0.09	16	3	56	21	7.2	5.6	20.6	7.0	6.0	0.9	1.79	0.77	195	82
New Mexico	41	0.20	0.07	11	3	47	14	6.1	5.7	16.9	5.0	8.2	υ.7	0.58	0.19	142	48
Washington	122	0.20	0.08	9	4	66	19	7.3	10.5	29.0	16.7	6.3	0.1	1.48	2.71	141	59
Pennsylvania	45	0.21	0.24	24	25	30	27	5.3	28.4	10.4	7.8	6.1	0.9	1.36	0.59	90	24
Louisiana	133	0.22	0.14	16	16	64	51	2.1	16.6	25.3	18.7	5.7	0.6	1.37	0.65	238	174
Indiana	80	0.23	0.14	13	5	51	25	7.0	9.8	16.5	8.6	5.7	0.7	1.35	0.55	130	57

<sup>\*</sup>Soils were selected from sites removed from mobile and point source contamination; values reported for metals may not be representative of U.S. soils in general. Data are expressed on a dry weight basis.

tN = number of soil sites analyzed

**<sup>\*</sup>SD** = standard deviation

TABLE 2. TRACE ELEMENT CONCENTRATIONS FOR SOILS FROM SELECTED SITES IN THE CONTINENTIAL UNITED STATES (HOLMGREN ET AL., 1985).\*

	Cd	Pb	Zn	Сu	Ni
			- mg/kg -		
Mean	0.27	17	57	30	24
Median	0.20	11	54	19	18
Geometric mean	0.17	11(16)†	43(48)	18(17)	16(13)
Std. deviation	0.26	141	39`	42`´	27` ´
Maximum	2.3	4,109	402	735	269
Minimum	0.01	0.2	1.5	0.3	0.3
50th percent.	0.20	11	54	19	18
95th percent.	0.79	26	127	98	56

<sup>\*</sup>Soils were selected from sites removed from mobile and point source contamination; values reported for metals may not be representative of U.S. soils in general. Data are expressed on a dry weight basis.

<sup>†</sup>Geometric mean of U.S. soils from Shacklette et al. (1984).

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TABLE 3. TRACE ELEMENT CONCENTRATIONS (DRY WEIGHT) IN THE EDIBLE PART OF CROPS GROWN ON UNTREATED SOILS.\*

Crop	Cd†				Zn≠				Pbt			
	Min	Max	Media	n 95th§	Min	Max	( Media	n 95th	Min	Max	Media	an 95th
	µg/kg			mg/kg				μg/kg				
Lettuce	34	3800	435	2100	13	110	46	78	36	1700	190	994
Spinach	160	1900	800	1480	17	200	43	128	240	2300	530	1180
Potatoes	9	1000	140	360	5.1	35	15	27	1	2200	25	97
Wheat	5	220	36	125	11	76	29	48	1	770	21	168
Rice	<1	250	5	34	7.7	23	15	20	<1	80	5	26
Sweet corn	0.5	230	8	57	28	55	25	46	7.6	260	9	62
Field corn	<1	350	4	67	12	39	22	30	<1	<b>3</b> 600	6	32
Carrots	15	1200	160	786	3.8	61	20	48	10	1100	55	236
Onions	11	340	90	240	6.1	33	16	26	2	720	38	95
Tomatoes	45	790	220	610	12	35	22	29	<1	460	27	108
Peanuts	11	660	68	219	17	63	31	42	< <u>1</u>	200	8	27
Soybeans	î	1200	45	180	32	70	45	59	3	350	36	99

<sup>\*</sup>Data are for crops grown in areas removed from mobile and point sources of contamination.

#Wolnick et al. (1985)

§95th percentile

<sup>†</sup>Wolnick et al. (1983a,1983b)

TABLE 4. GENERAL CHARACTERISTICS OF SOILS USED IN W-124 STUDY.

Location	Soil	Sand	Silt	Clay	Cation Exch. capacity	Organic C
			g/kg		(mmo1(+)/kg)	(g/kg)
AL	Decatur clay (Rhodic Paleudult)	240	340	420	91	11.3
AZ	Pima clay loam (Typic Torrifluvent)	330	340	330	280	11.0
CA(D)	Domino loam (Xerollic Calciorthid)	200	380	420	140	8.9
CA(G)	Greenfield sandy loam (Typic Haploxeralf)	100	270	630	90	5.1
CO	Nocono clay loam (Aridic Argiustoll)	250	400	350	230	13.2
FL	Lake fine sand (Typic Quartzipsamment)	942	35	23	14	7.0
IL	Ipava silt loam (Aquic Argiudoll)	-	-	-	266	35.1
IN	Chalmers silt loam (Typic Haplaquoll)	120	$\epsilon$ 10	270	251	23.7
MD*	Christiana sandy loam (Typic Paleudult)	-	•	-	<b>5</b> 9	10.5
MI	Celina silt loam (Aquic Hapludalf)	510	380	110	120	12.3
MN	Port Byron silt loam (Typic Hapludoll)	90	660	250	220	24.4
NE	Sharpsburg silty clay (Typic Argiudoll)	50	610	340		20.3
	Celina silt loam (Aquic Hapludalf)	230	610	160	120	13.5
OR	Willamette silt loam (Pachic Ultic Argixeroll	360	442	198	150	13.4
	Millville silt loam (Typic Haploxeroll)	340	530	130	190	11.6
WI	Plano silt loam (Typic Argiudoll)	40	820	140	197	26.2

<sup>\*</sup>MD(L) and MD(H) refers to unamended and  $CaCO_3$  amended soils, respectively.

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TABLE 5. CONCENTRATIONS OF Cd IN DTPA SOIL EXTRACTS AND IN LEAF AND GRAIN OF BARLEY GROWN IN 15 LOCATIONS.

		079		Ban	ley Lea	f	Barley Grain			
Location*	Year	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK
				*	1110711077	maying		me/ma/y	1007110	
						-mg/kg				
AZ	1	1.64	4.86	0.30	0.61	1.19	0.46	0.21	0.32	0.03
•	2	1.59	4.00	0.03	2.88	3.46	0.83	0.05	0.11	0.03
	3	3.00	4.03	0.25	1.43	2.47	1.95	0.04	0.06	0.05
	4	3.41	3.52	0.25	1.63	2.13	2.00	0.03	0.03	0.03
	5	3.52	3.63	0.19	NOT	ND	NO	NO	NO	NO
CA(G)	1	0.83	3.85	0.10	0.07	0.08	0.03	0.08	0.16	0.03
	2	0.95	1.35	0.10	0.03	0.03	0.03	0.03	0.04	0.03
	3	1.45	1.40	0.10	0.03	0.03	0.03	0.05	0.04	0.04
	4	1.52	1.26	0.10	0.03	0.04	0.03	0.09	0.04	0.06
	5	2.39	1.18	0.06	0.13	0.24	0.06	0.05	0.03	0.05
CA(D)	1	0.75	4.80	0.10	0.03	0.07	0.03	0.03	0.06	0.03
• •	2	1.48	3.05	0.10	0.03	0.03	0,03	0.03	0.03	0.03
	3	2.05	2.40	0.10	0.03	0.03	0.03	0.04	0.03	0.03
	4	1.95	1.77	0.10	0.03	0.03	0.03	0.03	0.03	0.03
	5	2.13	1.40	0.06	0.04	0.33	0.05	0.09	0.08	0.03
СО	1	ND	ND	NO	ND	ND	ND	0.08	0.16	0.05
	1 2 3	0.34	1.65	0.09	0.33	0.55	0.25	0.16	0.22	0.12
	3	0.50	2.05	0.13	ИD	MD	Ю	0.11	0.33	0.04
	4	1.77	2.29	0.12	ND	ND	ND	0.27	0.73	0.03
	5	1.25	2.63	0.21	NO	ND	NO	0.03	0.03	0.03
FL	1	0.76	4.58	0.06	0.44	0.67	0.33	0.11	0.13	0.03
	. 2	2.06	4.50	0.07	0.37	0.45	0.30	0.25	0.38	0.06
	3	2.93	4.05	0.05	0.69	0.73	0.20	0.32	0.47	0.03
	4	4.35	5.45	0.15	ND	ND	NG	0.27	0.37	0.09
	5	2.70	3.70	0.17	0.15	0.19	0.07	0,66	0.85	0.16
IN	1	ON	0.39	0.51	ND	NO	ND	ND	0.36	0.07
	2	NO	0.41	0.16	ND	סא	ON	סא	0.86	0.3
	2 3 4	ND	2.64	0.12	NO	NO	NO	ND	0.50	0.1
	4	ND	2.41	0.14	ND	0.43	0.33	ND	0.25	0.13
	5	ND	5.35	0.24	ND	0.22	0.08	NO	0.19	0.0
MD(L)	1	ND	OM	ND	0.86	1.22	0.13	0.27	0.78	0.0
	1 2 3	0.70	3.75	0.11	0.41	1.22	0.36	0.21	0.54	0.13
	3	0.80	3.04	0.09	0.22	0.43	0.11	0.30	0.32	0.1
	4	2.13	3.20	0.06	0.80	2.65	0.28	0.54	0.43	0.0
	5	3.54	3.13	0.06	0.56	0.79	0.35	0.32	0.42	0.0

TABLE 5 (continued)

			PA Extra			ley Lea		Bar	ley Gra	in
Location*	Year	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK
						mg/kg -				-,
MD(H)	1	NO†	ND	ND	0.49	1.21	0.15	0.25	0.59	0.06
	2	0.91	4.09	0.06	0.46	0.92	0.18	0.11	0.33	0.03
	3	0.83	2.57	0.05	0.58	0.17	0.07	NO	0.26	0.20
	4	2.28	3.08	0.07	0.60	0.68	0.18	0.40	0.46	0.07
	5	4.09	3.34	0.06	0.41	0.27	0.18	0.30	0.31	0.09
MI	1	0.68	2.15	0.28	0.89	2.39	0.85	0.55	1.08	0.65
	2	1.70	3.43	0.08	0.82	1.15	0.59	0.93	0.98	0.52
	3	1.55	2.47	0.36	ND	NO	ND	NO	MD	ND
	4	ND	NO	NO	1.20	1.13	0.61	0.84	0.85	0.72
	5	3.14	3.49	0.38	1.30	0.64	0.41	1.35	0.91	0.43
MN .	1	0.77	3.85	0.08	0.46	1.78	0.14	0.12	0.36	0.07
	2	2.27	5.58	0.17	0.34	0.65	0.14	0.18	0.33	0.03
	3	2.23	3.06	0.20	0.39	0.53	0.07	0.18	0.21	0.07
	4	2.59	4.97	0.24	0.33	0.34	0.12	0.23	0.28	0.05
	5	3.61	3.50	0.25	0.21	0.21	0.08	0.22	0.24	0.06
NE	1 2 3 4 5	2.90 0.88 4.00 4.62 2.55	5.00 1.39 3.89 4.53 3.82	. CM . OM OM OM OM	0.68 0.07 0.03 0.20 0.14	1.31 0.12 3.03 0.19 0.10	NO NO NO NO	0.95 0.25 0.03 0.18 0.06	1.70 0.42 0.03 0.18 0.07	NO NO NO NO
ОН	1	1.02	3.30	0.15	0.28	0.81	0.15	0.21	0.36	0.12
	2	2.20	5.11	0.18	0.34	0.64	0.18	0.26	0.32	0.13
	3	2.42	3.63	0.25	0.38	0.42	0.23	0.15	0.17	0.04
	4	2.83	3.10	0.25	0.27	0.22	0.14	0.11	0.08	0.03
	5	3.43	1.68	0.40	0.78	0.76	0.67	0.07	0.05	0.04
OR	1 2 4 5	ND ND 1.77 3.36	ND ND 1.06 2.75	ND ND 0.27 1.05	ND NO NO 0.36	ND ND ND 0.34	ND ND ND 0.26	0.23 0.03 0.55	0.40 0.03 0.66	0.12 0.03 0.41
UT	1	1.95	9.23	0.33	ND	ND	ND	0.21	0.63	0.11
	2	2.08	4.57	0.22	0.44	0.57	0.36	0.16	0.41	0.08
	3	3.33	5.68	0.30	0.56	0.49	0.50	0.14	0.23	0.05
	4	3.90	4.47	0.40	0.74	0.48	0.55	0.18	0.21	0.13
	5	6.58	6.13	0.57	1.47	2.79	0.65	0.17	0.32	0.11
WI	1	1.42	4.16	ND	0.33	0.74	0.21	0.27	0.47	0.17
	2	1.73	5.07	0.17	0.25	0.79	0.15	0.21	0.55	0.11
	3	1.43	5.62	0.29	0.36	0.79	0.27	0.20	0.45	0.14

<sup>\*</sup>See Table 4 for description of soils.

tND signifies not determined.

TABLE 6. CONCENTRATIONS OF Zn IN DTPA SOIL EXTRACTS AND IN LEAF AND GRAIN OF BARLEY GROWN IN 15 LOCATIONS.

			PA Extra		Bar	ley Lea		Barley Grain		
Location*	Year	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK
						mg/kg -				
		17.0	<b>50.5</b>		46.4		20.6	r, r	<b>.</b>	
AZ	1	17.9	59.5	5.1	46.4	58.1	30.6	51.5	59.2	46.7
	2 3	19.3	45.4	1.1	53.7	71.2	38.9	55.4	58.9	40.4
	3	44.2	52.9	4.4	35.5	36.7	16.4	55.7	66.4	46.7
	4	44.3	54.8	3.5	31.4	36.8	19.2	37.4	37.4	40.3
	5	46.6	55.1	3.3	ND †	ND	ND	ND	ND	NO
CA(G) ·	1	14.7	35.0	2.0	36.3	39.3	28.8	45.8	55.8	33.8
	2 3	18.3	23.6	1.8	22.0	29.5	20.8	34.0	40.5	26.3
	3	18.5	19.2	1.6	39.1	33.4	32.8	37.2	31.9	29.8
	4	21.4	15.8	1.6	32.4	24.5	26.6	33.3	39.4	28.9
	5	37.1	17.3	1.6	24.9	19.7	20.1	32.8	32.4	27.1
CA(D)	1	13.0	43.8	1.5	21.0	34.3	21.8	37.8	49.5	31.3
(-)	2	34.0	55.9	2.3	25.0	27.8	22.5	34.3	37.0	37.3
	2	26.4	27.5	1.5	37.1	36.3	30.0	41.5	42.0	32.0
	4	26.5	22.4	1.7	25.8	23.2	20.1	30.6	33.7	25.
	5	30.8	19.0	1.6	26.3	23.6	21.5	47.0	39.5	26.
co	1	ND	NO	NO	NO	NO	ND	57,8	58.8	51.
••	2	4.7	23.0	2.2	45.4	60.7	25,7	66.5	64.5	56.
	2	6.7	28.0	2.8	ON	NO	NO	51.8	66.8	41.8
	4	14.5	38.9	2.4	ND	ND	NŪ	4.8	76.1	47.
	5	18.9	40.0	2.6	МО	ND	NO	100.0	112.0	60.
FL	1	11.3	74.5	1.7	23.9	33.4	13.9	13.5	18.0	10.
		31.3	69.5	2.2	66.0	70.8	36.0	77.3	65.3	41.
	2	48.3	76.5	2.5	153.1	93.3	38.5	55.3	63.8	43.
	4	60.5	76.5	2.9	ND	ON	ND	67.1	73.8	30.
	5	41.1	61.8	2.5	30.9	28.7	20.8	85.0	94.1	50.
IN	1	ND	119.3	3.6	ND	ND	ND	ND '	82.1	57.
<del>-</del> · ·		ND	33.1	1.3	ND	ОИ	ND	OM	60.2	37.
	2 3 4	ND	31.0	2.0	ND	NO	ND	ND	61.6	35.
	4	ND	41.4	3.3	ND	31.4	21.0	ND	43.5	27.
	5	ND	68.5	2.3	ND	42.9	21.8	ND	44.0	31.
MD(L)	1	ОN	МО	NO	25.3	69.9	17.4	38.6	52.1	23.
\_ ,	2	9.3	53.6	2.4	21.4	58.7	17.0	32.2	46.7	27.
	2 3.	10.4	35.0	3.0	26.3	32.6	21.9	37.4	38.9	25.
	4	23.7	39.5	3.4	54.0	112.4	17.2	55.6	66.1	34.
	Ś	32.5	33.1	1.9	20.0	23.0	14.9	35.1	40.7	21.

TABLE 6 (continued)

			PA Extr	act		rley Lea			rley Gr	ain
Location*	Year	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NFK
						mg/kg ~				
MD(H)	1	NOT	NO	NO	25.0	74.9	18.5	33.0	49.7	24.4
	2	8.4	54.5	1.6	19.6	32.8	11.3	27.9	38.2	22.1
	3	9.6	32.4	1.5	22.4	26.7	17.4	NO	30.5	20.5
	4	23.1	31.3	1.8	40.7	43.0	13.4	45.3	43.8	27.6
	5	64.5	30.0	1.3	20.9	23.4	14.7	33.4	35.1	16.5
MI	1	8.4	25.8	1.5	67.9	112.3	48.7	64.8	78.5	47.5
	2	21.2	44.3	1.8	48.3	63.2	27.9	110.0	85.3	59. <b>5</b>
	3	24.3	35.7	3.9	ND	NO	NO	NO	NO	NO
	4	NO	Ю	ND	89.2	52.1	43.4	65.7	64.7	46.8
	5	52.8	56.4	5.4	79.5	57.5	44.0	118.3	155.0	121.5
MN	1	32.5	162.7	5.0	46.3	106.0	24.3	67.0	111.8	48.5
	2	34.0	85.5	2.0	38.8	52.0	28.5	67.5	80.3	41.3
	3	33.5	43.3	2.1	41.1	44.3	28.9	<b>69.3</b>	70.1	52.0
	4	35.8	67.4	2.8	45.0	57.7	40.7	49.4	46.4	36.0
	5	73.7	56.7	2.3	39.2	34.3	25.5	65.0	67.1	49.6
NE	1	31.5	102.0	NO	39.0	96.0	NO	94.0	118.0	DM
	2	18.1	54.8	NO	19.0	26.7	NO	98.1	111.0	ND
	3	45.5	43.8	NO	ND	ND	ND	64.3	63.5	NO
	4	59.4	62.4	NO	22.7	18.9	ND	75.5	69.5	ND
	5	54.8	52.6	NO	25.6	24.7	ND	49.7	47.7	NO
ОН	1	23.9		9.4	30.0	40.5	26.0	53.0	70.3	47.3
	2	37.8	70.2	9.5	32.2	32.3	24.8	50.1	49.9	35.8
	3	41.9	49.2	8.3	31.0	28.2	24.7	36.0	36.0	28.1
	4	55.8	36.0	19.9	35.6	35.9	29.6	38.0	32.6	27.8
	5	43.6	21.9	10.1	19.5	15.6	18.6	24.8	23.6	19.9
OR	1	МО	NO	ОИ	OM	ND	ND	18.3	21.9	13.4
	2	ND	DM	NO	NO	NO	ND	56.0	70.0	52.5
	4	31.4	22.3	8.8	ND	ND	ND	53.3	57.8	43.8
	5	55.3	40.2	18.9	56.3	41.1	57.0	68.2	57.6	63.0
UT	1	29.0	129.8	2.5	СИ	NO	ND	52.0	99.5	48.5
	2	29.9	58.5	2.8	26.0	32.3	21.5	65.3	77.5	47.5
	3	46.5	75.8	3.6	29.3	29.3	43.3	61.5	65.0	44.0
	4	50.4	57.5	5.4	37.3	35.8	22.0	65.5	58.8	71.0
	5	78. <i>€</i>	72.5	4.8	52.5	108.0	57.5	57.8	62.3	68.3
WI	1	29.6	74.6	3.5	61.2	80.4	54.4	54.4	78.4	36.5
	2	30.0	89.8	3.7	38.0	71.8	57.8	57 <b>.8</b>	86.4	36.1
	3	23.6	95.1	3.9	28.5	42.1	53.5	53.5	68.8	40.3

<sup>\*</sup>See Table 4 for description of soils.

<sup>†</sup>ND signifies not determined.

TABLE 7. CONCENTRATION OF CU IN DTPA SOIL EXTRACTS AND IN LEAF AND GRAIN OF BARLEY GROWN IN 15 LOCATIONS.

	DTPA Extract				Bar	·ley Lea	Barley Grain			
Location*	Year	20 mt/ha/y	100 mt/ha	NPK	20 mu/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK
						- mg/kg				
AZ	1	6.4	19.5	7.1	8.6	9.0	7.6	9.1	9.6	8.9
	2	8.3 13.8	15.5 17.7	2.4 3.8	10.4 12.4	12.1 11.8	9.9 12.4	10.8 13.4	11.8 14.1	9.9 17.
	4	13.8	15.7	2.8	10.9	11.8	11.9	12.7	13.2	13.
	5	14.6	15.7	2.5	NDT	ND	ND	NO	ND	ND
CA(G)	1	4.7	20.5	1.0	8.3	8.4	6.4	4.4	4.2	2.
	2	5.5	6.1	1.1	6.8	7.0	5.7	2.5	3.0	1.9
	3 4	5.8 6.0	5.3 4.3	1.0	7.8 12.0	6.1 9.0	6.2	3.9	3.6	3.6
	5	9.7	4.3	0.8	10.1	8.8	8.4 8.4	4.5 9.6	4.7 10.1	2.8
CA(D)	1	4.9	26.8	1.4	6.7	9.1	6.1	3.6	4.0	3.2
	2 3	10.1	17.9	1.8	6.9	7.0	6.5	3.2	3.9	4.1
	3 4	8.9	10.3	1.4	10.4	8.8	7.7	3.7	3.4	3.1
	5	7.8 9.2	7.1 5.9	1.4	8.3 11.4	6.3 10.9	7.7 12.2	4.6 18.4	6.0 17.0	5.4 15.8
co	1	NO	NO	ND	מא	ND	NO	12.5	15.9	13.8
	2 3	3.2	9.1	2.3	9.5	11.7	9.6	44.4	39.7	44.8
	3 4	4.2 7.2	12.3 17.2	2.9 2.5	ND ND	ND ND	ND Nů	6.4 6.7	5.3 6.2	5.3 5.6
	5	6.1	18.9	2.0	ND	ON	סא	14.2	13.1	8.2
FL	1	4.1	26.5	0.5	5.5	7.2	3.0	NG	NO	ОМ
	2	10.5	25.2	1.3	9.4	10.8	5.4	4.8	5.2	3.
	3 4	13.9 14.7	19.8 23.5	0.6 0.7	11.8 ND	9.3 ND	4.7 ND	4.0 ND	4.1 2.4	0.
	5	15.7	22.8	0.5	6.9	5.7	4.7	6.5	6.4	2.9
IN	1	ND	30.4	4 1	ND	סא	ND	סא	5.7	4.6
	2 3 4	ND	3.8	2.0	NO	ND	ND	ND	7.4	6.
	3	ОИ ОИ	18.0	3.2 3.7	DN DN	ND 5.1	ND 3.3	ND ND	5.9 5.0	4. 3.
	5	טא	24.7	3.5	NO	6.7	5.6	DN	7.1	6.
MD(L)	1	ND	NO	ND	8.9	13.1	5.0	4.9	3.9	2.
	2	3.7	1.8	1.1	5.9	9.7	5.4	4.4	5.7	3.
	3	4.5 11.8	14.9 17.6	1.3 1.8	5.7 11.5	7.2 12.5	5.4 6.3	7.7 6.1	4.4 2.9	3. 4.
	5	13.1	13.6	1.0	6.9	5.5	3.9	5.5	5.0	2.

TABLE 7 (continued)

			A Extra			ley Lea	f		ley Grai	
Location*	Year	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK
						- mg/kg	, <i>-</i>			
MD(H)	1	ND+	NO	ND	8.9	14.2	6.7	5.2	5.1	3,6
	2	4.2	19.7	0.9	6.2	7.6	4.5	2.6	4,5	3.7
	2	4.7	13.4	0.9	5.5	6.6	4.9	ЙĎ	5.9	3.3
	4	11.7	17.5	1.2	12.3	12.8	6.4	5.4	7.0	4.6
	5	14.0	14.4	0.9	5.4	4.9	4.1	4.5	5.2	2.8
MI	1	2.8	9.6	0.6	27.9	27.4	31.3	22.3	14.0	26.0
	2	9.2	19.8	0.7	13.4	14.0	13.2	19.0	23.8	22.4
	3	7.5	11.5	1.5	ND	NO	ND	NO	ND	ND
	4	ND	ND	ND	37.0	25.0	27.3	18.0	16.1	21.1
	5	15.3	17.1	2.0	17.9	18.3	13.6	32.4	19.3	16.5
MN	1	8.8	43.3	2.0	10.4	11.1	16.8	0.5	0.5	3.8
	2 3	8.5	21.8	1.0	5.7	7.1	4.8	0.5	0.5	0.5
	3	8.1	11.4	0.9	6.9	7.4	6.7	2.5	2.0	3.4
	4 5	8.4 15.8	16.8	1.1	8.8	10.1 9.8	7.4 7.2	3.2	3.0	3.0
	5	15.0	11.3	1.3	7.4	9.0	1.2	6.0	5.9	5.9
NE	1	8.6	24.3	ND	5.6	8.3	ND	10.2	9.8	NO
	2	4.9	13.2	ND	5.8	7.2	ND	8.7	13.6	סא
	3	10.3	11.6	ND	6.8	9.0	HD	4.3	4.3	ND
	4	14.2	15.2	NO	4.8	5.0	ND	4.1	5.6	ND
	5	14.1	15.9	ND	4.7	4.6	ND	2.6	3.6	Ю
OH .	1	8.0	19.2	2.9	26.8	34.3	24.0	5.4	6.4	5.1
	2	13.1	27.9	2.5	19.4	16.1	22.1	7.3	7.2	5.7
	3	13.5	20.0	2.7	21.9	18.1	18.6	4.8	4.1	3.8
	4 5	14.2 14.7	16.0	3.5	23.9	27.3	25.8	3.6	3.3	3.0
		14./	7.8	3.0	5.8	5.5	5.5	2.7	2.7	2.6
OR	1	ND	GM	DM	ND	ND	ND	4.5	5.0	2.3
	2	ND	ND	ND	ND	ND	ND	ND	NO	NO
	4	9.5	7.0	2.2	NO	NO	ND	4.2	5.1	2.4
	5	16.9	14.2	4.7	9.5	7.5	8.1	5.0	4.8	4.5
UT	1	13.2	54.0	1.4	ND	ON	, ND	7.0	12.8	5.8
	2 3	12.6	24.2	1.3	9.7	9.5	8.3	7.6	7.6	5.7
	3	17.4	27.6	1.3	9.7	9.8	13.0	6.1	6.3	5.2
	4 5	19.3 30.3	22.6 28.8	2.0 1.6	12.6 10.1	11.4 13.1	8.8 8.5	6.0 5.4	5.7 6.0	5.2 4.8
			40.0	1.0	10.1				0.0	
WI	1	9.7	24.8	1.3	12.4	12.7	9.6	7.1	8.6	5.5
	2	10.1	29.4	1.7	9.4	12.2	7.7	5.5	7.5	4.8
	3	6.9	28.7	1.9	10.5	11.0	9.2	5.9	6.8	4.9

<sup>\*</sup>See Table 4 for description of soils.

<sup>&</sup>lt;sup>†</sup>ND signifies not determined.

TABLE 8. CONCENTRATIONS OF Ni IN DTPA EXTRACTS AND IN LEAF AND GRAIN OF BARLEY GROWN IN 15 LOCATIONS.

		DTPA Extract			Bar	ley Leaf	-	Barley Grain		
Location*	Year	20 mt/ha/y	100 · mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100 mt/ha	NPK
						- mg/kg	<b>-</b> .			
AZ	1	4.9	7.9	1.3	4.1	4.6	3.2	3.5	4.5	2.9
	2	1.5	3.3	0.1 .	2.2	2.3	1.3	6.5	6.5	5.
	4	2.5 2.3	2.3	0.7	7.7 10.2	7.2 4.5	7.2 5.2	1.7 1.7	1.7 1.7	1.
	5	2.4	2.7	0.6	NOT	NO -	ND	NO	ND	ND
CA(G)	1	1.8	7.6	0.6	ND	ОМ	ND	ND	. ND	ND
	2 3 4	2.5	2.9	0.8	0.1	0.1	0.1	0.1	0.1	0.
	3	2.2	1.9	0.4 0.4	2.0 ND	2.0 ND	2.0 ND	2.0 ND	2.0 ND	2. ND
	5	1.9 4.1	1.2 1.7	0.7	1.0	1.0	1.0	1.0	1.0	1.
CA(D)	1	1.6	9.3	0.6	0.1	0.1	0.1	0.1	0.1	0.
• •	1 2 3 4	2.9	5.8	0.6	0.1	0.1	0.1	0.1	0.1	0.
	3	2.4 1.8	2.5 1.3	0.4 0.4	2.0 ND	2.0 ND	2.0 ND	2.0 ND	2.0 ND	2. ND
	5	3.0	1.8	0.8	1.0	1.0	1.0	1.0	1.0	1.
co	1	ND	ND	NO	:«D	NO	ND	1.6	1.4	1.
	2	0.8	2.2	0.5	1.3	1.4	0.7	1.6	2.0	2.
	2 3 4	0.9 1.9	2.8 4.8	0.7 0.6	DN CN	ND ND	ND ND	0.9 0.7	0.8 1.1	0. 0.
	5	1.4	7.8	0.6	ND	ND	ON	0.7	3.7	1.
FL	1	1.3	19.3	ND	ND	ND	ND	ND	ND	ND
	2	2.3	5.3	0.2	ND	ND	ND	ND	ND	NO
	1 2 3 4	2.8 5.7	5.0 5.6	0.2 0.4	NO ND	ND ND	ND ND	OM ND	DN DN	DM DM
	5	4.2	4.5	0.2	0.1	0.1	0.1	0.1	0.1	o.
IN	1	ND	43.3	6.8	ND	ND	ND	ND	2.7	1.
	2	ND	4.0	7.8	ND	NO NO	ND ON	ИD	1.0 0.6	0. 0.
	2 3 4	DM DM	6.9 2.0	1.9 1.7	ND ND	1.1	1.3	ND ND	2.9	1.
	5	NO	2.2	1.8	NG	0.7	0.5	ND	0.8	ō.
MD(L)	1	МО	סא	NO	0.3	0.5	0.5	0.5	1.0	0.
	2	1.6	7.2 4.2	1.4	2.6 1.0	2.8 1.0	2.6 1.5	0.6 0.4	0.7 0.5	0.
	2 3 4 5	2.3 6.3	7.8	3.3	1.8	1.9	1.9	0.4	0.7	0.
	5	4.5	5.2	0.9	1.2	1.5	0.9	1.0	1.3	Ō,

TABLE 8 (continued)

			A Extra		Bar	ley Lea	f	Bar	ley Grai	n
Location*	Year	20 mt/ha/y	100 mt/ha	NPK	20 mt/ha/y	100	NPK	20 mt/ha/y	100 mt/ha	NPK
						- mg/kg				-
MD(H)	1	ND†	NO	NO	0.3	0.4	0.5	6.0	0.7	0.5
	2	1.7	5.7 2.9	0.6 0.7	2.5 1.2	2.6	2.6	0.4	0.6	0.4
	4	5.0	5.0	3.0	2.0	1.2 2.2	1.3 2.0	ND 0.5	0.3 0.6	0.6
	5	3.8	3.8	1.1	1.0	1.1	1.2	1.3	1.1	0.7
MI	1	0.9	3.8	0.2	1.2	2.3	1.2	11.2	5.3	13.0
	2 3	2.7	5.7	0.5 3.3	1.3	1.5	1.4	1.2	1.1	0.5
	د 4	2.3 ND	3.4 ND	3.3 ND	ND 1.1	NO 1.0	ND 1.1	ND 0.3	NO 0.4	0.2
	5	3.8	4.4	0.5	0.9	1.0	0.7	1.3	0.9	0.5
MN	1	7.5	24.3	4.1	1.1	1,3	1.3	0.4	2.7	0.3
	2	6.5	12.3	2.3	0.5	0.9	0.5	1.4	3.2	1.2
	2 3 4	4.6 6.2	5.4 11.2	2.0 2.4	0.7 1.1	0.8 2.0	1.0	0.9 0.9	1.1 1.5	0.9
	5	9.7	7.8	2.8	0.8	0.7	0.4	1.1	0.8	0.8
NE	1	3.9	6.5	NO	2.8	5.0	ND	1.6	1.6	ND
	2	3.5 11.6	8.1 7.8	NO ON	1.1 1.2	1.1 5.0	ND ON	1.2 0.3	4.8	ND
	2 3 4	5.7	5.8	ND	1.2	1.1	ND	0.3	0.5 1.0	DM OM
	5	11.1	9.6	ND	10.1	6.3	ND	0.8	1.2	ПD
ОН	1	1.9	6.7	0.5	1.5	1.7	0.9	0.5	1.2	0.5
	2	3.5 3.2	8.7 4.3	1.0	1.0 2.0	2.3 1.7	1.4	0. <i>6</i> 0.5	0.7 0.3	0.5
	2 3 4	3.7	3.5	0.7	0.6	0.5	0.5	0.3	0.3	0.2
	5	3.7	3.5 1.8	0.9	0.9	1.0	0.6	0.4	0.5	0.7
OR	1	ND	NO	ND	ND	ND	ND	3.8	5.4	2.5
	2 4	ND 4.4	NO 2.8	ND 1.3	NO NO	NO ON	ON DN	2.0 2.5	2.0 4.7	2.0
	5	6.4	4.2	2.2	0.1	1.3	0.1	0.1	1.3	2.5 0.1
UT	1	3.4	12.7	0.7	ND	ND	ND	0.3	0.9	0.2
	2	3.0	5.6	0.6	2.0	2.3	1.9	0.2	0.3	0.1
	3 4	4.5 4.3	6.4 4.5	0.7	3.8 2.3	3.6 1.6	4.6	0.1 0.3	0.1 0.4	0.1
	5	3.9	5.3	0.5	1.8	2.3	1.6	0.2	0.4	0.1
WI	1	8.6	14.6	5.5	3.1	3.5	3.4	2.7	4.2	1.9
	2	8.4	14.9	5.0	2.2	2.8	2.0	1.8	2.8	1.4
	3	7.9	14.8	5.4	0.8	1.0	0.7	0.5	1.3	0.4

<sup>\*</sup>See Table 4 for description of soils.

TND signifies not determined.

### SECTION 3

### EFFECTS OF SLUDGE PROPERTIES ON

## ACCUMULATION OF TRACE ELEMENTS BY CROPS

### INTRODUCTION

Loading limits for trace elements from municipal sewage sludges applied to land should be based on sludge and soil characteristics that affect plant availability of those elements. Current evidence indicates that the rate at which a plant root absorbs a trace element such as Cd, Zn, or Cu depends on the activity of the free-ion form of that metal in solution at the root surface. The activity at the root surface, in turn, depends on equilibrium reactions between solution and solid phases and the rate of transport to the root. Therefore, if we can predict the trace element uptake that a specific application of a given sludge on a particular soil will produce, we should be able to establish long-term loading limits, that will assure that additions of trace elements to the food chain are within tolerable limits and phytotoxicities of other trace elements are not a problem.

Sludges by themselves support certain trace element activities when equilibrated with the soil solution. Absorbing sites on the soil immobilize some of the dissolved trace element ions, causing more ions to be released from the sludge. If the trace element adsorption capacity of the applied sludge is small compared to the adsorption capacity of the soil, the soil properties will be very important in determining the equilibrium solution activity. However, if the trace element adsorption capacity of the sludge is high compared to that of the soil (usually associated with high sludge rates), the soil adsorption sites that can be filled at the activity supported by the sludge will result in only a small decrease in solution activity, and the sludge properties will dominate. Under these conditions the soil's effect on the pH of the mixture may still be significant, and the pH will affect trace metal concentrations in equilibrium with the sludge.

The fact that trace element activity, and thus plant uptake of trace elements from sludged soils, tends to approach a maximum as the sludge rate increases suggests that this behavior could be used to differentiate sludges that would support potentially harmful concentrations in plant tissue from sludges that would not, regardless of the application rate. Therefore, trace element activity could be used to differentiate sludges

that do an do not require regulation in terms of trace element loading limits. (Fig. 1). In this figure, a constant pH is assumed, and the dashed horizontal line (C\_) represents the maximum concentration of trace element that would be allowed in a test species based on the maximum allowable dietary intake calculations or on phytotoxicity. In the case of Cd, this line would represent the maximum allowable concentration in a test plant (such as lettuce), based on the assumption that uptake by different plant species grown on soils amended with sludge will show proportionate differences in Cd uptake, and the total intake from a "market basket" mix of species grown on a given sludge treatment can be estimated from the concentration determined in the test species. The letters A, B, C, and D represent Cd uptake curves derived from sludges that support different maximum Cd concentrations in plants, and the subscripts 1 and 2 represent uptake curves for the designated sludge applied to soils with low and high Cd-adsorption properties, respectively (if the maximum plant-Cd concentration that a given sludge loading would depend on soil properties (curves  $A_1$  and  $A_2$ )). Sludges B, C, and D, at or above the pH usec in the test. would not require loading limits regardless of the soil properties. In fact, sludge D represents a sludge that supports a lower plant Cd concentration than does the nonamended soil. This is a rare occurrence, but it has been observed.

Estimates of plant availability of the trace elements in sludges could be obtained from field or greenhouse studies with a specific variety of lettuce (or other crop that tends to accumulate the elements of interest) grown on sludge-amended soils that had been allowed to equilibrate under aerobic conditions. Later, reliable methods for assessing trace element supplying properties of sludges and adsorption characteristics of soils may be developed for use in place of the bioassay. Interpretation of such tests will require research relating test results to plant uptake. Use of chelating resins for desorbing trace elements from sludges and for establishing known trace element activities for soil adsorption curves appears promising.

FORMS AND AMOUNTS OF TRACE ELEMENTS IN MUNICIPAL SEWAGE SLUDGES

All domestic sewage sludges contain varying amounts of Cd, Cr, Cu, Pb, Ni, and Zn. Data presented by Sommers (1977) showed wide variations in metal concentrations and a fairly large number of sludges containing very high concentrations. With the implementation of the federal industrial waste pretreatment program as a control on the discharge of these trace metals into publicly-owned treatment facilities, the metal loads to municipal wastewater treatment facilities and subsequently the levels in municipal sludges can be expected to decrease with time. The federal program, which is likely to generate vast amounts of performance data in the future, is not mature enough to produce such data at this time. This can be illustrated by examining trends in sludge metal composition data from Chicago, Baltimore, and Philadelphia where local pretreatment programs have been implemented.

### Trends in sludge trace-metal concentrations

The Metropolitan Sanitary District of Greater Chicago (MSDGC) adopted a program in 1969 that was designed with objectives similar to those of the federal program which prevents pass-through of pollutants and produces higher quality effluents. The program specifies concentration limits for 13 contaminants and nine limiting conditions for the discharge of industrial wastes to the MSDGC sewerage system. The specific limits required by this program are as follows (in mg/L):

boron, 1.0; cadmium, 2.0; chromium (total), 25.0; chromium (hexavalent), 10.0; copper, 3.0; cyanide (total), 10.0; cyanide (readily releasable at 150°F and pH 4.5), 2.0; iron, 50; lead, 0.5; mercury, 0.005; nickel, 10; zinc, 15.0; fats, oils and greases, 100 (changed to 250 in 1983); and pH 4.5 to 10 units.

The levels of sewage-borne metals entering the MSDGC sewage treatment facilities (Tables 9 and 10) have decreased substantially since 1971 (Whitebloom et al., 1978; Lue-Hing, 1985, personal communication). As an example, for one POTW in the MSDGC system the influent Cd loading was reduced by 57.5% over the period of 1971 through 1977 (Table 9) and for another plant in the system by 69.4% over the period of 1971 through 1984 (Table 10). Similar results were obtained with pretreatment in Baltimore (Table 11) and Philadelphia (Table 12). Once the federal pretreatment program has been fully implemented, reductions on the national level can be expected to approximate those achieved by these programs.

### Forms of Metals in Raw Sewage

The solubility of a metal in the soil-sludge mixture is inherently governed by the particular chemical form in which it occurs in the sludge. To understand the chemical form of the various metals in municipal sewage sludge, one must first determine what forms of these metals are affected by the wastewater treatment process. Each metal will be distributed between the soluble and solid phase based upon a complex equilibrium controlled by the wastewater composition. However, in most cases, Cd, Cr, Cu, Pb, and Zn have been found to be predominantly associated with the solid phase in wastewater influents.

Elenbogen et al. (1984) compared the raw sewage metals concentration entering a pilot-scale primary settling tank with the metals concentration of activated sludge solids (mixed liquor) in an activated sludge pilot plant. Elenbogen et al. (1984) concluded that the metals in raw sewage entering primary settling tanks are bound to the wastewater solids in similar proportions to the metals bound to the mixed liquor solids of the activated sludge process, and that the distribution of metals between the solid and liquid phases is the same.

Metal adsorption by sludges has been demonstrated by spiking the sludge with inorganic metal salts. Elenbogen et al. (1984) studied the adsorptive capacity of the activated sludge process for metals and the

strength of the bond between metals and the activated sludge solids. In batch-scale experiments in Table 13, Elenbogen et al. (1984) "spiked" mixed liquor with known amounts of  $\mathrm{CdCl}_2$  up to 30 mg/L. Most of the soluble Cd was adsorbed to the mixed liquor solids in less than 15 minutes and over 90% was adsorbed after 1 hour of aeration. In a separate series of controlled pH batch experiments, Elenbogen et al. (1984) also studied the uptake of Cd by mixed liquor spiked with soluble Cd ( $\mathrm{CdCl}_2$ ) concentrations of 2.3 and 5.0 mg/L, and found that uptake was not influenced by mixed liquor pH in the range of 5.0 to 8.0. Over 90% of the soluble Cd uptake occurred in 15 minutes for all of the pH levels tested.

Neufeld and Hermann (1975) also reported high adsorptive capacities in batch-activated sludge reactors dosed with 30, 100, and 300 mg/L soluble Cd. In their experiments, 65 to 70% of the added soluble Cd was adsorbed on the activated sludge floc within 1 hour after dosing, even at the 300 mg/L soluble Cd dose. Within 4 hours, 80% of the initial soluble Cd had been adsorbed on the solids.

Patterson (1979) reported that in Rockford, Illinois, the soluble fractions of Cd, Cr, and Zn were 24.0, 26.4, and 16.1%, respectively, of these total metals in the raw sewage. Patterson and Kodukula (1984) reported that in the raw sewage at a sewage treatment facility of the MSDGC, Cd was 12.9% soluble, Cr was 1.7% soluble, Cu was 5.0% soluble, Pb was 16.9% soluble, Ni was 28.3% soluble, and Zn was 12.1% soluble. Similarly, Lester et al. (1979) reported that 72% of the Cd, 70% of the Cu, and 73% of the Pb was associated with the primary settled solids at the Oxford, England treatment plant. In a companion study, Stoveland et al. (1979) reported that 73% of the Cr and 74% of the Zn was associated with the primary solids at the Oxford plant.

In general, therefore, studies of raw sewage metal speciation indicate that most metals are associated with the solid phase. This, coupled with the demonstrated affinity for soluble metals shown by the activated sludge process, would indicate that most of the metals contained in municipal sludge are associated with the solid phase rather than the liquid phase.

## Forms of Metals in Sludges

Because the chemical composition of municipal raw sewage and the types of metal compounds that may enter a wastewater treatment plant vary widely the chemical transformations that will occur in the plant are difficult to predict. However, a general understanding of aqueous metal chemistry would suggest that metals would be present in both organic and inorganic forms. Metals associated with organic matter are probably bound strongly to complexing sites. Inorganic forms could include metallic particles, relatively pure precipitates (phosphates. carbonates, sulfides, or silicates), solid solutions resulting from coprecipitation with precipitates of Fe, Al, or Ca, or as metal ions strongly adsorbed on surfaces of Fe, Al, or Ca minerals (Corey, 1981). If metals such as Cd or Zn enter the treatment plant in aqueous form, coprecipitation with phosphates, hydrous oxides, or

sulfides of Fe and Al, and with phosphates and carbonates of Ca would be expected (CAST, 1980; Logan and Chaney, 1983).

The scientific literature contains little specific analytical data on the various species or forms of metals contained in municipal sewage sludge. Investigators have focused on a determination of sludge metal forms through the use of various extractants. The amount of metal found in these various extractants is an indicator of the form of the metal in the sludge.

Stover et al. (1976) developed a sequential extraction scheme for fractionating Cd, Cu, Pb, Ni, and Zn in anaerobically digested sludge. In their scheme they suggest that KNO $_3$  extracts exchangeable metals, KF extracts adsorbed metals, Na $_4$ P $_2$ O $_7$  extracts organically bound metals, EDTA extracts metal carbonates, and HNO $_3$  extracts metal sulfides. For the 12 sludges they studied, Stover et al. (1976) found that Zn was predominantly found in the organically bound (Na $_4$ P $_2$ O $_7$ ) form, Cu in the sulfide (HNO $_3$ ) form, and Pb in the metal carbonate (EDTA) form. Nickel was distributed in many forms, and Cd was predominantly in the metal carbonate (EDTA) form.

A similar extraction procedure, incorporating 0.5M  $\rm KNO_3$ , "ion-exchange water", 0.5M  $\rm NaOH$ , 0.05M  $\rm Na_2EDTA$ , and 4.0M  $\rm HNO_3$  has been employed to fractionate Cd, Cu, Ni, and  $\rm Zn$  in anaerobically digested air-dried sludge into forms designated as exchangeable, adsorbed, organically bound, carbonate, and sulfide/residual, respectively (Emmerich et al., 1982). While the Cd, Ni, and  $\rm Zn$  occurred in sludge predominantly in carbonate form, the major forms of Cu extracted were in the order: organically bound > carbonate > sulfide/residual.

Six types of sludge from the MSDGC's West-Southwest Sewage Treatment Works were subjected to a sequential chemical extraction procedure in an effort to characterize the metal forms present in the sludges (Elenbogen, et al., 1983). The following conclusions were reached:

- 1. Lagoon sludge, waste-activated sludge, and filter cake had similar chemical distributions of Cd, Cu, and Zn with the predominant species (48 to 69%) of these metals being in the water soluble and readily exchangeable (KNO $_3$  extractable) forms. These values seem very high compared with those of other studies.
- 2. Digested sludge had the highest percentage (22.5 to 25.4%) of Cd and Zn in the sulfide form (1M HNO<sub>3</sub>) compared to the other sludges.
- 3. Heat-dried sludge and Nu Earth (air-dried MSDGC sludge) had similar chemical distributions for Cd and Zn, with a relatively small percentage (less than 10%) of these metals found in the water soluble form when compared to the other sludges (21 to 56%), and the greatest amount (34 to 75%) of Cd and Zn being recovered in the organically bound form (Na $_4$ P $_2$ O $_7$  extractable) for Nu Earth. However, in the case of heat-dried sludge, the greatest amount

(29%) of Cu was found in the highly insoluble (concentrated HNO $_3$  extractable) form, with considerably less (17%) Cu being found in the organically bound form (Na $_4$ P $_2$ O $_7$  extractable) compared to the Nu Earth.

# Metal Speciation in Soils

Metals in soils may be present in many forms. The application of sewage sludge to soils may alter the speciation of a metal, which, in turn, may affect its availability to plants.

The use of chemical extractants in studying metal speciation in soils has been focused mainly on the so-called plant available forms. Metals have frequently been extracted with simple aqueous solutions to determine plant available forms (Adams, 1965; Gupta and MacKay, 1966). In all cases, metal concentrations in water extracts were low.

Sequential chemical extraction schemes, considered to be of greater value than single extractants in determining metal distribution in wastewater sludge (Stover et al., 1976), have frequently been applied to fractionate trace metals in sludge-amended soils. A modified version of a sequential extraction procedure developed by Stover et al. (1976) was used by Emmerich et al. (1982) to determine the chemical forms of metals in loamy soils amended with anaerobically digested sludge. Emmerich et al. (1982) observed that less than 3% of the total Cd, Cu, Ni, and Zn in a sludge-amended loam soil were extracted by 0.5M KNO3 and "ion-exchange water". Sposito et al. (1982) extracted 1.1 to 3.7% of these same metals using the same extractants. These results are consistent with exchangeable-plus-adsorbed forms of Cd, Pb, and Zn in sludges (Stover et al., 1976), and with water-soluble plus-exchangeable forms of Cd, Cu, Pb, and Zn in silt loam soils amended with digested sludge (Silviera and Sommers. 1977).

The diversity of reagents used to extract specific metal forms in soils make comparison of such studies difficult. Even if the reagent used is the same, the rate of leaching will be a function of the sample size, duration of extraction, temperature, and other factors (Sterritt and Lester, 1984).

Speciation of the metals in soils which receive sludge application is also important, as it will determine availability (Sterritt and Lester, 1984). However, Lake et al. (1984) conclude that no comprehensive or reliable speciation schemes for determining discrete heavy metal species or groups in sewage sludge and soil-sludge mixtures has yet been developed.

# Plant-Availability of Sludge-Borne Trace Elements

The rate at which an element is taken up by a plant root appears to be a function of the activity of the free ion at the root surface (Checkai et

al., 1982; Baker et al., 1984). However, as the concentration at the root surface decreases because of uptake, transport to the root may limit the rate of uptake.

The interacting factors that determine the rate of element uptake are most easily presented in a mathematical model. As most trace elements, particularly at low loadings, are delivered to the root surface primarily by diffusion (Barber, 1984), a diffusion model is used for illustrative purposes.

Factors Controlling Trace Element Uptake--Theoretical Considerations

Soil factors that affect diffusive transport of a solute include water content, solute concentration in solution, and the ability to resupply absorbed solute (buffer power). Important plant factors include root geometry (root radius, presence of root hairs/mycorrhizae) and root uptake physiology, i.e., root absorbing power and effects of root exudates. How these factors interact is shown in Eq. (1), which is a modification of an uptake equation derived by Baldwin et al. (1973) that describes the diffusive radial flux of solute from an isotropic medium (soil) to a cylindrical sink (plant root), assuming depletion of a cylindrical volume of soil surrounding each segment of root.

$$U = [C_{1i}b]1 - exp \left\{ \frac{-2\pi\alpha A_1 r_0 L_v t}{b(1 + \frac{\alpha A_1 r_0}{D_1 ef} \ln \frac{r_h}{1.65 r_0})} \right\}$$
 (1)

Soil factors:

 $C_{1i}$  = initial concentration (mol/cm<sup>3</sup>) of nutrient in soil solution

b = buffer power--the change in concentration of total labile form [adsorbed + dissolved] (mol/cm³ soil) per unit of change in concentration of dissolved form (mol/cm³ soil solution)

 $A_1$  = fractional area of soil solution

 $\theta$  = volumetric water content (cm<sup>3</sup> water/cm<sup>3</sup> soil)

 $D_1$  = diffusion coefficient in soil solution (cm<sup>2</sup>/sec)

### Plant factors:

The conductivity factor, f, decreases with a decrease in 0 because of greater tortuosity of the diffusion path at lower water contents. The buffer power, b, is equal to the change in concentration of total labile solute per unit change in concentration of that solute dissolved in the soil solution. The labile form includes dissolved and reactive adsorbed forms. Soils with high adsorption capacities for specific solutes generally show high buffer powers for those solutes. Commonly found ranges in buffer power for specific nutrients range from less than 1 for nonadsorbed species to more than 1000 for strongly adsorbed species, and generally decrease with increasing saturation of the adsorbing sites with a particular solute (Nye and Tinker, 1977; Barber, 1984).

The root density,  $L_V$ , is equal to the length of root per unit volume of soil. The value of  $L_V$  is readily determined for roots without root hairs or mycorrhizae, but their presence makes the geometry of the nutrient absorbing system more difficult to describe quantitatively. The root absorbing power,  $\alpha$ , is equal to the uptake flux density divided by the nutrient concentration at the root surface. In some cases,  $\alpha$  can be described by a Michaelis-Menten plot of flux density (mole per cm² per sec) vs. concentration at the root surface (mol/cm³). This relationship has been shown to depend on the pre-existing nutrient status of the plant (Lauchli, 1984), speciation of dissolved solute (Checkai et al., 1982), and antagonistic effects of other metals (Logan and Chaney, 1983). A confounding factor is the effect of root exudates on rhizosphere pH (Marschner et al., 1982) and possible complexing of trace metals.

Sludge applications affect both  $C_{1j}$  and b in a soil. As the sludge application rate increases to the point where soil adsorption sites that can be filled at the activity supported by the pure sludge are nearly saturated, further increase in sludge application results in little additional change in either  $C_{1j}$  or b, and U should approach a maximum. The maximum uptake rate obtained with a given sludge should differ from soil-to-soil because pH differences affect both  $C_{1j}$  and b, and  $\theta$  and f vary.

Similarly, uptake will vary with plant species because of differences in  $\alpha$ ,  $L_{\nu}$ ,  $r_{o}$ , and the nature of root exudates.

Most trace elements, particularly trace metals, added to soils appear to be immobilized mainly by adsorption reactions, which can usually be described by a Langmuir or Freundlich adsorption isotherm (Cavallaro and McBride, 1978; Garcia-Miragaya and Page, 1978; McBride et al., 1981; Kotuba, 1985). If the solute adsorption curve (adsorbed concentration vs. concentration in solution) for the soil and the desorption curve for the sludge have been determined, the equilibrium solute activity for any mixture of sludge and soil can be calculated. For example, if both curves can be described mathematically, in this case, by a Langmuir equation, the variables  $C_{1i}$  and b in the uptake equation can be calculated in the following way for a trace metal:

$$M = \frac{M_T C}{K + C} \text{ or } C = \frac{M K}{M_T - M}$$
 (2)

where M is metal adsorbed, MT is the metal adsorption capacity, K is a constant equal to the dissolved metal concentration at one-half saturation of the adsorption sites, and C is the equilibrium metal activity. If a given amount of sludge is added to a given amount of soil, an amount of adsorbed metal, X, will be transferred from the sludge to the soil (if the sludge supports a higher metal activity than the soil), and a new equilibrium metal activity, CM will result. If the amount of metal in solution is insignificant compared with that adsorbed, the reaction can be represented by the following equation, where the subscripts A and B represent soil and sludge, respectively.

$$C_{M} = \frac{(M_{A} + X)K_{A}}{M_{TA} - (M_{A} + X)} = \frac{(M_{B} - X)K_{B}}{M_{TB} - (M_{B} - X)}$$
(3)

At equilibrium, equation (3) can be used to solve for X and C<sub>M</sub>. If X is small compared with M<sub>B</sub>, the equilibrium metal activity will be very close to that for the sludge alone. In theory this approach would permit estimation of both metal solution activity and buffer power for input into an uptake model. The fact that sludge properties change with time presents some difficulty in practice.

## Experimental Results

Few, if any, investigators have evaluated in a single study the interactions among plant uptake and sludge properties, sludge rate, and metal

activity in the soil solution. Fujii (1983) found that application of a sludge containing 180 mg Cd/kg on sand and silt loam soils at pH  $6\pm0.3$  maintained Cd activities in the sand 2.5 to 4 times those in the silt loam at rates up to 18.4 kg Cd/ha. Concentrations of Cd in corn tissue grown on these soils in the greenhouse were 1.5 to 2 times higher in the sand than the silt loam (Shammas, 1978). Adsorption characteristics of the sludge and soil were not studied.

Chelating resins have been used for determining Cd-adsorption characteristics of a muck and a sandy soil (Turner et al., 1984), and for characterizing dissolved metal complexes (Henderson et al., 1982; Hendrickson and Corey, 1983). Because of the large metal-ion buffering power of these resins, metal ions can be adsorbed on or desorbed from soils or sludges without significantly changing the metal-ion activities supported by the resin, if the proper ratio of resin to soil or sludge is used. Adaptation of this chelating-resin methodology to the routine determination of metal adsorption/desorption characteristics of soils and sludges appears promising.

Factors affecting the lability of metals in sludges have not been determined directly, but rather inferred from theoretical considerations, fractionation studies, or from greenhouse or field experiments with sludges of different chemical compositions. Greenhouse and field studies have generally supported the hypothesis (Corey, 1981; Corey et al., 1981) that much of the immobilization of trace metals in sludges is caused by coprecipitation with Fe, Al, and Ca precipitates during the treatment process, but no way of quantifying this effect has yet been devised.

In comparing the Cd uptake from equal rates of two noncalcareous sludges in the greenhouse, Cunningham et al. (1975a) found that the average concentration of Cd in plant tissue was about the same for the 2 sludges (1.5 vs. 1.4 mg/kg), even though the Cd content of the sludges differed by a factor of 3 (76 vs. 220 mg/kg). The sludge with the lower Cd content had a lower Fe content (1.2 vs. 7.9%) and also a lower P content (2.9 vs. 6.1%). Thus, the effect of the higher Cd content in the one sludge may have been offset by a relatively high content of substances such as FePO, with which Cd could coprecipitate. In a field study, the Cd concentration in corn leaves from plots treated with a sludge containing 229 mg Cd/kg, 3.0% Fe, 1.1% Al, 4.7% Ca, and 1.6% P was nearly 3 times as high (1.7 vs. 0.6 mg/kg) as in corn leaves from plots treated with the same amount of Cd supplied by a sludge containing 180 mg Cd/kg, 7.8% Fe, 2.5% Al, 1.5% Ca, and 3.0% P (Keeney et al., 1980). The isotopically exchangeable Cd was also found to be 3 times higher for the sludge low in Fe and P, even though the total Cd concentrations in the 2 sludges were similar.

In a greenhouse study, Bates, Soon and Haq (1979, personal communication) added sludges to soils cropped to annual ryegrass over a period of about 5 years. Fourteen successive crops of ryegrass were grown, with sludge being added prior to seeding each crop. The cumulative Cd loadings were 10.6 kg/ha for the Sarnia sludge and 12.1 kg/ha for the Guelph sludge.

The sludges had similar ratios of P to Cd at the start of the fourteenth crop, but the ratios of Fe to Cd were 889 and 195 for the Sarnia and Guelph sludges, respectively. The average Cd concentrations in the fourteenth crop of ryegrass were 1.35 mg/kg for the Sarnia sludge and 2.35 mg/kg for the Guelph sludge. With nearly equal additions of total Cd, the lower Cd availability was associated with the sludge having the higher Fe content. In fact, there was a measurable, though not statistically significant, decrease in plant-Cd concentration compared to the control treatment with a sludge in which the Fe and P contents were 8 and 5%, respectively, even though the total sludge-applied Cd was 1.63 kg/ha (Bates, 1986, Personal commununication).

Beli, et al. (1985, Personal communication) added 2 sludges with equal Cd but different Fe concentrations (sludges A and B in Table 14) to a fine sandy loam at rates high enough to show maximum Cd concentration in tobacco. The sludge with higher Fe showed lower Cd uptake even though the pH was slightly lower with that sludge.

Additional evidence that metal concentration in plants may be affected by the form of metals in sludge can be implied from the work of King and Dunlop (1982). Sludges from Wilmington, North Carolina (13 mg Cd/kg) and Phi-ladelphia, Pennsylvania (225 mg Cd/kg) were applied to several soils in which corn was grown in the greenhouse. Sludges were applied at rates to supply equal amounts of Cd. The effect of sludge type was significant, as evidenced by different slopes in models of Cd concentration in corn stover regressed on Cd loading rate:

Wilmington: plant Cd (mg/kg) = 0.11 + 0.18 Cd rate (kg/ha)Philadelphia: plant Cd (mg/kg) = 0.18 + 0.45 Cd rate (kg/ha)

The relationship between sludge rate and metal uptake by plants has been investigated by many researchers. In general, for Cd and Zn the metal concentration in tissue approaches a maximum and has shown a logarithmic or Langmuir-type relationship with sludge rate. In many cases, Cu uptake is not affected significantly. In studies when the sludge addition was not high enough to approach a constant metal activity in solution, the relationship between sludge rate and tissue concentration often approached linearity. For example, Pietz et al. (1983) and Hinesly et al. (1984a) found a near linear relationship between Cd or Zn in corn leaves and Cd application rate up to 111 kg/ha applied to a calcareous strip mine spoil in a sludge containing about 300 mg Cd/kg. The sludge was applied over a period of 6 years. When the same sludge was applied for 12 years to an acid Blount soil, the year-to-year variability in leaf analyses obscured any relationship to cumulative additions (Hinesly et al., 1984a). Vlamis et al. (1985) also noted a linear response with Cd and Zn in barley with 2 sludges applied at rates up to 225 mt/ha on an acid soil. However, metal uptake at the high sludge applications may have been augmented by the effects of lower pH found in these treatments.

In contrast to the linear responses reported above, Soon et al. (1980) found that Cd in corn stover was logarithmically related to sludge rate

(up to 4 kg Cd/ha) for 3 sludges receiving either Fe, Al, or Ca additions during treatment. All relationships were logarithmic, but the plots of the Cd concentration in stover and the log of Cd applied differed in slope by a factor of 3, emphasizing the effects of sludge properties on Cd availability.

In a field study, Chaney et al. (1982) used sludges with different Cd concentrations to determine the effect of concentration and application rate on Cd concentration in lettuce. The first increment of low-Cd sludge (13 mg/kg) increased Cd concentration in the lettuce slightly, but higher rates had no further effect on Cd concentration. The first increment of high-Cd sludge (210 mg/kg) had a pronounced effect on Cd concentration, but the response to additional increments was less pronounced, indicating a logarithmic response (Figure 2). Unpublished data on Cd accumulation in lettuce (Chaney, 1985, Personal communication) (Figure 3) also show a plateau effect along with an effect of pH and sludge composition.

In another field study (Bell et al., 1985, Personal communication), tobacco was grown on sludge-treated soil. Copper, Cd, and Zn contents of the tobacco were affected markedly by the first increment of sludge. Additional increments had no effect on Cu content, and only slightly increased Zn and Cd content.

Sommers (1985, Personal communication) presented data in which 3 sludges containing high concentrations of Cd (284, 1210, and 247 mg Cd/kg) were applied to a Chalmers soil at linearly increasing rates for 2 sludges, and logarithmically increasing rates for the third (Table 15). Over a period of 8 years, oats, winter wheat, soybeans, and corn were grown on these plots. In almost all cases, the relationship between tissue concentration of Cd or Zn and rate of metal applied in the form of sludge was logarithmic or approached a constant value at high sludge rates (Table 15). This was also the case with vegetable data from the Metropolitan Sanitary District of Greater Chicago (Table 16) and of Hinesly et al. (1984b). Crops studied by Hinesly et al. (1984b) included corn (Figure 4) and many other grass species (Table 17) grown for 3 years following sludge application. One interesting aspect of this latter study was the decrease in Cd concentration in corn tissue with time after application, particularly at the high rate (Figure 4). This effect is further illustrated in data from Dowdy et al. (1984) which show uptake by corn silage in grams per hectare over a period from 1979-1984 (Figure 5). These studies agree with the conclusions of CAST (1980) and Logan and Chaney (1983), that bioavailability of metals remains constant or decreases over a period of years at a constant pH. The marked decrease in metal uptake at the high rate after 1 year suggests that exposure of anaerobically-digested sludges to an aerobic environment, and/or interactions with soil may produce marked changes in lability of the metals during the first year following application.

In addition to metal forms and concentrations in sludges, the effects of sludge additions on soil pH must also be considered. Lime-stabilized sludges function as liming materials (Soon et al., 1980; Chang et al.,

1980). Noncalcareous sludges may either raise or lower soil pH, depending on such factors as amount of  $NH_4$ -N nitrified or sludge pH, and pH buffering capacity in relation to the pH and buffering capacity of the soil. Generally, plant uptake of metals increases with increasing acidity (CAST, 1980; Logan and Chaney, 1983).

### CONCLUSIONS

- 1. Trace metals in the influent to a sewage treatment plant are associated mainly with the solids, and they remain associated with solids in the sludge following treatment.
- 2. Concentrations of trace elements in many POTW sludges have decreased markedly in the past decade as a result of industrial waste pretreatment, and this trend is expected to continue.
- 3. Uptake of solutes by plants is influenced by soil and plant factors, and a simplified mathematical model is presented to indicate how these variables interact in affecting solute uptake.
- 4. The relevant soil variables related to solute uptake affected most by sludge application are concentration in solution, concentration of the labile adsorbed form, and distribution of dissolved species between free-ion and complexed forms.
- 5. The relevant plant variables related to solute uptake affected most by sludge application are root adsorbing power (related to speciation in solution and concentrations of other ions competing for uptake sites) and possibly root geometry.
- 6. The equilibrium trace element concentrations that a sludge supports depend on the chemical properties of the sludge, particularly the presence of trace-element precipitates, whether relatively pure or coprecipitated with Fe, Al, or Ca precipitates, the strength of bonding to organic and mineral adsorption sites, the proportion of potential adsorbing sites filled, and the presence of dissolved ligands capable of complexing the trace metals. If sludge matrix is constant, plant availability of a trace element increases with increasing concentration of that trace element in the sludge.
- 7. If a sludge supports a higher equilibrium solution concentration of an element than does a soil, mixing the 2 will result in an equilibrium concentration intermediate between the 2 that should be predictable if the desorption characteristics of the sludge and adsorption characteristics of the soil are known. This hypothesis has not been tested experimentally.
- 8. As increasing amounts of sludge are added to a soil, trace-element adsorption sites on the soil become progressively saturated (or

desaturated) to the point that the equilibrium concentration approaches that of the sludge alone. Further sludge applications above a particular level depending on soil properties should result in little, if any, change in equilibrium concentration. Below this critical sludge rate, soil adsorption characteristics affect the equilibrium concentration supported by a given addition of sludge. Above that critical sludge rate, the equilibrium concentration characteristic of the sludge should be maintained (sludge controls).

- 9. If the equilibrium trace-element concentration (and buffer power) supported by the sludge is less than that which will result in excessive concentrations in plant tissue or damage to the plant, there is no need to limit application rates of that sludge on the basis of metal content.
- 10. If the equilibrium metal concentration (and buffer power) supported by the sludge at a specified pH is high enough to cause excessive concentrations in plant tissue or plant damage, determining maximum loading rates based on both soil and sludge characteristics will be required.
- 11. Most research indicates that plant availability of sludge-derived metals stays the same or decreases with time following application. Therefore, any testing procedures developed to establish long-term metal-loading limits should be run after the sludge has been allowed to equilibrate with the soil. Presently the time(s) required to equilibrate sludges with soils are not precisely known, but limited data suggest a minimum of two cropping seasons.
- 12. Methods involving chelating resins for obtaining metal desorption curves for sludges and metal adsorption curves for soils appear promising.
- 13. Addition of Fe or Al salts or lime during the sewage treatment process appears to reduce equilibrium metal activities supported by sludge; however, research designed to test this hypothesis has not been done.

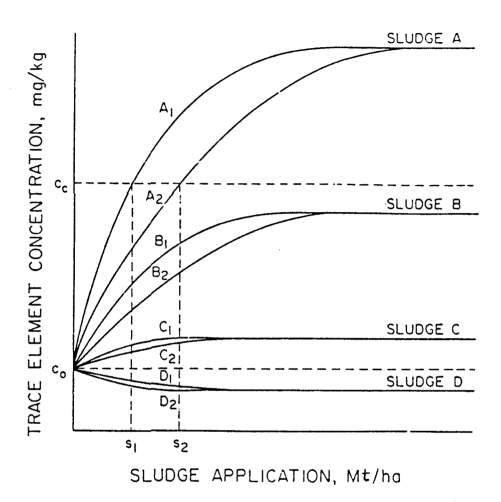
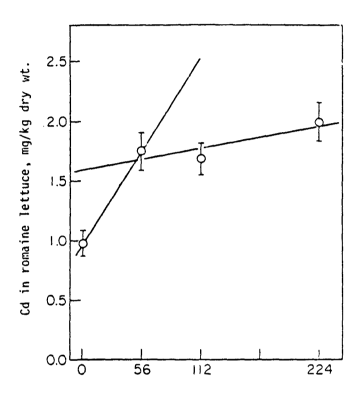


Figure 1. Basis for differentiating sludges that do not require loading limits to prevent harmful trace element accumulations in plants (sludges B, C, and D) from one that does (sludge A). The two curves for each sludge represent that sludge applied to a soil of relatively low adsorption capacity (subscript 1) and a soil of higher adsorption capacity (subscript 2). C<sub>3</sub>, is the concentration of a given element in plant grown on the unamended soil, is shown as being the same for both soils. C<sub>c</sub> is the critical concentration in the plant, and S and S are loading limits for sludge A applied to soils (1) and (2), respectively).



Sludge application rate (mt/ha)

Figure 2. Uptake of Cd by romaine lettuce from soils treated with municipal sewage sludge at various rates (mean data over 8 years).

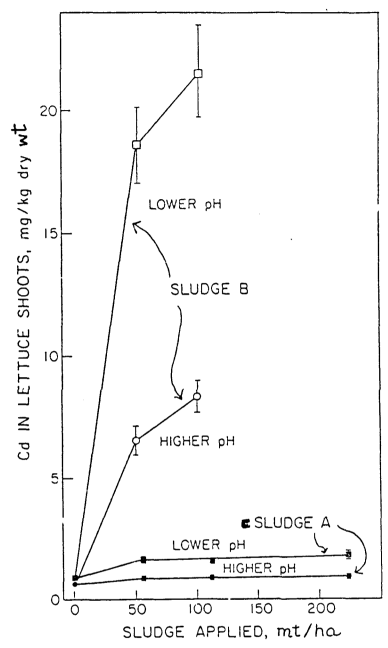
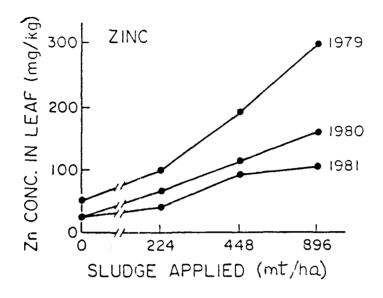


Figure 3. Effect of sludge application rate on Cd in lettuce leaves. Sludge A (13.4 mg Cd/kg) and 8.3% Fe) applied in 1976 and lettuce grown in 1976 to 1983; sludge B (210 mg Cd/kg) and 2.5% Fe) applied in 1978 and lettuce grown in 1978 to 1983. Results shown are geometric means over years ± standard error (Chaney, 1985, Fersonal communication).



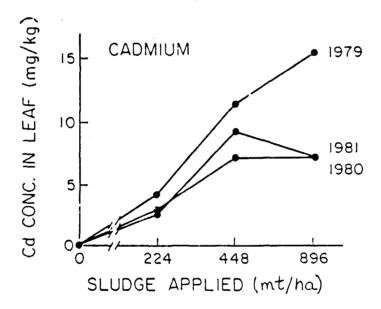


Figure 4. Effect of a one-time application of municipal sewage sludge containing 4230 mg Zn/kg and 300 mg Cd/kg on the Zn and Cd contents of corn leaf tissue in each of 3 years after application to a calcareous strip-mine spoil. Data for 1979 and 1980 are from Hinesly et al. (1984c). Data for 1981 are from Table 17.

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Figure 5. Decrease in Cd uptake by corn silage with time after application of sewage sludge at three rates in 1979 (Dowdy et al., 1984).

Table 9. METAL LOADINGS AND CUMULATIVE PERCENT REDUCTIONS TO CHICAGO AREA TREATMENT FACILITIES, 1971 THROUGH 1977.\*

	Cd	Cr	Cu	РЬ	Ni	Zn
			<b> k</b> gs/d	day		
1971	398	5,197	2,166	2,049	2,443	6,972
1972	343	3,321	1,996	1,793	1,377	4,641
1973	301	2,463	961	1,063	957	4,260
1974	213	1,894	652	735	643	3,403
1975	113	1,522	538	497	386	2,537
1976	132	1,527	<b>6</b> 85	368	416	2,400
1977	168	1,422	588	536	436	2,587
Cumulative % reduction	57.7	72.6	72.9	73.8	82.2	62.9

<sup>\*</sup>From Whitebloom et al. (1978).

Table 10. METAL LOADINGS AND CUMULATIVE PERCENT REDUCTIONS TO CHICAGO AREA TREATMENT FACILITIES, 1971 THROUGH 1984.\*

	Cd	Cr	Cu	РЬ	Ni	Zn
			kgs/	day		
1971	875	11,434	4,765	4,508	5,374	15,338
1984	267	2,607	2,088	871	1,545	5,109
Cumulative % reduction	69	77	56	81	71	67

<sup>\*</sup>Lue-Hing (1985, personal communication). Data in Tables 9 and 10 are from two different POTWs within the MSDGC system.

Table 11. RESPONSE OF METALS CONCENTRATIONS IN DIGESTED SLUDGE FILTER CAKE AT THE BACK RIVER POTW, BALTIMORE, MARYLAND IN RESPONSE TO PRETREATMENT EFFORTS.\*

Year	Cd	Cu	Pb	Ni	Zn
		m	g/kg dry weigl	nt	
1980	18	2840	433	381	3400
1981	19	2070	493	374	3410
1982	18	1110	398	193	2360
1983	23	1060	324	214	2620
1984	26	1010	372	266	2750
1985	22	681	346	126	2030

<sup>\*</sup>Source identification began in 1980, and source reduction began in 1981. Based on monthly composites in early years, then biweekly and weekly. Spencer, E. (1985, Personal communication).

TABLE 12. RESPONSE OF METALS CONCENTRATIONS IN SLUDGES AT TWO PHILADELPHIA POTWS IN RESPONSE TO PRETREATMENT PROGRAM.\*

Year	Cd	Cu	РЬ	Ni	Zn
			mg/kg dry wei	ght	
		Sout	hwest		
1974	31	825	1540	100	3043
1976 1977 1978 1980 1981 1982 1983 1984 1985	27 27 16 18 25 20 12.5 14.3 15.0	1110 1400 1020 986 971 940 736 1140 880	2710 2170 1800 740 562 1030 421 427 373	103 185 275 98 117 113 79 111	2650 3940 4050 2780 2300 2440 1700 1830 1730
		Nort	heast		
1974	108	1610	<b>2</b> 270	391	5391
1976 1977 1978 1980 1982 1983 1984 1985	97 71 57 26 14 10.9 12.4 17.3	2240 2320 1240 1210 985 1020 1200 1270	2570 2680 1620 728 423 351 360 382	372 459 319 275 185 130 130	5070 3920 5910 3890 2570 2110 1980 2100

<sup>\*</sup>Source identification began in 1976. Liquid sludge analyzed until 1982, and sludge filter cake in 1983 and later. Semske, F. (1985, Personal communication).

TABLE 13. CADMIUM UPTAKE OF MIXED LIQUOR SEWAGE SLUDGE (MLSS) AT VARYING SOLIDS TO CD RATIOS.\*

Run	Cd added†	Time of aeration	MLSS	MLSS to Cd ratio	Soluble Cd remaining	Cd uptake
	mg/L	min	mg/L		mg/L	%
A	1	0 30 60	1900 1900 1900	1900:1 1900:1 1900:1	0.0153 0.0189 0.0138	98.5 98.1 98.6
В	2	0 15 60 120 180	1000 1000 1000 1000 1000	500:1 500:1 500:1 500:1 500:1	0.375 0.186 0.122 0.112 0.066	81.2 90.8 93.9 94.6 96.7
С	10	0 15 30 60	9680 9680 9680 9680	968:1 968:1 968:1 968:1	0.49 0.31 0.15 0.14	95.1 96.9 98.5 98.6
D	30	0 60 120 960	2600 2600 2600 2600	87:1 87:1 87:1 87:1	7.50 2.28 1.82 0.78	75.0 92.4 94.0 97.4

<sup>\*</sup>From Elenbogen et al. (1984).

tAdded in the form of  $CdCl_2$ .

TABLE 14. EFFECT OF SLUDGE PROPERTIES ON PLATEAU CONCENTRATIONS OF CD IN TOBACCO LEAVES GROWN IN THE FIELD LONG AFTER SLUDGE APPLICATION.\*

Sludget	Concentration in sludge Cd Fe			Maximum application		Cd concentration above control		
	mg/kg	%	mt/ha	kg Cd/ha		mg Cd/kg dry wt.		
A	13.2	2.5	224	2.90	5.4	7.5		
В	13.4 13.4	8.3 8.3	224 224	3.00 3.00	5.2 5.8	2.4 0.2		

<sup>\*</sup>Bell, et al. (1985, Personal communication).

tSludge A applied in 1972 to Beltsville silt loam. Sludges B applied in 1970 to Christiana fine sandy loam. Tobacco ('Maryland 609') grown in 1983 and 1984 for Sludge A, and 1984 for Sludge B.

TABLE 15. EFFECT OF SLUDGE RATE AND YEAR AFTER SLUDGE APPLICATION ON CONCENTRATIONS OF CADMIUM, ZINC, COPPER, AND NICKEL IN OAT STRAW AND LEAVES OF WINTER WHEAT, SOYBEAN, AND CORN.\*

	Metal	Oats	Winte			tration bean		orn
Sludge		(1)	(2)	(उ)	(1)	(8)	(1)	(8)₹
	kg/ha			m	g/kg -			
				Cad	mium			
A	0 16 32 64 127	0.9 1.3 2.0 2.2 3.3	0.5 1.5 2.0 5.4 6.5	0.3 0.3 0.4 0.5 0.5	1.6 2.2 1.8 1.8 2.4	1.4 1.8 1.9 3.4 4.9	1.3 1.6 1.5 1.6	0.6 1.2 1.0 1.0
FK	0 68 136 203	0.9 9.9 15.9 20.3	0.5 12.1 14.6 15.8	0.3 1.2 1.4 1.5	1.6 4.6 5.0 6.0	1.4 5.0 8.9 11.0	1.3 5.1 7.8	0.6 5.6 4.7
MA	0 14 28 42	0.9 0.9 1.1 1.1	0.5 1.1 1.0 1.4	0.3 0.3 0.3	1.6 2.1 1.7 2.1	1.4 1.2 1.3 2.4	1.3 1.1 1.4 0.9	0.6 1.1 1.1 0.8
				<u>Z</u>	inc			
A	0 38 1 76 2 15 2 3	20.2 30.3 41.4 48.9	21.9 56.3 67.4 63.5	35.3 38.1 44.2 44.4	41.6 53.0 66.8 55.0	72.2 103.5 92.6 110.0	25.7 39.1 53.7 48.7	69.8 62.7 71.3 77.0
FK	0 106 213 319	20.2 27.0 30.3 32.0	21.9 52.8 50.5 48.8	35.3 40.1 36.9 40.1	41.6 54.4 59.1 63.2	72.2 79.1 80.1 82.9	25.7 50.5 48.5	69.8 56.2 54.1
MA	0 <i>2</i> 9 1 58 2 8 7 5	20.2 20.2 26.0 22.6	21.9 38.0 33.1 30.3	35.3 32.8 38.5 35.6	41.6 44.1 48.4 51.2	72.2 76.7 88.5 86.7	25.7 44.4 44.7 52.7	69.8 78.1 67.8 83.9

TABLE 15 (continued)

			Leaf metal concentration t							
	Metal	Oats	Winte	r wheat	Soyt	pean	Co	rn		
Sludge		(1)	(2)	(8)	(1)	(8)	(1)	(৪)		
	kg/ha			mg/	kg					
				Cop	per					
A	0 67 134 269	2.0 2.2 3.0 4.9	4.0 4.1 5.0 13.0	2.6 2.5 3.4 4.0	7.4 8.7 8.1 7.7	9.1 9.4 9.4 9.7	8.3 8.2 9.2 9.4	5.5 5.9 6.4 7.5		
	538	6.0	7.8	4.8	7.4	10.2	10.3	7.7		
FK	0 74 149 223	2.0 2.6 4.2 3.7	4.0 4.6 4.3 4.5	2.6 4.8 3.7 4.7	7.4 9.8 10.1 10.3	9.1 10.1 10.6 11.2	8.3 11.4 10.6	5.5 6.7 6.8		
A	0 25 50 76	2.0 2.5 2.5 2.6	4.0 3.7 4.9 4.0	2.6 3.2 3.1 2.7	7.4 8.2 8.4 10.0	9.1 10.0 10.9 10.2	8.3 7.9 8.9 9.6	5.5 7.1 6.7 4.7		
				Nic	kel					
A	0 114 228 451 914	1.1 2.0 3.3 5.1 12.5	2.9 3.2 4.0 9.8 7.5	0.3 0.3 0.4 0.4	49.3 18.5 20.2 28.5 20.8	5.7 9.5 9.3 10.9 16.6	3.1 3.9 1.6 5.9 2.5	0.2 0.3 0.2 0.3 0.3		
FK	0 24 48 72	1.1 1.6 2.5 3.4	2.9 4.1 2.8 3.6	0.3 0.4 0.6 0.5	49.3 44.1 28.4 19.2	5.7 6.9 8.6 9.2	3.1 2.3 2.6	0.2 0.3 0.2		
A	0 12 24 36	1.1 1.1 1.2 1.3	2.9 2.5 2.7 2.7	0.3 0.4 0.5 0.4	49.3 21.3 24.2 11.4	5.7 3.7 5.2 4.2	3.1 1.3 6.8 1.4	0.2 0.5 0.5 0.4		

<sup>\*</sup>Sommers (1985, Personal communication).

tleaf tissue for winter wheat, soybean and corn; straw for oats.

 $<sup>\</sup>pm \text{Numbers}$  in parentheses denote crop year following sludge application.

TABLE 16. CADMIUM, ZINC, COPPER, AND NICKEL CONCENTRATIONS IN EDIBLE PARTS OF VEGETABLES GROWN AT WEST-SOUTHWEST SEWAGE TREATMENT WORKS, METROPOLITAN SANITARY DISTRICT OF GREATER CHICAGO.\*

		Sludge added, mt/ha						
Crop	Year	0	60	120	240	300		
			mg me	tal/kg edit	ole tissue ·			
			<u>c</u>	admium				
Beets	1979	0.2	1.4	1.6	2.7	2.9		
	1983	0.7	1.8	2.0	1.8	4.9		
Tomatoes	1979	1.1	1.8	2.4	2.2	3.4		
	1983	2.0	2.2	2.1	2.6	2.9		
Swiss	1979	2.3	8.0	12.2	16.8	22.1		
chard	1983	1.3	4.0	8.1	8.4	12.2		
Carrots	1979	0.7	1.4	1.2	1.9	2.3		
	1983	1.3	1.8	2.8	3.4	3.1		
Green	1979	0.4	0.3	0.4	0.4	0.5		
beans	1983		0.1	0.2	0.1	0.2		
Spinach	1979	6.4	12.6	10.3	14.4	12.1		
	1983	8.5	21.4	28.3	31.7	33.8		
	(1982)	10.3	11.3	17.2	10.6	11.8		
			Zinc	-				
Beets	1979	34	45	62	93	90		
	1983	39	55	59	80	97		
Tomatoes	1979	27	30	30	34	35		
	1983	27	32	31	29	32		
Swiss	1979	69	129	176	237	302		
chard	1983	62	91	120	129	251		
Carrots	1979	20	22	25	27	32		
	1983	31	33	30	31	33		
Green	1979	32	33	37	37	37		
beans	1983	38	35	33	35	35		
Spinach	1979	147	249	265	309	311		
	1983	209	433	404	451	472		
	(1982)	201	276	315	226	258		
			54		(c	ontinued		

TABLE 16 (continued)

			<u></u>	udge added, mt	/ha	
Crop	Year	0	60	120	240	300
		~ ~ ~	mg	metal/kg edibl	e tissue	
			Copper			
Beets	1979	8.8	10.4	9.8	11.1	12.8
	1983	9.7	11.9	12.3	14.2	14.1
Tomatoes	1979	13.2	16.6	14.5	16.2	16.6
	1983	10.4	12.4	11.9	11.0	12.0
Swiss	1979	23.7	20.8	25.6	30.9	29.2
chard	1983	25.2	26.6	27.6	26.7	29.4
Carrots	1979	5.4	6.0	5.8	5.9	6.5
	1983	7.9	7.8	7.5	7.8	7.7
Green	1979	8.6	8.9	9.2	8.2	8.5
beans	1983	8.5	9.1	7.6	8.6	7.9
Spinach	1979	13.8	14.2	17.3	18.9	19.1
	1983	12.3	14.2	15.1	16.3	17.4
			<u>Nickel</u>			
Beets	1979	0.5	0.6	0.9	1.4	1.5
	1983	2.1	2.6	2.8	3.3	6.6
Tomatoes	1979	1.1	1.3	1.4	4.1	2.7
	1983	6.8	7.1	27.1	7.2	8.4
Swiss	1979	1.3	1.4	2.5	3.5	4.3
chard	1983	0.8		2.4	2.1	3.3
Carrots	1979 1983	0.6 1.9	1.1 0.7	1.4	2.2	2.9 1.0
Green	1979	3.3	1.2	2.4	3.6	3.3
beans	1983	2.3		2.9	5.1	4.7
Spinach	1979	1.4	1.5	1.7	2.7	2.6
	1983	10.6	5.8	8.8	6.0	10.0

<sup>\*</sup>Nu Earth was applied from 1977 through 1979 in three equal applications (C. Lue-Hing, 1985, Personal communication).

TABLE 17. EFFECT OF SLUDGE RATE APPLIED IN 1979 ON CONCENTRATIONS OF CADMIUM, ZINC, AND COPPER IN WHEAT, RYE AND FOUR GRASSES IN 1981.\*

-		Sludge rate	, mt/ha	
Crop	0	224	448	896
		– – mg meta	1/kg tissue -	
		<u>Cadmium</u>		
Wheat (leaf)	<0.1	0.7	1.6	1.5
Rye (leaf)	0.1	0.4	0.7	0.9
Redtop	0.1	0.3	0.8	1.2
Brome	0.1	0.8	1.0	1.6
Orchard grass	0.2	0.5	1.4	1.7
Western wheat grass	<0.1	0.4	0.8	0.7
Reed canary grass	0.1	0.2	0.6	0.8
Perennial rye	<0.1	0.1	0.9	0.7
Timothy	0.1	0.1	0.2	0.8
Tall fescue	0.1	0.4	0.9	1.8
		Zinc		
Wheat (leaf)	15	26	39	46
Rye (leaf)	20	42	71	61
Redtop	19	45	68	86
Brome	17	29	40	48
Orchard grass	21	32	39	49
Wetern wheat grass	19	35	41	44
Reed canary grass	28	62	104	121
Perennial rye	17	40	90	88
Timothy	23	50	65	74
Tall fescue	23	35	48	62
		Copper		
Wheat (leaf)	7	8	9	9
Rye (leaf)	10	13	18	18
Redtop	5	9	12	14
Brome	5	7	8	10
Orchard grass	4	6	7	9
Western wheat grass	4	7	S	8
Reed canary grass	6	9	11	13
Perennial rye	4	6	10	17
Timothy	5	9	10	12
Tall fescue	5	7	8	10

<sup>\*</sup>Hinesly and Redborg (1984b)

# SECTION 4

# EFFECTS OF LONG-TERM SLUDGE APPLICATION ON ACCUMULATION OF TRACE ELEMENTS BY CROPS

#### INTRODUCTION

Since the last comprehensive review of elemental uptake by plants grown on sludge treated soils (Logan and Chaney, 1983), considerable data from long-term field experiments have become available. Most experiments were designed to assess the effects of sludge applications on plant accumulation of metals (e.g., Cd, Cu, Ni, Zn, etc.). This report will concentrate on newly available long-term field data in terms of their implications on land application of sludges.

In the following sections, we will attempt to answer:

- (1) What is the quality of experimental data?
- (2) Do repeated annual sludge applications affect the metal accumulation in plant cissue?
- (3) Do the plant tissue metal accumulation patterns of a single sludge application differ from those of multiple sludge applications having equal total metal input? and
- (4) Does metal uptake by plants change following termination of sludge application?

# Nature of the Experimental Data

The data used for the analyses and the results presented in this report are, for the most part, derived from replicated field experiments. Logan and Chaney (1983) pointed out that common errors in the study of toxic element uptake by plants grown in sludge treated soils are (1) substituting inorganic metal salts for sludge or spiking the sludge with inorganic salts in preparing the growth medium; (2) relying on short term small pot experiments in the greenhouse rather than field observations to

predict metal concentrations in plants. Recently data from field studies have become available which allows us to greatly reduce our reliance upon information flawed by the above mentioned errors. During the course of our deliberation, only in the absence of field data did we draw upon "large" pot, greenhouse or growth chamber findings. It has not been necessary to use any data derived from studies that used "salts" as the metal source.

The drawing of "general" conclusions from a pool of information derived from unrelated field studies have limitations that must be recognized (plant species exhibit different abilities to accumulate metals). Only plant species that showed a positive metal uptake resulting from sludge applications can be considered for evaluation of factors that affect response. Environmental factors that influence the metal accumulations are reviewed in detail in Sections 2 and 3. Where possible, these differences have been recorded along with data presented.

The recognition of these constraints served as the impetus for the USDA, CSRS Regional Technical Committee, W-124, to conduct a "uniform" field study at 15 locations across the United States in 1979 (see Section 2 for details). Data from this study were used in our delineation.

#### CUMULATIVE EFFECTS FROM ANNUAL SLUDGE APPLICATIONS

From the data provided by several researchers (Vlamis, et al., 1985; Soon and Bates, 1981; Chang et al., 1983; Hinesly et al., 1984) it was apparent that cumulative effects from annual sludge applications may be broken down into two categories, according to the metal inputs, namely: (1) zinc and cadmium when introduced with sludge at high levels (>100 kg Zn/ha/yr and >1 kg Cd/ha/yr) resulted in an increase in plant tissue metal over the years of sludge application, but the rate of increase decreased with time; and (2) typical sludges applied at agronomic rates to satisfy N requirement for crop growth cause Cd and Zn concentrations in plants to become greater than those of the control but Cd and Zn contents of plant tissue remained at a low, nearly constant level with each successive sludge application.

When sludges were applied to barley at rates equivalent to 8.6 kg Cd/ha and 714 kg Zn/ha annually over a 7-year period, the concentration of Cd in the barley straw increased from 0.26 to 3.39 mg/kg (Table 18, Vlamis et al., 1985). At the the same time the Zn concentration of the plant tissue increased from 113 to 820 mg/kg. During the experiment, the pH of the sludge-treated soil decreased from 5.5 to 4.8 which could account for some of the increase in metal accumulations by the barley. While the Cd and Zn accumulation in the vegetative part of barley was substantial, the barley grain harvested from sludge-treated soils contained considerably lower levels of Cd and Zn and frequently were not significantly different than those of the control. Swiss chard shows a similar Cd and Zn uptake response, except in greater amounts. With an

average input of 8 kg Cd/ha annually, Swiss chard took up 0.9 mg Cd/kg plant tissue the first year and increased to 18.0 mg Cd/kg by the tenth year. Again, as with the barley/sludge system, the pH decreased from its original 7.0 to 6.5. The long-term effects of sludge applications on the Cd and Zn levels in affected soils and Swiss chard are illustrated in Fig. 6 and 7 (Chang and Page, 1985, Personal communication). Soon and Bates (1981) measured Cd and Zn contents of corn and bromegrass on sludge-treated plots with a total accumulative Cd and Zn additions of 5.6 and 680 kg/ha, respectively, over an 8-year period. As with barley and Swiss chard, addition of sludge resulted in an increased metal content in the corn and bromegrass. Although the Cd and Zn inputs were high the successive additions of sludge did not result in a continuous increase of Cd and Zn concentrations in plants. The use of iron (Fe<sup>+++</sup>) and aluminum (Al<sup>+++</sup>) treated sludges in this experiment might have affected the results.

Table 19 summarizes the Cd and Zn contents in crops that were grown with sludges applied to satisfy nitrogen requirements of plant growth. In all cases, the concentration of Cd and Zn in the affected plant tissue remained constant over the years of application. The levels of Cd and Zn were greater for plants in the sludge-treated plots than the control plots.

There appeared to be a slight but statistically significant increase of Cu and Ni in plants grown on sludge-treated soils when they were compared to plants grown on untreated soils. Their levels did not appear to rise annually with the successive sludge inputs (Soon and Bates, 1981, Table 20). The crops (corn and bromegrass) were grown on calcareous soils which undoubtedly reduced plant availability of the added metals. An earlier report (Vlamis et al., 1978) on sludge application to a noncalcareous soil, however, supported the observations that Cu and Ni are generally not accumulated in plant tissue. In this study 324 kg Cu/ha and 97 kg Ni/ha were applied in the form of sludges over a three-year period and the barley grown on sludge-treated soils did not accumulate significant amounts of Cu and Ni. Studies by other investigators (Chaney, 1985, Personal communication) also showed little detrimental effect to plants at Cu and Ni input levels considerably higher than those reported by Soon and Bates (1981) and Vlamis et al. (1978). In sludge treated soils maintained at pH > 6.0, phytotoxicity from sludge applied Cu and Ni accumulation has rarely been reported (Marks et al., 1980).

From the data reviewed, there are pathways in the sludge-soil-plant system by which potentially harmful metal elements can accumulate in plant tissue through land applications of sludge. Amounts absorbed by plants are small and usually accounted for <1% of the inputs from sludge.

# SINGLE VS. MULTIPLE APPLICATION

Depending on the way sludges are applied, plants often respond to Cd and Zn introduced into soils in a different manner. Response curves (i.e., metal input from sludge applications vs. metal levels in plants) generated from single sludge additions usually have steeper slopes than response curves generated from multiple sludge additions which have the same total input spanned over a period of time. This would imply that the relationship between total applied metal and the resulting metal content in plants is not necessarily unique. Based on the results of a greenhouse pot experiment, the relationship appeared to be a function of the annual application rate (Ryan, 1982).

Results from the W-124 experiment were used to illustrate the patterns of 2n and Cd concentrations in plants with multiple sludge applications and a single sludge addition which had the equal total input. To summarize the data from various locations into a single diagram, we converted metal concentrations in plant tissue into "relative metal increment of plant tissue" (RMI) which is the ratio of metal increment of plants for a given year (i.e., metal concentration of affected plant tissue minus metal concentration of the control plant) to the first year metal increment of plants receiving 20 mt/ha treatment.

The graphic illustration of the data may be divided into several regions. Under the multiple sludge applications, the line of RMI=0 represents the metal concentration of plants equal to the background metal concentration (Fig. 8). The line RMI=1 represents non-additive effect which indicate, with subsequent sludge inputs, the increment of metal concentration in affected plant tissue are equivalent to that of the first year. The additive effect of multiple sludge inputs on metal contents of plant tissue is represented by the 1 to 1 line that passes through RMI=0. There was a wide range in the relative metal increments of each location, and in one occasion the RMI even exceeded the strictly additive regime. The mean annual RMI for all locations, however, were approximately 1 (0.86-1.08) indicating non-additive effects due to multiple sludge applications. Sometimes, the relative metal increments of the plant tissue in subsequent years was significantly lower than increments of the first year.

A large single sludge application (100 mt/ha in this case) produced a high plant tissue metal concentration in the crop immediately following the sludge application (Fig. 9). This large single sludge application produced a sharp rise of Zn levels in plant tissue. In 3 out of the 11 cases, the first year metal increment of plant tissue exceeded those calculated by the strictly additive rule with first year metal increment of the 20 mt/ha/yr as the reference point. The RMI of successive crops from the single sludge addition, however, decreased. By the time when inputs from the multiple applications had reached the same amounts as with the single application (year 5 in this case), the plant tissue metal increment of the single sludge application was not significantly different from that of the multiple sludge applications.

In the early days of land application studies, several researchers hypothesized that organically complexed metals in soils were less available to plants than uncomplexed metals. When sludge applications were terminated, soil microbial activity would reduce organic matter levels of the sludge-amended soils resulting in a higher availability of sludge-borne metals (Chaney, 1973; Haghiri, 1974; Brown, 1975). But the long-term observations made in field experiments show that the plant availability of metals in sludge-treated soils either remained unchanged or was reduced with time after cessation of sludge applications (Touchton et al., 1976; Dowdy et al., 1978).

Data from a field experiment in Illinois showed that where sludge was applied annually for three consecutive years at the agronomic nitrogen rates on silt loam soils, Cd and Zn concentrations in leaves, stover and grain of corn were increased significantly by the sludge addition. After sludge applications were terminated, concentrations of these metals in aerial parts of corn plants, although still higher than the control, decreased with each successive corn crop (Hinesly et al., 1979). Three years after sludge additions ceased, Cd concentrations of corn grain from the sludge-affected soils had receded to levels similar to those from control plots and levels of the metals in leaves and stover from sludge-treated plots were slightly higher than those from control plots. Webber and Beauchamp (1979) and Dowdy et al. (1978) reported similar patterns of metal accumulation in plant tissue.

Crops grown on a soil which received annual sludge applications exhibited a slight but significant increase in Cd and Zn concentrations of plant tissues (Hyde et al., 1979) immeditiely following the termination of sludge applications at this location. Chang et al. (1982) grew two winter wheat crops and observed that Cd and Zn concentrations of wheat grain and straw from sludge-treated plots were slightly higher than those from control plots. Concentrations of these metals, however, were well within normal ranges of concentrations found in wheat grown on uncontaminated soils. Similar results were found by Hinesly (1985) for Cd, Cu, Ni, and Zn contents in soybeans and wheat grown on plots of silt loam soil where sludge applications were terminated after four and six years of annual applications.

Even for soils that received repeated heavy sludge applications and for plants that were sensitive to metals in the soil, there was little indication that the availability of sludge-borne heavy metals would rise upon termination of sludge applications. In one field trial, the spring and fall split applications of composted sludge at rates ranging from 22.5 to 180 mt/ha/yr on one-half of the experimental field was discontinued after the 6th year. The Cd and Zn contents of plants (Swiss chard and radish) harvested from the area no longer receiving sludges remained elevated but concentrations were lower than those obtained from the area where sludge applications continued (Chang and

Page, 1985, Personal communication). For the six croppings (3 years) following the interruption of sludge applications, the metal concentrations of harvested plants remained at levels similar to or less than those at the time sludge application was terminated (Tables 21 to 26).

Based on the data, there is no evidence that the bioavailability of metals in sludge-treated soils will rise with time after terminating the sludge application. Unless chemical conditions of the sludge-treated soils are altered or a metal sensitive plant species is planted, there is no indication that plant uptake of metals should increase with time following termination of sludge application.

#### CONCLUSIONS

- Application of Cd and Zn to soils from municipal sludge will cause the Cd and Zn concentration of plants grown on these soils to exceed those of the untreated controls. When the sludge is applied at rates to satisfy the N requirement of the crop grown the Cd and Zn contents of plant tissue remain at nearly constant levels with successive sludge applications.
- 2. In sludge treated soils maintained at pH >6.0, Cu and Ni contents of the tissue from plants grown on these soils may become slightly elevated. Phytotoxicity from sludge-applied Cu and Ni, however, has rarely been reported.
- 3. Given adequate time for sludge to equilibrate with the soil, metal concentration of the affected plant tissue would be determined by the total amounts of metals in the soil and would not be affected by the methods of sludge application (e.g., single addition vs. multiple applications to yield the same total application as the single addition).
- 4. Plant availability of sludge-borne metals is highest during the first year sludge is applied. Using the first year response curve generated by a large single sludge addition will overestimate metal accumulation in vegetative tissue from plants grown in well stabilized sludge/soil systems.
- 5. There are no field data to indicate that trace element concentration in plant tissue will rise after the termination of sludge applications if chemical conditions of the soil remain constant. Cadmium and zinc levels of plants grown in soils which are no longer receiving sludges either were not significantly different from the pretreatment levels or decreased with time.

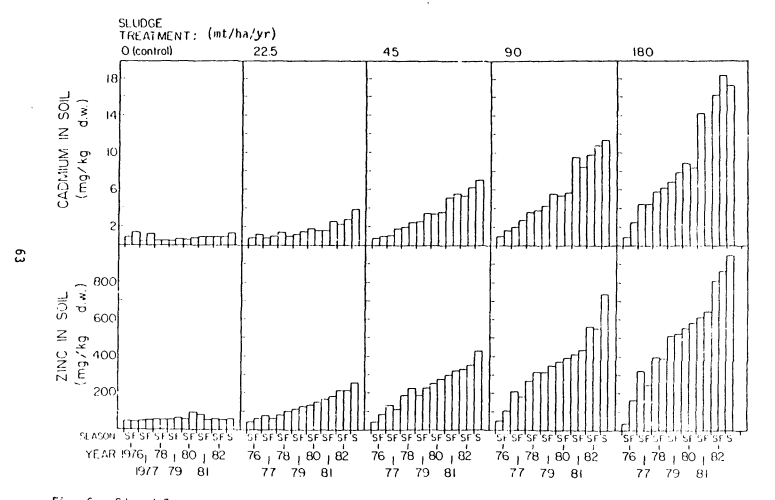


Fig. 6. Cd and Zn concentrations of composted sludge treated Ramona sandy loam (Chang and Page, 1985).

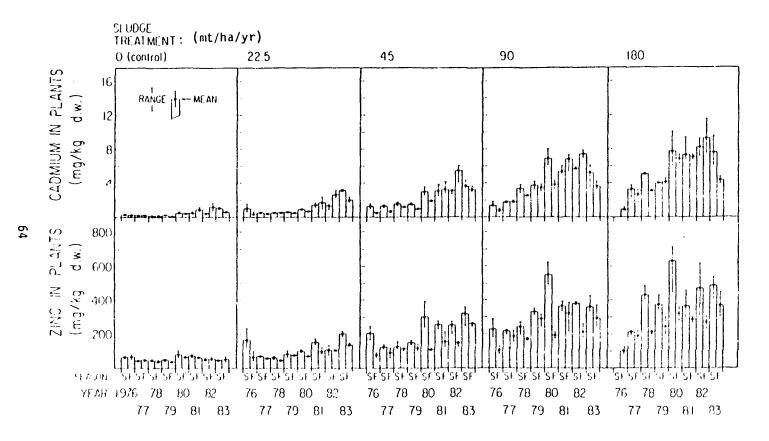


Fig. 7. Cd and Zn content of Swiss chard harvested from soils receiving biannual (spring and fall) sludge application from 1976-1983 (Chang and Page, 1985).

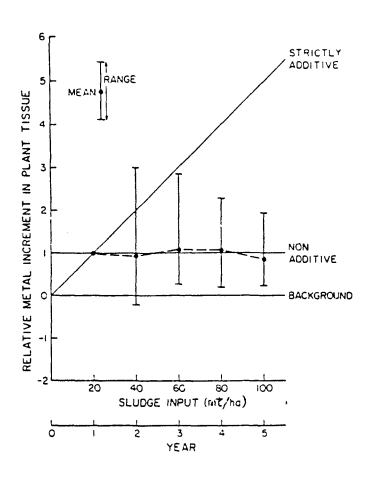


Fig. 8. Relative Zn increments of barley leaf receiving annual sludge addition of 20 mt/ha for five years (calculated with data from 11 of the 15 experimental sites of W-124).

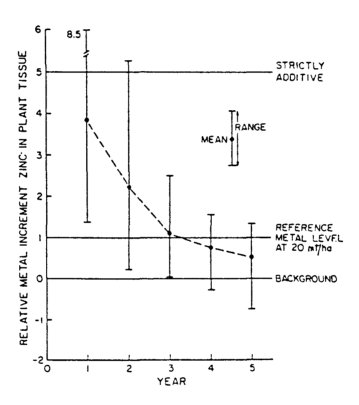


Fig. 9. Relative Zn increments of barley leaf receiving 100 ml/ha onetime sludge application (calculated with data from 11 of the 15 experimental sites of W-124).

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TABLE 18. CADMIUM AND Zn CONTENTS OF PLANT TISSUES WHEN SLUDGES WERE APPLIED ANNUALLY AT HIGH RATES.

						No.	of succ	essive a	nnual app	lications				
	Plant	Total metal	pH Initial/	1	2	3	-4	5	6	7	8	9	10	
Metal/Crop	parts	inputs (kg/ha)	final				Hetal	in plant	tissue (	mg/kg)				Reference
Cd/Barley Barley	straw straw	60 0	5.5/4.8 5.5/5.5	0.26 0.09	0.23 0.04	0.55 0.07	0.63 0.06	0.85 0.04	1.61	3.39 0.06	-	-	-	Vlamis et al. 1985
Corn Bross-	stover above	5.44	7.5/6.7	0.38	0.54	0.80	0.70	0.53	0.36	0.39	0.57	-	-	Soon & Bates 1981
grass	ground	6.08	7.4/7.2	0.09	0.39	0.23	0.20	0.28	0.27	0.30	0.45	-	-	Soon & Bates 1981
Swiss	above													
chard Swiss	ground above	80	7.0/6.2	0.9	2.7	3. 2	4.2	6.9	7.1	9.4	4.4	13.1	18.0	Chang & Page 1985
chard Swiss	ground above	20	7.1/6.5	0.5	0.7	1.2	1.0	2.0	3. 3	5.5	3. 2	7.2	8.7	Chang & Page 1985
chard	ground	0	7.2/7.5	0.3	0.3	0.2	0.1	0.4	0.9	1. 2	0.6	0.8	1.7	Chang & Page 1985
Zn/Barley	straw	5000	5.5/4.8	113	150	248	34 1	402	455	820	-	-	-	Vlamis et al. 1985
Barley	straw	0	5.5/5.1	12	58	71	51	37	116	68	-	-	-	
Corn Brom-	stover above	680	7.5/6.8	41	69	103	88	85	77	74	65	-	-	Soon & Bates 1981
grass	ground	672	7.4/6.9	28	41	39	34	40	42	44	65	-	•	Soon & Bates 1981
Swirs	above													
chard Swiss	ground above	6400	7.0/6.2	105	19 1	216	249	324	209	275	37 2	. 844	997	Chang & Page 1985
chard Swiss	ground above	1600	7.1/6.5	79	90	111	116	110	155	146	26 1	345	567	Chang & Page 1985
chard	ground	0	7.2/7.5	67	50	40	39	66	67	56	52	68	34	Chang & Page 1985

TABLE 19. CADMIUM AND Zn CONTENTS OF PLANT TISSUE WHEN SLUDGES WERE APPLIED AT AGRONOMIC RATES.

		Yrs. of				No.	of succ	essive a	nnual ap	plicatio	ns		
	Plant	sludge applica-	Total metal	pH Initial/	1	2	3	4	5	6	7	8	
Metal/Crop	parts	tion	inputs	finel			Metal 1	n plant	tissue (	mg/kg)			Reference
Cd/Barley	straw	7	2	5.5/7.0	0.23	0.07	0. 16	0.20	0.12	0.26	0.24	_	Vlamis et al. 1985
Barley	straw	7	0	5.5/5.8	0.08	0.04	0.06	0.08	0.09	0.08	0.04	-	
Corn	stover whole	8	0.72	7.4/7.3		0.30	0.29	0.27	0.25	0.17	0.16	0.18	Soon & Bates 1981
Brome-													
grass	plant	8	1.6	7.4/7.4	0.04	0.16	0.08	0.12	0.11	0.08	0.12	0.11	Soon & Bates 1981
Barley	grain	6	5.5	6.1/6.7	0.07	0.05	0.04	0.04	0.04	0.05	-	-	Chang et al. 1983
Barley	grain	6	0	6.3/7.1	0.07	0.04	0.01	0.04	0.04	0.04	-	-	•
Barley	grain	6	5.5	7,1/7,1	0.06	0.02	0.01	0.04	0.04	0.05	-	-	Chang et al. 1983
Barley	grain	6	0	7.1/7.1	0.03	0.02	0.05	0.04	0.04	0.04	-	-	,
In/Barley	straw	7	133	5.5/7.0	66	46	52	46	44	93	54	-	Vlamis et al. 1985
Barley	straw	7	0	5.5/5.8	45	41	52	36	30	98	57	-	
Corn Brom-	stover whole	8	112	7.5/7.3	26	23	35	30	31	30	29	27	Soon & Bates 1981
grass	plant	8	192	7.4/7.4	20	24	25	23	27	24	25	32	Soon & Bates 1981
Barley	leaf	6	80	6.1/6.7	19	22	23	32	32	47	-	_	Chang et al. 1983
Barley	leaf	6	0	6.3/7.0	16	13	25	21	18	22	-	-	<b>,</b>
Barley	leaf	6	80	7.1/6.9	24	20	17	29	25	26	-	_	Chang et al. 1983
Barley	leaf	6	0	7.1/7.1	20	14	18	22	21	21	-	_	

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TABLE 20. CADMIUM AND NI CONTENTS OF PLANT TISSUES FROM SLUDGE-TREATED SOILS.

			Yrs. of				No.	of suc	cessive	annual	applicat	ions		
Metal	Crop	Plant parts	sludge applica- tion	Total metal inputs	pH Initial/ final	7	2	3 Metal	in plant	tissue	6 (mg/kg)	7	8	Reference
Hi	Carn	staver	8	507	7.5/8.0		1.5	1.5	0.5	2.3	1.0	1.2	1.9	Soon & Bates 1981
	Corn Brom-	stover	8	63	7.5/7.8	-	1.0	1.6	0.4	1.9	0.7	0.6	0.7	
	grass Brom-	whole plant	8	624	7.4/8.0	-	0.2	1.2	0.6	6.0	4.3	4.2	4.8	Soon & Bates 1981
	grass	whole whole	8	156	7.4/7.7	-	0.3	1.6	0.5	3.5	2.0	2.2	1.9	
Cu	Corn	stover	8	354	7.5/6.7	-	12	10	7	11	8	9	8	Soon & Bates 1981
	Corn Brom-	stover	8	88	7.5/7.3	-	8	8	6	9	6	6	6	
	grass Brom-	whole plant	8	302	7 5/6.7	-	11	15	14	10	15	14	14	Soon & Bates 1981
	grass	•	8	98	7.4/7.3	-	8	10	9	10	10	7	8	

TABLE 21. CADMIUM CONCENTRATIONS (mg/kg) OF SWISS CHARD GROWN ON SLUDGE-TREATED SOILS (CHANGE AND PAGE, 1985).

			22.5 mt/h	a/yr#	45 mt/ha	/y <b>r</b> ¥	90 mt/h	a/yr#	180 mt/h	a/yr#
Year	Season	Control	Continued Application	Termi- nated	Continued Application	Termi- nated	Continued Application	Termi-	Continued Application	Teran-
1976	Spring	0.72	1.05	<del></del>	1.78		1.30	<del></del>		<del></del>
	Fall	0.28	0.35	-	0.52	•	0.88	-	0.88	-
1977	Spring	0.20	0.58	-	1. 35	•	1.75	-	3. 28	-
	FALL	6.25	0.40	-	0.72	-	1.78	-	2.12	-
1978	Spring	0.20	0.48	-	1.60	•	3, 38	-	5.12	_
	Fall	0.20	0.50	-	1. 20	-	2.58	-	3. 18	-
979	Spring	0.28	0.62	-	1.55	-	3.75	-	4.02	_
	Fall	0.12	0.52	•	0.98	-	3.48	-	4.15	-
980	Spring	0.52	0.90	-	3.05	-	6.85	-	7.75	-
	Fall	0.42	0.72	-	1.95	-	3.80	-	6.88	-
981	Spring	0.50	1.45	-	3. 15	_	5.40	-	7.35	-
	Fall	0.90	1.75	-	3, 36	-	6.32	-	7.12	-
982	Spring	0.50	1.35	1.70	3. 15	3.80	5.75	6.10	8.32	11, 10
	Fall	1. 20	2.68	2.30	5.50	5.20	7.45	6.70	9.38	8.70
983	Spring	0.70	3, 18	2.00	3.72	3.80	5.29	4.60	7.65	7.20
	Fall	0.50	2.02	1.30	3. 32	2.50	3.65	3.80	4.38	3.70
984	Spring	1.40	4.82	2.30	5.68	5.60	6.30	6.80	9.32	5.80
<del>-</del> -	Fall	0.8	3.00	1.40	7.20	4.80	9.80	7.40	13. 10	10.70
1985	Spring	1.4	3. 30	1.00	11.3	3.60	18.30	10.90	21.60	17.20
-	Fall	1.5	4.70	2.80	8.70	5.60	13.60	8.60	18.00	9.50

M<sub>t</sub>xperimental field was split after fall 1981 where one-half continued to receive sludge and the other half no longer received sludge but was cropped.

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TABLE 22. ZINC CONCENTRATIONS (mg/kg) OF SWISS CHARD GROWN ON SLUDGE-TREATED SOILS (CHANG AND PAGE, 1985).

			22.5 mt/ha	/yr*	45 mt/ha/	yr*	90 mt/ha/y	r*	180 mt/ha/ Continued	y-*
Year	Season	Control	Continued Application	Termi-	Continued Application	Termi- nated	Continued Application	Termi-	Continued Application	
1976	Spring Fall	65 67	167 67	-	170 79		229 105	-	105	-
1977	Spring Fall	45 50	72 60	-	124 90	-	215 189	-	215 191	-
1978	Spring Fall	48 40	62 47	-	127 111	-	24 1 17 2	:	4 32 2 16	
1979	Spring	51 39	85 78	-	149 116	- -	332 289	-	378 249	-
1980	Spring Fall	81 66	10 1 7 1	-	30 1 110	:	550 19 <i>2</i>	-	633 324	-
1981	Spring Fall	76 67	15 2 96	-	297 155	-	368 322	-	366 290	-
1982	Spring Fall	61 67	107 106	127 88	25 1 146	278 174	38 2 213	337 240	475 275	465 282
1983	Spring Fall	52 54	20 3 1 38	16 3 94	320 26 1	268 225	36 1 29 3	313 386	490 3/2	404 304
1984	Spring Fall	84 66	20 2 188	95 98	176 345	218 290	319 554	348 554	37 3 844	230 730
1985	Spring Fall	7 3 37	232 209	10 1 99	389 567	25 2 2 18	725 768	385 5 10	1000 997	548 562

<sup>\*</sup>Experimental field was split after Fall 1981 where one-half continued to receive sludge and the other half no longer received sludge but was cropped.

TABLE 23. CADMIUM CONCENTRATIONS (mg/kg) OF RADISH LEAF GROWN ON SLUDGE-TREATED SOILS (CHANG AND PAGE, 1985).

			22.5 mt/ha		45 mt/ha/		90 mt/ha/y	r*	180 mt/ha/	
Year	Season	Control	Continued Application	Termi- nated	Continued Application	Termi- nated	Continued Application	Termi- nated	Continued Application	
1976	Spring Fall	0.2 0.2	0.4	-	0.4	-	0.5 1.1	-	- 1.5	-
1977	Spring Fall	0.3 0.5	0.6 0.7	:	0.9 1.9	- -	1.5	-	3.4 5.0	-
1978	Spring Fall	0.2 0.3	0.2 1.0	-	1.2 1.4	-	1.4 2.0	-	1.9	-
1979	Spring Fall	0.5 0.7	2.0 2.5	-	2.5 3.6	-	3.1 5.9	-	3.9 7.4	-
1980	Spring Fall	1.0 0.6	1.4	-	2.0 3.8	-	3.2 3.8	<u>-</u>	5.2 8.8	-
1981	Spring Fall	1.1	1.8	-	2.7 4.2	-	5.4 6.1	-	7.0 8.4	-
1982	Spring Fall	0.8 1.8	2.1 3.7	1.2	3.1 5.6	1.9 3.5	5.9 7.2	2.9 4.8	6.6	6.3 3.0
1983	Spring Fall	1.4	2.8 2.2	2.1 1.5	3.7 2.9	2.7 2.2	4.9 3.7	3.6 2.4	6.8 5.3	6.1 3.7
1984	Spring Fall	0.9	2.5 2.8	1.2	6.6 6.7	2.6 2.1	8.0 8.5	3.2 3.6 :	8.8 14.2	7.1 7.5
1985	Spring Fall	0.7 0.6	4.2 3.4	1.7 1.6	11.8 5.5	3.7 3.2	13.7 7.9	9.5 4.7	14.9 10.8	11.3

<sup>\*</sup>Experimental field was split after Fall 1981 where one-half continued to receive sludge and the other half no longer received sludge but was cropped.

TABLE 24. CADMIUM CONCENTRATIONS (mg/kg) OF RADISH TUBER GROWN ON SLUDGE-TREATED SOILS (CHANG AND PAGE, 1985).

			22.5 mt/ha	ı/yr¥	45 mt/ha/	yr*	90 mt/ha/y	r*	180 mt/ha/	yr*
Year	Season	Control	Continued Application	Termi-	Continued Application	Termi- nated	Continued Application	Termi-	Continued Application	
1976	Spring Fall	0.2 0.1	0.2 0.2	-	0.2 0.3	-	0.2 0.4	-	- 0.7	-
1977	Spring Fall	0.1 0.2	0.3 0.4	-	0.3 0.7	-	0.4 0.9	-	0.9	-
1978	Spring fall	0.2	0.3	<del>-</del>	0.4	-	0.5 1.2	-	0.7 1.8	-
1979	Spring F411	0.2 0.3	0.4 1.0	-	0.5 1.1	-	0.7 1.5	-	1. 1 2. 2	:
1980	Spring fall	0.3 0.3	0.5 0.8	-	0.5 1.4	-	0.8 2.5	-	1.0 2.7	-
1981	Spring Fall	0.3 0.5	0.4 0.5	-	0.6 0.6	-	0.9 0.8	-	1.0 0.9	-
1982	Spring Fall	0.4	0.6 0.5	0.4	1.0 0.8	0.7 0.5	1.4 1.3	1.0 0.9	1.4 1.4	1.4
1983	Spring Fall	0.8 0.6	0.7	0.7 0.9	0.8 1.6	0.7 1.1	0.7 2.1	0.7 1.5	0.9 2.2	0.7 2.1
1984	Spring Fall	0.3 0.3	0.9 1.1	1.0 0.6	1.0 1.6	0.8 1.0	1.2 1.9	0.9	1. 2 2. 7	1.0 2.3
1985	Spring Fall	0.6 0.3	1. 1 1. 9	1.0 1.1	1.0 3.1	1. 1 1. 9	2.5 4.0	1.3 3.8	3.5 5.6	2.6 4.3

<sup>\*</sup>Experimental field was split after Fall 1981 where one-half continued to receive sludge and the other half no longer received sludge but was cropped.

TABLE 25. ZINC CONCENTRATIONS (mg/kg) OF RADISH LEAF GROWN ON SLUDGE-TREATED SOILS (CHANG AND PAGE, 1985).

			22.5 mt/ha	/yr¥	45 mt/ha/	уг#	90 mt/ha/y	r*	180 mt/ha/	yr <del>.t</del>
Year	Season	Control	Continued Application	Termi-	Continued Application	Termi-	Continued Application	Yermi-	Continued Application	Termi-
1976	Spring Fall	55 42	56 58	-	62 75	-	55 78	-	173	-
1977	Spring Fall	45 55	46 82	-	56 64	- -	73 102	-	79 138	-
1978	Spring Fall	46 39	46 64	<b>-</b> ,	71 90	-	93 134	•	121 177	-
1979	Spring Fall	38 4.7	5 <i>2</i> 95	-	74 232	-	129 223	-	198 254	-
1980	Spring Fall	42 40	58 66	-	81 121	-	129 201	-	19 3 294	-
1981	Spring fall	46 50	73 84	-	117 134	-	216 176	:	266 275	-
1982	Spring Fall	45 42	74 79	56 63	109 142	78 82	206 192	120 116	234 283	200 184
1983	Spring Fall	41 48	74 95	51 57	107 159	76 90	167 230	110 133	27.5 35.4	196 225
1984	Spring Fall	50 70	97 147	62 73	195 26 2	97 120	316 38 3	180 171	421 627	24 <b>4</b> 286
1985	Spring Fall	32 44	10 <b>3</b> 106	53 62	186 166	85 93	305 232	14 3 120	45.2 36.3	231 179

<sup>\*</sup>Experimental field was split after fall 1981 where one-half continued to receive sludge and the other half no longer received sludge but was cropped.

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TABLE 26. ZINC CONCENTRATIONS (mg/kg) OF RADISH TUBER GROWN ON SLUDGE-TREATED SOILS (CHANG AND PAGE, 1985).

			22.5 mt/ha	/yr*	45 mt/ha/	yr#	90 mt/ha/y	r*	180 mt/ha/	yr*
Year	Season	Control	Continued Application	Termi-	Continued Application	Termi-	Continued Application	Termi-	Continued Application	Termi
1976	Spring	34	38 57	-	39 70	-	39 47		-	-
	fall	48	. 5/	•	70	•	4/	-	92	-
1977	Spring	38	31	-	37	-	51	-	61	-
	Fall	31	46	-	56	-	62	-	84	-
1978	Spring	20	32		57	-	94	-	54	_
	Fall	42	63	-	86	-	124	-	158	-
1979	Spring	36	59	-	56	-	71	-	88	-
	Fall	37	46	-	71	-	97	-	135	-
1980	Spring	34	41	-	51	-	63	-	76	
	Fall	49	58	-	82	•	131	-	144	-
1981	Spring	33	39	-	52	_	70	-	·. 11	
•	fall	37	40	•	52	-	67	-	79	-
1982	Spring	42	50	46	71	57	99	73	105	93
	Fall	31	40	30	53	38	80	46	95	12
1983	Spring	30	42	36	44	40	60	41	82	58
	Fall	46	58	39	75	51	10 1	76	117	101
1984	Spring	31	44	40	69	48	89	63	113	76
	Fall	40	84	50	118	68	154	90	224	157
1985	Spring	24	38	33	68	39	94	49	116	12
	Fall	42	78	52	88	64	110	79	132	84

<sup>#</sup>Experimental field was split after Fall 1981 where one-half continued to receive sludge and the other half no longer received sludge but was cropped.

# SECTION 5

# TRANSFER OF SLUDGE-APPLIED TRACE ELEMENTS TO THE FOOD CHAIN

#### INTRODUCTION

Assessment of the likelihood of risks to humans, livestock, and wildlife from potentially toxic constituents in sewage sludge applied to land requires a knowledge of the potential for transfer of each constituent from the sludge or sludge-soil mixture to crops and to animals (including humans) which ingest sludge, sludge-soil mixture, or crops grown on the sludge-amended soil. Transfer of sludge constituents from soil to crops is predominantly a function of: (1) the constituent; (2) soil pH; (3) characteristics of the applied sludge and cumulative sludge application rate; and (4) the crop species and cultivar grown. Each of these factors can be associated with a 2-fold or greater change in plant concentration of some trace elements.

Earlier attempts to estimate food-chain transfer have used plant uptake slopes obtained by linear regression of the constituent's concentration in edible crop tissue and the amount of the constituent applied [(mg constituent/kg dry crop) per (kg constituent applied/ha)]. In the long-term, plant uptake of sludge-applied Cd and Zn is a curvilinear (plateauing) response to the cumulative application rate of applied constituent in a sludge. Further, the plateau reached is a function of the concentration of the constituent in sludge and other sludge properties such as Fe added during sludge processing.

Estimation of food-chain transfer is critical to valid estimation of the potential for risk. At present, these estimates are best made by considering (1) the relative increased uptake of constituents by various crops from sludge-amended soil under responsive conditions; (2) the rate of ingestion of different crops by the U.S. population (e.g. Pennington, 1983); (3) the demonstrated bioavailability of the increased amounts of an element in sludge-grown crops or ingested sludge; and (4) an appropriate transfer coefficient from sludge-amended soil to edible crop tissues [i.e., the increase in crop trace element residue (above that in the same crop grown on background soils) at the plateau reached on sludge-amended soil].

Ryan et al. (1982) developed an approach for estimating food-chain transfer of sludge-applied Cd. While this model has to be modified to

account for curvilinear response to sludge-applied Cd, many other parts of the approach remain valid. In particular, the aggregate increased foodchain exposure to a sludge-borne constituent for the exposed population can be expressed in terms of a constant times the element transfer coefficient (height of the plateau above that for untreated soil) for an indicator crop such as lettuce. Individuals who grow, on acidic sludge-amended soils, a significant portion of the garden vegetables they ingest are generally believed to be the individuals most likely to have increased dietary Cd due to land application of sludge. For chronic lifetime (50 year) Cd exposure, estimation also relies on U.S. food intake estimates (q fresh weight/day) from Pennington (1983), and solids content of foods from USDA data bases (Adams, 1975). Because response curves or plateaus have not yet been evaluated for every crup consumed by the U.S. population, uptake by various food crops must be represented by the FDA food groups. For Cd, based on many data sources, the response of minor food crops remains well represented by the FDA food groups.

On the other hand, the use of a "Cd-accumulator" crop to represent increased Cd uptake by all crops in a food group has been criticized. For example, lettuce and broccoli were both listed in the leafy vegetable food group, but have at least 10-fold different response slopes. Root vegetables and garden fruits also include high and low element accumulating crops. These wide crop differences have caused an excessive estimated increase in food-chain Cd transfer (providing a hidden safety factor). Thus, food group aggregate transfer response slopes have to be adjusted for the proportion of low and high accumulating crops in each food group. Each food group can still be represented by a constant times the response of lettuce. This approach appears to be appropriate based on the findings of this workshop.

Many studies have shown significantly higher uptake of Cd, Zn, and Ni at lower soil pH compared to neutral soil pH (see Logan and Chaney, 1983; CAST, 1980). Exceptions were reported by Pepper et al. (1983) and Hemphill et al. (1982) in which corn silage was unchanged or slightly higher in Cd on limed sludge-amended soil. One possible reason for this exception is that corn differs from other crops in its mechanism of obtaining Fe from soil. Recent research on the mechanism plants use to obtain Fe from soil has shown that the Graminae (corn, wheat, barley, rice. oat, sorghum, etc.) excrete an organic chelating agent which facilitates Fe diffusion to the root and Fe uptake (Sugiura and Nomoto, 1984; Römheld and Marschner, 1986), while plants in other families do not excrete chelators. These compounds also chelate Cd, Cu, Zn, and other microelements in the presence of Fe (unlike bacterial siderophores, which have very high selectivity for ferric iron). Non-specific chelating agents added to soils are also known to facilitate diffusion and plant uptake of Zn, Cd, and other elements. Thus, Cd uptake by Graminae can have a lesser response to soil pH change than other species, depending on the availability of soil Fe.

Most studies of soil pH effect on metal uptake by crops have studied the pH range from about 5.0 to 7.5. In a report by Francis et al. (1985),

S in a coal gasification waste caused soil pH to fall to 4.0, and the ryegrass crop grown on this extremely acidic soil was stunted and very high in metals even though metals in the soil and coal waste were not very high. Extreme acidic soil pH levels associated with severe pH mismanagement may allow crop metal residues not otherwise considered in this report.

Important problems remain in estimating the variance in potential risk due to unusual individual dietary selection patterns, and to individual variation in retention of potentially toxic constituents in foods. For example, it would be useful to know the statistical distribution of increased Cd intake among individuals consuming self-selected diets using crops grown on sludge-amended acid gardens. If these data were available, one could design Cd limits to protect individuals with the 95th percentile of increased exposure. Several papers have noted that the relevant information needed on variation in food intake for evaluation of chronic risk from food Cd is lifetime (50-year) variation in intake of foods, rather than the 1-day variation now available (Dean and Suess, 1985). Although the mean 1-day ingestion of foods for a population is estimated well by a large survey of individual 1-day intakes, the variance in long-term average daily intake is greatly over-estimated by the variance of 1-day intakes for a population (Beaton et al., 1983; Todd et al., 1983; Block, 1982). Beaton et al. (1983) and Sempos et al. (1985) found that intra-individual variance was greater than the inter-individual variance for multiple 1-day observations. Much smaller variances among individuals are associated with long-term dietary intakes, especially if one considers major dietary subgroups (e.g. vegetarians) separately. Thus, present approaches for estimating the fraction of the population ingesting more Cd than some limit (e.g., 95th percentile, or fraction > 71  $\mu g$  Cd/day), based on variances in population 1-day food intakes, would greatly overestimate the lifetime Cd intake by individuals.

Further, the protection of individuals varying in retention of ingested elements must consider the effects of important nutritional interactions on element bioavailability. There are several clear examples of this source of error in estimating risk. One is the health effects to Japanese farmers who ingested rice grown in soils rich in Cd and In due to contamination by Zn-ore wastes. In contrast to essentially all other food crops, rice is grown in flooded soils. Cadmium and Zn uptake by rice is normally kept very low because insoluble metal sulfides are formed in the soil. However, some metal uptake occurs because the roots obtain oxygen by air channels within the plants. It was found that rice metabolized Cd and In differently in these anaerobic rice soils than patterns common to other food crop species in aerobic soils. Although both soil Cd and soil Zn were greatly increased (up to 10 mg Cd/kg and 1200 mg Zn/kg), rice grain Cd rose up to 100-fold while grain Zn was unchanged (Tsuchiya, 1978, page 237). Further, during preparation of polished rice, much of the Ca, Zn, and Fe in brown rice is removed during milling, while a much lower fraction of the Cd is removed (Pedersen and Eggum, 1983; Chino, 1981; Chino and Baba, 1981, Yoshikawa et al., 1977). Lastly, rice Fe has very low bioavailability (Hallberg et al., 1974, 1977). All these factors [crop metal uptake characteristic, food processing, and effect of nutrient (Fe, Zn, and Ca)

status on human Cd retention] favored Cd retention by the farmers. Bioavailability of Cd was thus high, and human disease resulted.

In another Cd exposure case, individuals who consumed large amounts of Cd-rich oysters had higher dietary levels of Zn, Fe, and Ca, levels which were more like those in the normal U.S. diet. The oyster Cd had very low apparent bioavailability based on Cd in blood and urine compared to the effect of smoking on Cd in blood and urine (Sharma et al., 1983; McKenzie et al., 1982). These findings suggest that predictions of human retention of Cd from Western-type diets way be less than values currently in use.

Another example comes from the availability of sludge Cu to live-stock. In contrast to Cu salts mixed with diets, Cu in sludge fed to livestock has low bioavailability. Generally, ingested sludge lowers liver Cu stores rather than causing Cu toxicity, even though equal levels of soluble Cu salts would poison the animals (Decker et al., 1980; Bertrand et al., 1981; Baxter et al., 1982).

MODELING THE EFFECT OF CROP VARIATION IN INCREASED TRACE ELEMENT ACCUMULATION IN RESPONSE TO SLUDGE APPLICATION

The extent of increase in trace element concentration above control for crops grown on a sludge-amended soil is very strongly affected by crop species. Besides crop species variation in response, sludge Cd concentration and soil pH very strongly affect the plant Cd:soil Cd relationship. Other factors, such as crop cultivar and Fe level in the sludge, may have a substantial effect on the plant Cd:soil Cd relationship, while many other factors have affected results in some studies (soil organic matter, soil sesquioxides, pH buffering capacity of the soil, soil fertility, crop mycorrhizal infection, type of N fertilizer, and climatic factors).

An approach to estimate the relation between sludge-applied Cd, for example, and increased dietary Cd exposure is to integrate all crop response in terms of Cd uptake by a responsive reference crop such as lettuce. Relative to the responsive crop, the increased Cd uptake among crops due to sludge application has been reasonably consistent. However, few individual experiments have included a large number of crop species in a responsive sludge treatment to provide the relative increases in crop levels of potentially toxic constituents needed for dietary exposure assessments. The studies by Davis and Carlton-Smith (1980) and Carlton-Smith and Davis (1983) report the response of many crop species in one experiment. The conditions of these studies meet the constraints for appropriate techniques for sludge trace element risk assessment. They grew many crop species (some with multiple cultivars) on 2 soils collected from long-term sludge farms in England. They used large pots of soil (10 kg) in the greenhouse. Soil A had pH 6.7 and contained 5.8% organic matter and 7.4 mg Cd/kg soil. Soil B had pH 6.8 and contained 26% organic matter and 68.7 mg Cd/kg soil. Crop Cd ranged from near zero to about 8 mg/kg dry

weight. Relative crop Cd concentrations were similar between these 2 soils, and in good agreement with other research results.

In an effort to make maximum use of these data, Davis and Carlton-Smith developed tables of relative element concentrations for Cd, Zn, Cu, Ni and Pb. The concentration in each crop was expressed as a percentage of that in the crop with highest uptake of a given element, and the data were averaged across the 2 soils. The raw data for this study were obtained from Dr. R. D. Davis. In evaluating the raw data, it was noted that several data points were outliers, and these data points were deleted. The geometric mean element concentrations for the 2 soils (Table 27) were calculated; then a normal background Cd concentration in each group was subtracted from the geometric mean Cd level in that crop, and the ratio of Cd in crop X to the mean level of Cd in lettuce [(Cd in Crop X):(Mean Cd in lettuce)] was calculated (Table 28). The background Cd levels were estimated for crop groups, mainly relying on Wolnik et al. (1983, 1985); other field results were also considered and summarized by Korcak (1986, Personal communication).

Table 27 shows the geometric mean concentration of Cd, Ni and Zn in edible crop tissues of vegetable and grain crops. Similar results for relative uptake for total shoots of forage crops grown on 2 sludge-amended soils and 1 control soil are presented for Cd, Cu, Mo, Ni, and Zn in Table 29 from Carlton-Smith and Davis (1983). (We do not report their Pb data because the crops were grown near an urban area, and the Pb results are not representative of agricultural production areas).

The relative crop uptake tables remove factors other than crop species and cultivar. From the summarized data, it is not possible to ascertain the effects of other parameters (such as soil organic matter, soil pH, or sludge application rate) on relative metal uptake among crops.

Relative Cd uptake by crops was also evaluated in areas of naturally Cd-rich soils in Salinas Valley, Monterey Co., California (Table 30; Burau, 1980). Many paired samples of mature vegetable produce were obtained along with plow layer soil. The slope of the relationship between crop and soil Cd was reported for crops for which linear regression showed a significant slope. The Cd-enriched soils have quite similar properties, with Cd coming from geologic sources rather than sludge. Table 30 shows slopes and a relative uptake calculation similar to those used in Tables 28 and 29, with leaf lettuce set = 100.

Giordano et al. (1979) reported crop uptake of trace elements as affected by sludge application rate and soil heating. They continued the study for 2 years following application. Their results (unheated soil data) corroborate the extremely low increase of Cd in beans, cabbage, pepper. tomato, and the curcurbit family and low Cd increase in potato.

Although it has been clearly shown that crops differ in uptake of trace elements from the same soil, the biochemical/physiological basis for crop differences has not been explained. Basic research by Jarvis et al.

(1976) indicated that crops differed in Cd uptake by roots, and also differed in the fraction of root Cd translocated to shoots. Recently, Grill et al. (1985) found that many plant species made a family of cysteine-rich peptides (related to glutathione) which strongly chelate Cd, Zn, Cu, Pb, and Zn. Their "phytochelatins" may be synthesized in the fibrous roots and chelate absorbed metals, and thereby protect root metabolism and reduce trace element translocation to edible plant tissues (Grill et al., 1985; Rauser and Glover, 1984).

CROP CULTIVAR DIFFERENCE IN METAL UPTAKE FROM SLUDGE-AMENDED SOIL

Cultivars (also referred to as varieties, genotypes, selections or strains) within a crop species vary significantly in uptake of sludge-applied trace elements. Cultivar variation in Cd and other element uptake was evaluated because in agronomic management unrelated to sludge use (Foy et al., 1978) this source of variation had been found to be important in correction of trace element deficiencies (Fe, Cu, Mn, Zn) and in tolerance of plants to trace element toxicity (Al, Mn, Zn). The expected benefits from cultivar difference in tolerance or uptake of sludge-borne trace elements include: (1) the ability to select relatively metal-tolerant or metal-excluder (non-accumulator) cultivars for use in management of designed sludge farms; (2) determination of whether cultivar differences are great enough to require adjustment of dietary element risk assessments; and (3) identification of cultivars which could be used to reduce background levels of element ingestion from the general food supply.

At this time, a few crops have been studied under the conditions which generate results considered reliable for evaluation of the long-term effects of sludge-borne trace elements. Cadmium uptake by cultivars of corn, soybean, and lettuce have been studied in appreciable detail. Carrot, wheat, and some forage species have been studied, but to a lesser extent. Other studies are needed, particularly for crops which strongly absorb or exclude particular trace elements. As a general rule, cultivars have been found to vary by at least 2- to 5-fold from lowest to highest uptake response. However, a 30-fold variation was found in corn inbreds.

An extensive characterization of relative corn cultivar variation in uptake of a sludge-applied trace elements was reported by Hinesly et al. (1978, 1982). Uptake of Cd and Zn by 20 corn inbreds grown on long-term sludge-amended soils in the field were reported. These soils provided substantially increased plant-available Cd and Zn. The relative Cd and Zn concentration in the cultivars were recalculated as the geometric mean for three sludge rates. Leaf Cd ranged from 0.88 mg Cd/kg dry weight in inbred R805 to 30.3 mg Cd/kg dry weight in inbred B37. Grain Cd ranged from 0.05 mg Cd/kg dry weight in inbred H96 to 1.81 in inbred B37. The grain and leaf Cd concentrations were highly correlated as were the ranks among inbreds of grain Cd concentration and leaf Cd concentration. However, the grain Cd to leaf Cd concentration ratio of an inbred varied from 1.8 to 10.4. This wide range in grain Cd to leaf Cd ratio indicates that one

should not base a breeding program to lower grain Cd concentrations only on measuring seedling leaf Cd concentrations, nor should one use grain Cd results alone to select for Cd-excluder silage corn cultivars. On these same field plots, corn leaf Zn ranged from 44.2 to 152 mg/kg dry weight and grain Zn varied from 31.5 to 58.4 mg/kg. Although ranks and concentration of both Cd and Zn in corn inbreds were each well correlated over sludge rates, concentrations of Cd and Zn were not significantly correlated, nor were cultivar ranks for Cd and Zn concentrations significantly correlated. Thus, mechanisms controlling uptake or translocation to corn grain may vary for Zn and Cd.

Hinesly et al. (1982) then used the results from study of corn inbreds to make selected corn single-cross hybrids, and evaluated Cd and Zn uptake to leaves and grain, Cd and Zn distribution in milling fractions (Hinesly et al., 1979), and Cd bioavailability to laying hens (Hinesly et al., 1985). Single-cross corn hybrids were prepared which accumulated low or high levels of Cd into grain. Significant reduction in food-chain transfer of Cd is possible by selecting for Cd-excluder corn hybrids.

Bache et al. (1981) compared the accumulation of Cd and other elements by 8 corn hybrids adapted to upper New York State. In a greenhouse study with freshly applied air-dried sludge, a 6-fold range in grain Cd and 2-fold range in stover Cd were found among the hybrids tested. The possible range of response is probably fairly widely expressed in the wide range of corn hybrids of commercial importance in the U.S., but this range is not nearly as wide as seen for corn inbreds.

The Cd uptake to shoots of seedling plants of soybean cultivars grown on sludge-amended soil showed about a 4-fold range for 10 cultivars (Boggess et al., 1978). The investigators also reported Cd uptake from soils amended with Cd salts; although relative Cd concentrations among cultivars were similar, they were not related closely enough to rely on salt-Cd results to predict cultivar uptake on sludge-amended soils. Hinesly (1986, Personal communication) found a smaller range in concentration of Cd in grain of soybean cultivars grown to maturity in the field, and the ranks of grain Cd concentration among 3 cultivars was similar to the findings of Boggers et al. (1978) for seedling soybean shoots.

Following a study with 3 lettuce cultivars (Harrison, 1986a), Yuran and Harrison (1986) studied relative Cd uptake by 60 lettuce genotypes. They grew the 60 genotypes in 1 soil type, with 1 rate (90 mt/ha/yr) of 2 sludges (12 and 57 mg Cd/kg dry sludge), for 2 years. Soil Cd in year 2 was 0.09, 1.1, and 9.0 kg Cd/ha for the catrol, lower Cd sludge, and higher Cd sludge, respectively. Soil pH was adjusted to 5.7 by S addition. Some genotypes varied in response between years, while others did not (genetic studies are being conducted). In 1984, the geometric mean response for all genotypes on the untreated soil was 1.7 mg Cd/kg dry weight (range 1.2-2.5), 2.6 mg Cd/kg for the lower Cd sludge (range 1.4-5.5), and 5.6 mg Cd/kg for the higher Cd sludge (range 3.7-11.9). Thus, the field-determined variation in lettuce foliar Cd concentration was only about 3-fold.

Other less intensive studies with sludge-amended soils confirm this relatively narrow range (Giordano et al., 1979; Davis and Carlton-Smith, 1980 [Table 27]; Feder et al., 1980 [see CAST, 1980, Table 8] for leafy type lettuces. Head lettuce had approximately half the Cd concentration found in leafy lettuces grown in Salinas Valley (Burau, 1980) (Table 30). Chaney and Munns (1980, unpublished) tested the effect of sludge source and soil pH on Cd uptake by 6 lettuce cultivars. They also found cultivar differences in Cd uptake were small. Relative cultivar response was similar on lower and higher pH soils, but the range was much narrower on limed soils. Davies and Lewis (1985) and Crews and Davies (1985) compared trace element concentration in egible tissues of lettuce cultivars grown on metal rich soils contaminated with mine wastes in Great Britain. They found that relative cultivar response was similar on different soils, and the range of lettuce Cd concentration was about 2- to 3-fold on the several soils.

The Cd concentration in potato tubers did not vary significantly among 6 cultivars grown on a metal-rich soil at a long-term sludge utilization farm (Harris et al., 1981). The soil was pH 6.6 and contained 19.6 mg Cd/kg, while the mean Cd level for the washed unpeeled potato tubers was 0.28 mg Cd/kg dry weight [slightly greater than background Cd level in U.S. potatoes, 0.165 mg/kg dry weight (Wolnick et al., 1983)].

Meyer et al. (1982) found substantial differences among wheat types grown on U.S. scils containing background Cd levels. Durum type cultivars contained 0.140 mg Cd/kg dry grain, while soft red spring, soft red winter, hard red spring, and white wheat cultivars contained only 0.044 mg Cd/kg. Grain Cd was not significantly correlated with soil total Cd across all wheat cultivars, but was correlated if wheat types grown on similar soils were examined. Additional information has been provided by Hinesly (1986, Personal communication) on grain Cd in different cultivars of wheat grown on sludge-amended soil. 'Beau' grain contained 3.4 mg Cd/kg, while 'Argee' contained only 2.4 mg Cd/kg (strongly acidic soil, pH 5.5; 0.1 M HCl-extractable soil Cd about 33 mg/kg).

In a cooperative field trial on sludge metal availability, 4 barley cultivars from different regions of the U.S. were compared in a greenhouse experiment (Chang et al., 1982); no significant differences were found among the cultivars in Cd or Zn uptake to leaves or grain from plants grown on sludge-treated soil.

The effect of sewage sludge and carrot genotype on Cd accumulation in edible carrot roots was reported by Harrison (1986b). Two sludges were applied to 1 soil in 3 bed configurations. The mean Cd level was 0.38 mg Cd/kg for the control carrots, 0.50 mg Cd/kg for carrots grown on the lower Cd level sludge, and 0.77 mg Cd/kg for the higher Cd sludge. Cultivars differed less than 2-10ld in Cd accumulation. Differences among hybrid selections were significant for Cd, Zn, and other elements, although not all elements were increased due to sludge application.

Several methods have been used in different nations and at different times to estimate the maximum cumulative Cd application which protects the health of individuals (Dean and Suess, 1935). This is a very complex issue, as has been noted by Ryan et al. (1982) and Logan and Chaney (1983). The analysis given in Ryan et al. (1982) was considered when the US EPA proposed the existing regulations on land application of sludge in 1979 (Environmental Protection Agency, 1979a). A background document (Environmental Protection Agency, 1979b) reporting the scientific basis for the regulations was released at the time the interim final regulations were published.

Based on FDA dietary Cd intake estimates (36 µg Cd/day) and WHC/FAO recommendations for maximum tolerable weekly Cd intake (52-71 µg Cd/day), EPA (1979a) concluded that sludge could safely add no more than 30 uq Cd/day to an individual's diet. The high-risk or high-exposure individual was to be protected by the regulation: "That high-risk situation is one where an individual receives 50% of his vegetable diet from sludge-amended soils for a period of 40 to 50 years." The U.S. EPA recognized the strong effect of soil pH on Cd uptake by crops. For soils with low background pH, it was considered likely that soil pH would fall (from the pH 6.5 required during the permitted period of sludge application) to background soil pH. Thus, data from crops grown on acidic sludge-amended soils were used to estimate the relative Cd uptake by different food groups. The background document cites work by Dowdy and Larsen (1975), Giordano and Mays (1977), Chang et al. (1978), Chaney and Munns (1986, Personal communication), and a pot study by Furr et al. (1976). EPA calculated the increase above control, relative to that for lettuce, for each sludge application rate. The relative increases were averaged across rates; radish and carrot were averaged to obtain "root vegetables", and pea fruits and pea pods were averaged to obtain "legume vegetables." Lettuce represented "leafy vegetables" and tomato represented "garden fruits."

Table 31 summarizes the presumed relative increased Cd uptake by crops in the relevant FDA food classes, and daily food intakes for the teenage male diet model used by EPA in 1979 (Environmental Protection Agency, 1979b). If one multiplies food intakes (g dry/day, column B) times relative increased Cd uptake (column D), one obtains relative increased daily Cd intakes (column E). Thus, if lettuce is increased by 1 mg Cd/kg dry weight, garden foods are increased by 7.90  $\mu g$  Cd/day for 100%, or 3.95  $\mu g$ /day for 50% of garden foods grown in acidic sludge-amended garden for 40-50 years. EPA judged that strongly acidic soils (pH 5.4 to 6.2) would not cause greater than 30  $\mu g$  Cd increase/day, although very acid soils (4.9) caused larger increases.

Several groups evaluat d the 1979 regulations, and other research provided new information on and a better understanding of Cd transfer and

food consumption. By 1981, when U.S. EPA's Office of Solid Waste was preparing regulations under The Clean Water Act Section 405d, it was clear that average adult dietary intake data rather than teen-aged male dietary intakes should be used. Pennington (1983) provided an early draft of her results, and these were summarized into food groups by Flynn at EPA (1986, Personal communication). Leafy vegetables included lettuce, spinach, collards, cabbage, coleslaw, and sauerkraut. Potatoes included french fries, mashed, baked, boiled, scalloped, and sweet potatoes, and potato chips. Root vegetables included carrots, onions, beets, radishes, onion rings, mushrooms, and mixed vegetable. Legume vegetables included pinto, lima, navy, green (snap), and red beans, pork and beans, cowpeas, peas, peanuts, and peanut butter. Garden fruits included cucumber, pickles, tomatoes, tomato sauce and juice, catsup, cream tomato soup, squash, and vegetable soups; broccoli, celery, asparagus, and cauliflower were included here by Flynn because they have Cd response more similar to garden fruits than leafy vegetables, which had been classified in the 1979 teen-age male diet mcdel.

The 36  $\mu g$  Cd/day average intake (from FDA) was subtracted from the 71  $\mu g$ /day WHO/FAO limit yielding 35  $\mu g$  Cd/day allowed increase. Using the average adult rather than the teenage male dietary intakes reduced the predicted increase from 7.90 to 3.79  $\mu g$  Cd/day (for 100% of garden vegetable foods grown on acidic sludge amended soil) when lettuce is increased by 1  $\mu g$  Cd/day dry weight (Table 31). If one divides 35  $\mu g$  Cd/day by 3.79  $\mu g$  increased dietary Cd (per 1  $\mu g$  Cd increase/g dry lettuce), one finds leafy vegetables could safely reach 9.23  $\mu g$  Cd increase/g for 100% of diet; or 18.5  $\mu g$  Cd increase/g lettuce for 50% of diet; or 27.7  $\mu g$  Cd increase/g for 33% garden foods diet grown on acidic sludge-amended soils.

EPA then attempted to connect these leafy vegetable Cd increases to cumulative soil Cd loadings. Results for cumulative soil loading versus leafy vegetables were separated into acidic (pH 5.4-6.2) [Y(lettuce Cd) = 0.48X(kg Cd/ha) + 5.6; R<sup>2</sup> = 0.11] and very acidic (pH 5.3) [Y(lettuce Cd) = 8.1 X(kg Cd/ha) - 1.1; R<sup>2</sup> = 0.50]; each had low R<sup>2</sup>. They then tested the annual soil Cd loading results for acidic soils and got a better correlation [Y(lettuce(Cd) = 1.24X(kg Cd/ha) + 0.12; R<sup>2</sup> = 84]. Thus, using this equation, and 33% of garden foods, one calculates that soil could contain 22.2 kg added Cd/ha, or about 10 mg/kg.

The effect of soil pH on relative increased Cd uptake by garden crops is demonstrated by the data in Table 32. Strongly acidic soil pH causes much greater Cd uptake than near neutral soil pH, especially for the Cd-accumulating leafy vegetables. The relative increased Cd uptake is greater for carrot, potato, and peanut at the higher soil pH. Because high crop uptake at acidic soil pH is required for appreciable risk from nearly all sludges, relative increased Cd uptake for acidic soils should be used for risk assessment. However, this source of variation should be considered in evaluating different sources of data on relative Cd uptake by crops.

Relative increased Cd uptake by food groups from studies reported above are summarized in Table 33. Results varied among studies due to soil

Fe level. The 2 sludges differed strongly in solids content, and much more sludge adhered at 7.6% solids than at 2.0% solids. This study was done during the active growing season, and growth rapidly diluted the sludge content of forage. Although orchardgrass had slightly higher sludge adherence initially, either more rapid growth diluted adhering sludge on orchardgrass than on the tall fescue, or sludges did not adhere as long on orchardgrass.

In the second study, 1 sludge was applied at 2 rates to unclipped or clipped stands of 5 forage crop species, and forage samples were harvested at 6 times until after the normal harvest age of the clipped areas (Table 37). Statistical analysis showed that the harvest date was the most significant treatment variable, although the highest significant interaction term was clipping-x-species-x-harvest date. Alfalfa had the highest sludge content initially, but the lowest by the final harvest. Bluegrass had the second highest content initially, and retained the highest content during growth. Alfalfa's growth pattern is different from that of the grasses; the new growth of cut alfalfa comes from the highest axillary nodes, and all new growth emerges well above the thatch layer and contains no sludge. Treated parts of the grasses rise as new growth occurs at their base; new leaves emerge at the base and grow through the thatch layer. Thus, alfalfa and other dicotyledonous forage species are unlikely to allow much sludge transfer to the food chain under good management practices. Bluegrass forms very tight bunches, and sludge particles are trapped within the harvestable portion of the crop. These studies, taken together, indicate that present advice of "clip before spray application, avoid high solids content sludges, and wait for normal regrowth of the crop before harvest or grazing" continues to reflect research findings. Bluegrass is a species particularly inappropriate for spray application of sewage sludge. Application rate, although statistically significant, was the least important factor studied.

Although forages can reach 15-30% sludge (dry matter basis) immediately after application, Environmental Protection Agency regulations (1979a) now require a 30-day waiting period before grazing; users are advised to apply sludge to well grazed or clipped forages. This requirement reduces initial adherence and growth rapidly dilutes the adhering sludge. Decker et al. (1980) and Bertrand et al. (1981) found sludge comprised only 2-3% of the dry diet (based on forage and feces analyses) [Table 39] in practical grazing management. Injection of sludge in the plow layer soil prevents sludge adherence to forage crops, and greatly reduces potential ingestion of sludge from the soil surface.

Ingestion of sludge from the soil surface was estimated for a compost amendment which did not adhere to the crop (Decker et al., 1980). Cattle consumed about 1-3% compost when the forage had no detectable compost adhering (Table 39). Lower compost ingestion occurred in 1978 when compost was applied only once during the growing season. Others have evaluated soil consumption by well managed dairy cattle, sheep and swine (Fries et al. 1982, Fries and Marrow, 1982). Up to 8% soil was ingested from pasture, and less from bare soil. Hogue et al. (1984) reported metal

residues in sheep tissues after the sheep grazed 152 days on a grass-legume pasture established on soil in which 224 mt/ha metal-rich sludge had been incorporated. Although the forage was increased in Cd, kidney Cd of sheep was not increased. Other element residues were not influenced by sludge incorporation. Similar results were obtained when sheep grazed pastures which received surface-applied high Cu swine manure during the previous grazing season. Neither liver Cu nor fecal Cu was consistently affected by previous manure application (Poole et al., 1983).

Ingestion of sludge-borne trace elements does not necessarily cause the health effects which are expected based on traditional toxicological studies with added metal salts. Sludge feeding studies have been conducted to evaluate element deposition in tissues of cattle, sheep, and swine. Low metal concentration sludges have not increased Cd, Zn, Pb, etc. in animal tissues in several studies (Decker et al., 1980; Baxter et al., 1982; Evans et al., 1979); while high Cd sludges have increased Cd in liver and kidney (Bertrand et al., 1980; Fitzgerald, 1980; Johnson et al., 1981; Kienholz et al., 1979; Baxter et al., 1982; Hansen et al., 1981) (see review in Hansen and Chaney, 1983). The most consistent potential problem resulting from sludge ingestion is reduced Cu concentration in the liver. Sludge Zn. Cd. Fe. and possibly Mo could interfere with Cu absorption. Ingestion of sludge rich in Fe induced Cu-deficiency in cattle in the only sludge feeding or sludge grazing study where animal performance or health was negatively affected (Decker et al., 1980). When ingested sludges are rich in Cd, Hg, F, or Pb, deposition occurs in bone or liver, but little change has been found in livestock tissues used as food (Hansen and Chaney, 1983).

Crops grown on sludge-amended soils can transfer trace elements to feeds and foods. However, the extent of increases of trace elements in crop tissues, and the bioavailability of these to animals varies with sludge properties. Crops grown on soils amended with low metal sludges had little effect on kidney Cd in several studies; however, high Cd sludges increased Cd in crops, which increased Cd in kidney and liver (Decker et al., 1980; Bertrand et al., 1980; Rundle et al., 1984; Miller and Boswell, 1979; Chaney et al. 1978a, 1978b; Boyd et al., 1982; Bray et al., 1985; Babish et al., 1979; Haschek et al., 1979; Heffron et al., 1980; Lisk et al., 1982; Telford et al., 1982, 1984; Williams et al., 1978).

Much of this work has focused on Cd, because Cd can be mobile in food chains. Humans and laboratory animals have been used to characterize Cd bioavailability. Problems have been identified with the experimental methods used in this research. The early studies (Rahola et al., 1973; Yamagata et al., 1975) measured retention after only a few days or weeks. McLellan et al. (1978) found that part of the diet Cd was absorbed by intestinal mucosal cells which were subsequently sloughed into the intestine and the diet Cd repeatedly recycled in intestinal cells for a prolonged time. This delayed excretion allowed true absorption long after the test diet was fed and other parts of the test meal excreted. Flanagan et al. (1978) found that Fe deficiency very strongly affected Cd retention. The Fe deficiency increased Cd absorption into the intestinal mucosal cells where it was largely trapped as Cd-metallothionein. Fox et al. (1984)

showed this aspect of Fe deficiency allows increase in true Cd absorption and movement to kidney long after the test diet residue is excreted. Shaikh and Smith (1980) were able to study subjects up to 800 days after the test dose (using 109Cd), and resolved whole body Cd into 3 Cd pools, now including the slowly excreted intestinal Cd turnover pool. The biological half-life of the slowest pool was 18 years to infinity rather than 100 days as previously reported for the shorter-term human studies. Again, Fe deficiency affected retention of dietary Cd, but the apparent retention was appreciably lower than in earlier studies. These tend to support the findings of Newton et al. (1984) and Snarma et al. (1983) that only low amounts of Cd are retained by humans ingesting Western-type diets.

# CONCLUSIONS

- 1. Conditions for valid assessment of relative increased crop concentration of an element due to sludge utilization are limited to long-term sludge amended soils, preferably 4 or more years after sludge is applied. Metal salts and metal salt-amended sludges do not provide valid data for assessment of food-chain element transfer.
- 2. Some trace elements can be increased in edible crop tissues when sewage sludges rich in the element are applied to acidic soils (Cd, Zn, Ni), or alkaline soil (Mo). Under these conditions which allow substantial increase of a trace element in crops (responsive conditions), the relative increase in element concentration among crop species are sufficiently consistent to be relied upon in dietary exposure modeling. Some variation in relative increased trace element concentration among crops may result from high soil organic matter, or from calcareous vs. acidic soil conditions. High organic matter and high soil pH both reduce element uptake (except for Mo and Se).
- 3. Except for corn inbreds, cultivar variation in element concentration has been found to be approximately 2- to 5-fold. Because of inclusion of various cultivars in the food supply, this variation would not significantly alter chronic exposure due to increased crop uptake of sludge applied elements. However, cultivar selection can be used to reduce food-chain transfer of elements from well managed sludge utilization farms.
- 4. If the FDA food groups are used in dietary Cd modeling, they should be adjusted for relative high and low Cd accumulating crop types (lettuce vs. cabbage; carrot vs. beet) within a food group. Food intakes should represent average adult intake for 50 years, not the maximum intakes of teen-age males. For evaluation of potential chronic Cd exposure from acidic sludge-amended garden soils, the adult food intakes reported by Pennington (1983) can be used, and the relative increases in Cd in crops or food groups summarized in this report. Increased Cd uptake by all garden foods can be integrated in terms of increased Cd uptake by a reference crop such as lettuce. Dietary Cd

increase can be predicted by the response of Cd concentration in lettuce grown in test soils (height of the plateau in lettuce Cd on sludge-amended soils above the control, as affected by sludge Cd concentration and other factors) times the integrated garden foods Cd intake factor. Thus, increase in dietary Cd due to growing 100% of consumed garden vegetables on sludge-amended acidic garden soils was estimated as 2.20  $\mu g$  Cd/d when lettuce is increased above background by 1 mg/kg dry lettuce.

- 5. Prediction of changes in kidney Cd due to increases in dietary Cd from foods grown in acidic sludge-amended gardens should consider effects of nutritional status and nutrients in the garden crops on Cd retention by humans.
- 6. Ingestion of sludge can allow exposure and/or risk which can be prevented by incorporation of sludge below the soil surface, or by tilling sludge into the soil. For some elements (e.g., Fe, F, Cu, Zn, Pb), this pathway may allow sufficient exposure to sludge-borne elements to cause risk, at least for element-rich sludges. Bioavailability of many elements in ingested sludge is very strongly influenced by concentration of the element and other elements present, sludge carbonate content, and sludge redox potential. For many elements which comprise potential risk if sludge is ingested, median quality sludges have not caused any problems with livestock at common exposure rates from surface-applied sludge.

TABLE 27. TRACE ELEMENT CONCENTRATION IN EDIBLE PLANT TISSUES, AND RELATIVE Cd CONCENTRATIONS IN EDIBLE TISSUES OF CROPS (DRY WEIGHT BASIS); GEOMETRIC MEANS FROM EACH OF 2 LONG-TERM SLUDGE-AMENDED SOILS) (DAVIS AND CARLTON-SMITH, 1980).

Crop	Cultivar	Cđ	Ni	Zn	Relative Cd Concentration
		mg/k	g dry we	ight	
Lettuce	Tom Thumb	8.1	1.2	95.	100.
Lettuce	Webbs	6.9	2.2	96.	85.
Lettuce	Paris White Romaine	6.2	4.0	82.	77.
Spinach	Bloomsdale	5.0	1.4	391.	62.
Kale	Maris Kestrel	1.3	6.0	105.	16.
Cabbage	Greyhound	0.97	5.7	105.	12.
Wheat	Spartacus	0.88	6.4	75.	11.
Mangold	Yellow Globe	0.74	2.2	131.	9.1
Turnip	Bruce	0.74	1.9	45.	9.1
Leek	Musselburgh	0.73	0.91	28.	9.0
Wheat	Sappo	0.62	4.9	75.	7.7
Turnip	Snowball Snowball	0.58	2.0	37.	7.2
Rape	Orpa1	0.54	8.2	54.	6.7
Onion	White Lisbon	0.52	0.81	40.	6.4
Beetroot	Detroit	0.41	2.3	103.	5.1
Tomato	Moneymaker	0.40	2.0	22.	4.9
Sugarbeet	Sharpes Klein Monobeet	0.35	2.1	130.	4.3
Bettroot	Boltardy	0.34	2.1	76.	4.2
Carrot	Standard Improved	0.33	1.5	38.	4.1
Radish	French Breakfast	0.33	3.1	48.	4.1
Barley	Julia	0.31	4.2	67.	3.8
Parsnip	Giant Exhibition	0.26	3.2	34.	3.2
Barley	Ark Royal	0.25	7.8	69.	3.1
Swede	Acme	0.24	1.2	28.	3.0
Potato	Desiree	0.20	0.66	20.	2.5
0at	Leander	0.18	7.2	53.	2.2
Squash	Zucchini	0.17	5.8	80.	2.1
Sweet corn	Golden Earley	0.16	0.37	41.	2.0
Sunflower	Tall Single	0.15	11.4	41.	1.9
Maize	Caldera	0.13	1.0	37.	1.6
French bear	ı Canadian Wonder	0.08	9.9	31.	1.0
Pea	Onward	0.05	4.1	63.	0.6

TABLE 28. RELATIVE INCREASED CD CONCENTRATION IN EDIBLE TISSUES OF CROPS GROWN ON LONG-TERM SLUDGE-AMENDED SOILS. MEAN CROP CD FROM TABLE 27 WAS CORRECTED FOR NORMAL BACKGROUND LEVELS OF CD IN CROPS. ALL INCREASED CD CONCENTRATIONS WERE DIVIDED BY 6.37, THE MEAN CORRECTED CD CONCENTRATION IN 3 CULTIVARS LETTUCE (BASED ON DATA FROM DAVIS AND CARLTON-SMITH, 1980).

Crop	Cultivar	Crop Cd	Back- ground Cd	Increased Crop Cd	Relative Increased Cd Uptake
		mg	Cd/kg dry	weight	
Lettuce	Tom Thumb	8.1	0.7	7.4	116.
Lettuce	Webbs	6.9	0.7	6.2	97.
Lettuce	Paris White Romaine	6.2	0.7	5.5	86.
Spinach	Bloomsdale	5.0	0.7	3.7	58.
Kale	Maris Kestrel	1.3	0.27	1.0	16.
Wheat	Spartacus	0.88	0.08	0.80	13.
Cabbage	Greyhound	0.97	0.27	0.60	9.4
Wheat	Sappo	0.62	0.08	0.54	8.5
Mangold	Yellow Globe	0.74	0.21	0.53	8.3
Turnip	Bruce	0.74	0.21	0.53	8.3
Leek	Musselburgh	0.73	0.27	0.46	7.2
Rape	Orpal	0.54	0.08	0.46	7.2
Turnip	Snowball Snowball	0.58	0.21	0.37	5.8
Onion	White Lisbon	0.52	0.21	0.31	4.9
Barley	Julia	0.31	0.08	0.23	3.6
Beetroot	Detroit	0.41	0.21	0.20	3.1
Barley	Ark Royal	0.25	0.08	0.17	2.7
Sugarbeet	Sharpes Klein Monobeet	0.35	0.21	0.14	2.2
Bettroot	Boltardy	0.34	0.21	0.13	2.0
Radish	French Breakfast	0.33	0.21	0.12	1.9
0at	Leander	0.18	0.08	0.10	1.6
Carrot	Standard Improved	0.33	0.25	0.08	1.3
Tomato	Moneymaker	0.40	0.32	0.08	1.3
Potato	Desiree	0.20	0.13	0.07	1.1
Sunflower	Tall Single	0.15	0.08	0.07	1.1
Squash	Zucchini	0.17	0.11	0.06	0.9
Parsnip	Giant Exhibition	0.26	0.21	0.05	0.8
Sweet corn	Golden Earley	0.16	0.11	0.05	0.8
Maize	Caldera	0.13	0.08	0.05	0.8
Swede	Acme	0.24	0.21	0.03	0.5
French bean	Canadian Wonder	0.08	0.06	0.02	0.3
Pea	Onward	0.05	0.06	-0.01	-0.2

Background Cd concentrations were based on field grown control crops reported in many studies, but mainly Wolnik et al. (1983, 1985); Korcak (1986, Personal communication) summarized these results in a draft report to EPA.

TABLE 29. RELATIVE UPTAKE OF TRACE ELEMENTS TO TISSUES OF FORAGE CROPS (DRY WEIGHT BASIS) (CARLTON-SMITH AND DAVIS, 1983).

1 m - 1 m - 1 m - 1 m

	4 4	% of	crop w		ghest	concentration
Crop .	Cultivar	Cd	Cu	Mo	Li	Zn
Agrostis tenuis	'Goginan'	15	49	38	73	36
Agrostis tenuis	'Parys'	12	55	50	85	44
Dactylis glomerata	<b>'</b> \$26'	39	100		57	17
Dactylis glomerata	'S37'	15	57		33	10
Dactylis glomerata	'S143'	12	47		30	11
Festuca arundinacea	'S170'	52	76	23	63	32
Festuca pratensis	'S215'	37	78		91	32
Festuca rubra	'Merlin'	10	51		34	19
Festuca rubra	'S59'	29	51		47	22
Lolium multiflorum	'Aubade'	15	<b>6</b> 8	29	44	24
Lolium multiflorum	'RVP'	17	72	28	55	25
Lolium multiflorum	'S22'	15	73	29	46	26
Lolium multiflorum	'Sabalan'	15	67	25	59	21
Lolium multiflorum	'Sabrina'	11	89	29	60	26
Lolium perenne	'Cropper'	16	86	37	80	31
Lolium perenne	'Melle'	15	86	27	92	27
Lolium perenne	'S23'	14	86	33	68	30
Lolium perenne	'S24'	14	90	29	100	30
Lolium perenne	'S321'	19	90	29	80	30
Lolium perenne	'Talbot'	14	89	35	82	32
Phleum pratense	'\$48'	31	96	25	73	30
Phleum pratense	'S51'	31	95	24	73	28
Phleum pratense	'S352'	32	88	21	80	33
Avena sativa	'Trafalgar'	9	29	16	42	8
Hordeum sativa	'Julia'	9	48	16	16	15
Triticum aestivum	'Sappo'	7	38	16	17	10
	'Caldera'	13	30		16	
Zea mays	'Maris Carmine'	32	30	11 12	16	10 14
Zea mays Medicaco sativa	'Europe'	14	35	34	31	
Medicago sativa	'Hungarapoly'	7	53	54 62	31 44	11 14
Trifolium pratense Trifolium pratense	'S123'	7	53 57	62 62	44	16
	'Kent Wild White'	8	43		34	18
Trifolium repens	'S100'	5		100	34 36	
Trifolium repens	'S184'	5	51 20	79	36 40	20
Trifolium repens		16	38	71 41		21
Brassica oleracea	'Maris Kestrel'		21	41	30	13
Brassica rapa	'The Bruce'	50	35	38	33	19
Beta vulgaris	'Sharpes Monobeet'		66	19	57 57	83
Beta vulgaris	'Yellow Globe'	95 	81	26	57	100
Highest concentrati						
(mg/kg dry weig	ht)	1.41	15	14	1.4	417

Concentration in plant shoots normalized across control and 2 sludge treatments. For the crop with the maximum normalized concentration, (100) = the listed mg element/kg dry matter.

TABLE 30. RELATIVE Cd CONCENTRATION IN CROPS GROWN ON NATURALLY Cd RICH SALINAS VALLEY SOILS (BURAU, 1980).\*

Crop	mg Cd/kg wet weight mg Cd/kg dry soil		mg Cd/kg dry weight mg Cd/kg soil	Relative uptake
Spinach	0.70	9.3	7.5	280
Endive	0.24	6.9	3.5	130
Lettuce. leaf	0.16	6.0	2.7	100
Lettuce, romai	ine 0.16	6.0	2.7	100
Lettuce, head	0.07	4.5	1.6	59
Chili pepper	0.10	8.0	1.2	44
Carrots	0.13	11.8	1.1	41
Artichokes	0.14	13.5	1.0	37
Potatoes	0.09	20.2	0.45	17
Garlic	0.17	38.7	0.44	16
Sweet corn	0.11	27.3	0.40	15
Cucumber	0.02	4.9	0.41	15
Squash, zucchi	ini 0.02	5.4	0.37	14
Red beets	0.04	12.7	0.31	11
Onions	0.03	10.9	0.28	10
Cauliflower	0.02	9.0	0.22	8
Parsley	0.03	14.9	0.20	7
Tomatoes	0.01	6.5	0.15	8 7 6 3 3
Broccoli	0.01	10.9	0.092	3
Beans, white	0.08	89.1	0.090	3

<sup>\*</sup>Soils were between pH 6-8 and contained 1-10 mg Cd/kg, and about 1.5% organic matter.

tFrom Watt and Merrill (1963).

TABLE 31. CADMIUM EXPOSURE MODEL FROM THE 1979 ENVIRONMENTAL PRCTECTION AGENCY SLUDGE APPLICATION REGULATION AND BACKGROUND DOCUMENT (EPA, 1979a, 1979b), AND THE 1981 DRAFT BACKGROUND DOCUMENT. TABLE SHOWS INTAKES OF FDA FOOD CLASSES BY THE HYPOTHETICAL TEENAGED MALE DIET MODEL (1979) OR AVERAGE ADULT DIET MODEL (1981), AND RELATIVE Cd UPTAKE BY FOOD GROUPS (EPA, 1979b).

	A	В	С	D	E = (B x D)	
Food Group	Food Intake		H <sub>2</sub> 0	Relative Increased Cd Uptake	Relative Daily Cd	
	g wet/day	g dry/day	%		μg C/day	
1979 Diet Model						
Leafy vegetables	55	4.95	91	1.00	4.95	
Potatoes	183	43.9	76	0.02	0.88	
Root vegetables	33	2.64	92	0.23	0.61	
Legume vegetables	69	13.1	81	0.04	0.52	
Garden fruits	69	5.52	92	0.17	0.94 7.90	

If 100% of garden foods diet were grown on acidic sludged land, diet would be increased 7.90  $\mu g$  Cd/day when lettuce increased by 1  $\mu g$  Cd/g dry weight. If 50% of garden vegetables, diet increases by 3.95  $\mu g$ /day when lettuce increases 1  $\mu g$  Cd/g dry weight.

1981 Diet Model					
Leafy vegetables	26	2.34	9	1.00	2.34
Potatoes	64	15.36	24	0.02	0.307
Root vegetables	13	1.04	8	0.23	0.239
Legume vegetables	38	7.22	19	0.04	0.289
Garden fruits	60	3.60	6	0.17	<u>0.612</u> 3.79

If 100% of garden foods diet were grown on acidic sludged land, diet would be increased 3.79  $\mu g$  Cd/day when leafy vegetables increased by 1  $\mu g$  Cd/g dry weight. If 50%, diet increases by 1.90  $\mu g$ /day; and if 33%, 1.26  $\mu g$  Cd/day.

TABLE 32. EFFECT OF SOIL pH ON RELATIVE INCREASE ABOVE CONTROL OF Cd IN EDIBLE CROP TISSUES (CHANEY, 1985. PERSONAL COMMUNICATION).

	Increase	Increased Crop Cd		Relative Increase		
Crop	Acidic	Limed	Acidic	Limed		
<del></del>	mg Cd/	kg dry				
Lettuce	29.3	5.73	1.00	1.00		
Carrots	2.15	1.48	0.073	0.26		
Potatoes	1.17	1.02	0.040	0.18		
Peanuts	0.54	0.41	0.018	0.072		

Sludge containing 210 mg Cd/kg applied at 50 and 100 mt/ha in summer, 1978. Carrot and lettuce results from 1979; potato and peanut results from 1980.

TABLE 33. COMPARISON OF RELATIVE INCREASED Cd UPTAKE BY FOOD GROUPS BASED ON DIFFERENT DATA SOURCES SUMMARIZED ABOVE.

	Reference <sup>†</sup>					
Food Group	EPA	Davis	Dowdy Larson	Giordano et al.	Burau	Chaney Acidic
	1979	(2)	1975	1979	(4)	(6)
Leafy Vegetables (lettuce)	1.00	1.00	1.00	1.00	1.00	1.00
Potato	0.02	0.020	0.052	0.00	0.17	0.040
Root Vegetables	0.23	0.07	0.36	0.37	0.21	0.073
Legume Vegetables	0.04	0.01	0.022	0.017	0.03	0.014
Garden Fruits	0.17	0.020	0.15	0.18	0.12	

<sup>†</sup>Numbers in parentheses indicate table in text where data are contained.

TABLE 34. AVERAGE ADULT DAILY INTAKES OF FOODS AGGREGATED INTO FOOD GROUPS ON WET WEIGHT AND DRY WEIGHT BASIS. AVERAGE WET G/DAY FOOD INTAKES OBTAINED FROM PENNINGTON (1983); SIX MALE AND FEMALE DIETS, FOR AGES 14-65, WERE AVERAGED. CONVERTED TO DRY WEIGHT USING DATA FROM ADAMS (1975).

<u>Pennington</u>		Adams Data	Adult Food	
Food	g wet/day	% dry wt.	g dry/day	% dry
Leafy Vegetables	- High Cd Uptake	<u>:</u>		
Lettuce	19.231	4.8	0.865	
Spinach	0.816	8.6	0.070	
Spinach	2.329	8.0	0.196	
op maon	22.376	3.0	1.189	5.3
Leafy Vegetables	- Low Cd Uptake:			
Collards	1.715	10.4	0.178	
Cabbage	2.849	6.1	0.174	
Coleslaw	2.530	19.4	0.491	
Sauerkraut	0.939	7.2	0.068	
Broccoli	2.403	8.7	0.209	
Celery	0.922	5.9	0.054	
Asparagus	0.836	6.4	0.054	
Cauliflower	0.772	7.2	0.056	
	12.966		1.284	9.9
Potatoes:				
French fries	20.026	55.3	11.074	
Mashed	16.232	20.7	3.360	
Boiled	12.202	20.2	2.465	
Baked	6.859	24.9	1.708	
Chips	2.963	98.2	2.901	
Scalloped	5.941	28.9	1.717	
Sweet	1.541	36.3	0.559	
Sweet	0.674	40.0	0.270	
	66.438		24.054	36.2
Root Vegetables -				
Carrots	3.401	17.8	0.605	17.8
Root Vegetables -				
Onions	2.473	10.9	0.270	
Mixed veg.	5.154	17.4	0.897	
Mushroom	0.787	9.6	0.076	
Redbeets Radish	1.069	10.7	0.114	
Onion rings	0.402	5.5	0.022	
union rings	0.710	8.2	0.058	
	10.595		1.437	13.6

(continued)

TABLE 34 (continued)

Pennington Data		Adams Data	Adult Foo	d Intake
Food	g wet/d	% dry wt.	g dry/day	% dry
egume Vegetables:				
Snap beans	2.924	7.6	0.222	
Snap beans	6.172	8.1	0.500	
Cowpeas	2.402	28.2	0.677	
Lima beans	0.870	35.9	0.313	
Lima beans	1.301	26.5	0.345	
Navy beans	1.567	31.0	0.486	
Red beans	1.748	31.0	0.542	
Peas	<b>5.5</b> 68	23.0	1.281	
Peas	1.533	17.9	0.274	
Peanut butter	2.269	98.3	2.230	
Peanuts	0.896	98.2	0.880	
Pinto beans	7.589	31.0	2.353	
Pork & beans	7.552	33.6	2.537	
			12.640	29.8
			12.00	23.0
Garden Fruits - Hi				
Tomatoes	16.186	5.5	1.053	
Tomato juice	3.658	6.4	0.234	
Tomato sauce	2.108	13.0	0.274	
Tomatoes	0.885	6.3	0.056	
Tomato soup	9.464	9.5	0.899	
Catsup	2.396	31.2	0.748	
Green pepper	0.830	6.6	0.055	
			3.319	9.3
Garden Fruits - Lo				
Cucumber	2.735	4.3	0.118	
Pickles	1.324	6.7	0.089	
Summer squash		4.5	0.099	
Winter squash	1.207	11.2	0.135	
Watermelon	4.748	7.4	0.351	
Canteloupe	4.416	8.8	0.389	
Strawberries	2.508	10.1	0.253	
Sweet corn	6.574	25.9 25.1	1.703 0.958	
Sweet corn	3.816	25.1	0.576	
Sweet corn	2.431	23.7	0.5/0	
	31.949		4.671	14.6

TABLE 35. FOOD GROUP AGGREGATION OF FOOD INTAKE RESULTS FROM PENNINGTON (1983). DATA FOR SIX AGE-X-SEX GROUPS (AGES 14-65) WERE AVERAGED; WET WEIGHT CONVERSION TO DRY WEIGHT CONDUCTED ON INDIVIDUAL FOOD BASIS USING DATA FROM ADAMS (1975). FOODS FROM THE PENNINGTON LISTS WERE THE SAME AS LISTED BY FLYNN EXCEPT STALK VEGETABLES WERE MOVED TO LEAFY VEGETABLES - LOW CATEGORY, AND SWEET CORN AND MELONS WERE ADDED TO GARDEN FRUITS - LOW.

Food Group <sup>†</sup>	Food wet g/d	Intakes dry g/d	% Dry Weight	Relative Increased Cd Uptake	Relative Increased Cd Intake
Leafy Vegetables-High Leafy Vegetables-Low Potatoes Root Vegetables-High Root Vegetables-Low Legume Vegetables Garden Fruits-High Garden Fruits-Low	22.376 12.966 66.438 3.40 10.60 42.39 35.537 31.949	1.121 1.284 24.063 0.605 1.437 12.640 3.319 4.671	5.0 9.9 36.2 17.8 13.6 29.8 9.3 14.6	1.00 0.13 0.020 0.20 0.052 0.010 0.020 0.010	1.121 0.167 0.481 0.121 0.075 0.126 0.066 0.047
					2.20

<sup>†</sup>Leafy Vegetables - High includes lettuce and spinach.

Leafy Vegetables - Low includes cabbage, kale, broccoli, etc.

Root Vegetables - High includes carrots.

Root Vegetables - Low includes radish, turnip, beet, onion, and leek.

Garden Fruits - High includes tomato products and pepper.

Garden Fruits - Low includes cucurbits, sweet corn, and strawberries.

TABLE 36. COMPARISON OF FOOD INTAKES, RELATIVE INCREASED Cd UPTAKE, AND ESTIMATED INCREASED DIETARY Cd IN THE EPA (1979b), 1981 EPA DRAFT, AND PRESENT DOCUMENT.

Food Group	EPA, 1979	EPA, 1981	Present
		g dry weight/day	
Food intakes:			
Leafy vegetables	4.95	2.34	2.405
Potato	43,9	15.36	24.063
Root vegetables	2.64	1.04	2.042
Legume Vegetables	13.1	7.22	12.640
Garden fruits	5.52	3.69	7.990
Relative food group in	ncreased Cd uptake	e (lettuce = 1.00)	<u>)</u> :
Leafy vegetables	1.00	1.00	0.536
Potato	0.02	0.02	0.02
Root vegetables	0.23	0.23	0.096
Legume Vegetables	0.04	0.04	0.010
Garden fruits	0.17	0.17	0.014
Estimated relative foo	od group increased	d Cd intake:	
Leafy vegetables	4.95	2.34	1.288
Potato	0.88	0.307	0.481
Root vegetables	0.61	0.239	0.196
Legume Vegetables	0.52	0.289	0.126
Garden fruits	0.94	0.612	0.113
Increased dietary Cd	(ug/day) if lettu	ce increased 1 mg	Cd/kg dry:
	7.90	3.79	2.20

Present estimate differs from previous estimates by separating leafy, root, and legume vegetable and garden fruits into high and low Cd-accumulating crop sub-groups. The overall food group relative increased dietary Cd intake was calculated using the increased food and Cd intake from high and low sub-groups (from Table 35).

TABLE 37. EFFECT OF SLUDGE SOURCE, AND TIME AFTER SLUDGE APPLICATION ON SLUDGE ADHERENCE TO TALL FESCUE AND ORCHARDGRASS (CHANEY AND LLOYD, 1986, PERSONAL COMMUNICATION).

			Days	After Slu	Sludge Applicati	
Sludge % Solids Cro	Crop	0	7	14	20	
<del></del>	<u></u>		%	sludge in/	on forages	
City 1	2.0	Tall Fescue Orchardgrass	3.0 7.4	3.1 4.4	1.5 2.8	0.82 0.90
City 23	7.6	Tall Fescue Orchardgrass	10.2 11.9	6.2 6.9	3.1 2.1	2.7 1.0

Forages were not clipped; sludge applied at 94 m<sup>3</sup>/ha using watering cans. Sludge content calculated based on increased levels of 6 elements above levels present in unsprayed control forage samples.

Anaerobically digested sludge from City 1 contained (in mg/kg dry solids): Zn, 3030; Cd, 549; Pb, 495; Cu, 665; Ni, 68; Fe, 11,000.

Anaerobically digested sludge from City 23 contained (in mg/kg dry solids): 2n, 750; Cd, 7.2; Pb, 170; Cu, 195; Ni, 27; Fe, 123,000.

TABLE 38. EFFECT OF FORAGE CROP SPECIES, CLIPPING CROP BEFORE SLUDGE APPLICATION, AND TIME AFTER APPLICATION ON ADHERENCE OF SPRAY-APPLIED FLUID SLUDGES TO FIVE FORAGE CROP SPECIES (CHANEY AND LLOYD, 1986, PERSONAL COMMUNICATION).

Crop	Harvest Date	Unclipped	Clipped
		% sludge	in/on forage
Tall Fescue	0	3.69 d-i	4.44 cd
'Kentucky 31'	7	3.02 f-l	3.77 c-h
	14	2.79 h-n	2.61 k-o
	28	1.89 m-q	1.19 q-v
	43	2.85 h-m	0.53 u-x
	70	1.01 q-w	0.30 vwx
Orchard grass	0	2.68 i-o	4.53 cd
'Potomac'	7	2.27 1-p	3.67 d-i
•	14	2.34 k-p	1.67 o-s
	28	1.61 p-t	0.60 t-x
	43	1.32 p-v	0.40 vwx
	70	1.11 q-x	0.13 wx
Kentucky bluegrass	0	6.36 b	5.68 b
'Merion'	7	4.18 cde	3.99 c-f
•	14	4.73 c	3.33 e-k
	28	4.37 cd	2.97 g-1
	43	2.64 j-o	1.32 p-v
	70	1.81 n-q	0.52 u-x
Smooth bromegrass	0	3.63 d-j	4.26 cde
'Saratoga'	7	3.29 e-1	4.31 cde
	14	2.75 h-n	3.97 c-g
	28	1.45 p-u	1.70 o-r
	43	1.56 p-t	0.62 t-x
	70	0.77 r-x	0.45 u-x

(continued)

TABLE 38 (continued)

Crop	Harvest Date	Unclipped	Clipped
		% sludge	in/on forage
Alfalfa	0	8.48 a	5.98 b
'Saranac'	7	6.17 b	<b>4.</b> 30 cde
	14	3.67 d-i	1.82 n-q
	28	1.21 q-v	0.51 u-x
	43	0.66 s-x	0.13 wx
	70	0.09 wx	-0.05 x

Forages were established in spring, 1976 on methyl-bromide treated field plots. After establishment, sludge was applied on May 11, 1977, at 51 and 103  $\rm m^3/ha$ , to unclipped and clipped (to 10-15 cm as recommended for species, with clippings removed) crops in three replications. Forage was harvested to 5 cm after 0-70 days growth. Normal harvest of the clipped forage would have occurred about day 43.

Sludge was 1.4% solids and contained (in mg/kg dry solids): Zn, 1140; Cu, 432; Pb, 394; and Fe, 36,000. Sludge content was estimated by increased levels of Zn, Cu, Pb, and Fe in sprayed forage. The speciesx-clipping-x-harvest date was the highest significant interaction in ANOVA. Sludge content results followed by the same letter were not significantly different (at  $\Gamma < 0.05$ ) according to the Duncan Multiple Range Test.

TABLE 39. Adherence of spray-applied liquid sewage sludge to tall fescue (Decker et al., 1980) or 'Pensicola' bahaigrass (Bertand et al., 1981) and sludge content of feces of cattle which rotationally graze these pastures.

Study and Treatment	Sludge Solids	Appli- cation Rate	S <sup>l</sup> udge in/on Forage	Sludge in Feces
	*	cm	%	%
Decker et al. (1980)*				
1976 - 21-day sludge 1976 - 1-day sludge	4.4 4.8	20 x 0.51 20 x 0.51	5.39 22.3	7.1 18.6
1977 - 21-day sludge 1977 - compost	2.9	20 x 0.51	2.18 (0.74) <sup>†</sup>	7.7 6.5
1978 - 21-day sludge 1978 - compost	3.7	24 x 0.51	2.91 (0.50) <sup>†</sup>	6.1 2.0
Bertrand et al. (1981)‡				
1979 - 7-26-day sludge 1979 - 7-13-day sludge	2.1	9 x 0.84 18 x 0.84	2.17 5.17	4.6 5.8

<sup>\*</sup>Four paddocks grazed on a rotation system; sludge was applied to clipped pasture 21 days before grazing (21-day sludge), or regrown pastures 1 day before grazing (1-day sludge). Compost applied 3 times in 1977 and 1 time in 1978, with at least 21 days before grazing began.

<sup>†</sup>Estimates based on individual elements were not in close agreement; no significant sludge content.

<sup>\*</sup>Data of Bertrand et al. (1981) recalculated using results for Cu, Fe, Pb, and Zn, elements substantially increased by sludge application. Two paddocks were grazed in rotation. Rotations were made every 12-14 days; depending on forage growth and weather, the sludge application occurred 7 to 13 days before grazing commenced. The two sludge treatments differed in number of sludge applications made during the grazing season.

## SECTION 6

# EFFECTS OF TRACE ORGANICS IN SEWAGE SLUDGES ON SOIL-PLANT SYSTEMS AND ASSESSING THEIR RISK TO HUMANS

## INTRODUCTION

Describing the impact of trace organics in sludge on soil-plant systems can be an even greater challenge than is faced with trace elements. One reason is the sheer number of compounds potentially involved. Literally thousands of trace organics exist and many, if not all, can be expected in sewage sludge at highly variable concentrations. At the same time, the literature discussing the effects of trace organics on soil-plant systems is much less voluminous than the trace element literature. Undoubtedly, the paucity of scientific studies on trace organics is due to the complexity of studying these chemicals and the expense of trace organic analyses.

An important difference between trace metal and trace organic additions to a soil is the time each may reside or persist in that soil. The half-life of the most persistent organics (e.g., PCBs) in soil was concluded to be 10 or more years (Fries, 1982), whereas the residence time for most metals was estimated to be a few thousand years (Bowen, 1977). Studies of trace organic behavior in soils must also consider assimilation mechanisms such as degradation (biotic and abiotic) and volatilization, in addition to factors such as solubility, adsorption/desorption, leaching and plant uptake. While these additional mechanisms make trace organic studies more challenging, they also lend themselves to management alternatives not available for trace elements. For example, long-term application programs with organics (e.g., food processing and petroleum wastes) attest to the soil's ability to receive and successfully assimilate wastes over time.

This attenuation capacity suggests that limiting the addition of trace organics to soils via sludge application should be based on "matching the total loadings of an organic(s) with the soil's assimilative ability" rather than "a specified concentration present in the sludge". Such an approach allows more flexibility to consider environmentally sound options on a case by case basis for local circumstances.

Some analogies of trace organics to agricultural pesticides can be made. But the influence of the sludge organic matter matrix in combination with a specific organic, when added to the soil, is poorly understood.

To facilitate discussion, the large number of organic chemicals were divided into groups which tend to have similar chemical and physical properties. Various organics could then be discussed by groups relevant to their prevalence in sludge, fates in the soil-plant system, and recent efforts at assessing the risk of trace organic additions to the soil via sludge application.

The following discussion focuses on the impact of trace organics to soil-plant systems. The toxicity of trace organics to soil organisms, animals or humans as a result of their addition to soils and potential pathways, whereby exposure of soil-applied organics to animals and humans might occur, are listed and briefly discussed.

### PREVALENCE OF TRACE ORGANICS IN SLUDGES

Any program to assess the risk from trace organics must begin by determining (1) which chemicals are the most likely to be present in sewage sludge and (2) what quantities may be added to soils by the application of sludges containing these trace organics. To assess the potential impacts, priority should be given to substances shown to be prevalent in sewage sludge through residue analysis and which have certain physical-chemical properties that could lead to unacceptable toxicological or environmental effects. The next priority are those chemicals heavily used in society, possessing similar undesirable physical-chemical properties but not yet identified in sludges. With the current data base only the first priority can be effectively considered.

Since municipal raw sewage contains virtually all the wastes from man's activities, one could expect the sludge resulting from the treatment of this sewage to also contain these same products. Because as many as 15,000-20,000 man-made chemicals (with an array of functional groups) exist, analyzing all the chemical constituents in a sewage sludge is impossible. Sludges have been analyzed according to predetermined lists of specific organic chemicals such as the organic priority pollutants list (NRDC, 1976). Another approach is to separate the organics into "chemical groups" which have similar physical-chemical properties and focus on selected groups anticipated to have a greater toxicological and/or environmental risk. With either approach, the wastewater treatment plant should first determine what organics are being discharged by users, particularly industry. This information can then be used as guidance for identifying which organic compounds should be tested to check for any unusually high concentrations in sewage sludges.

A third approach is to use short-term bioassays to test sludges or sludge-amended soil for mutagenicity (Brown et al., 1982; Hopke and Plewa,

1984; and Peters, 1985). Sludges failing such a test would then be evaluated more rigorously and analyzed for selected organics. This approach requires additional research on suitable bioassays followed by calibration of these bioassays with experiences in the field, before it could realistically be used.

Only a few studies have reported the analyses of trace organics in sewage sludges. These studies confirm the wide variety of trace organic compounds that can occur in sewage sludge, but significant problems exist in the analysis and interpretation of these data:

- Sludges are heterogenous and obtaining a representative sample can be difficult.
- · Day-to-day variations in composition occur.
- Analytical protocols vary widely in extractions, separations and cleanup procedures which in turn affect the number and types of compounds recovered.
- For some groups, recoveries from a complex matrix like sludge can be poor.
- Data are reported in various units ( $\mu g/1$ , mg/kg, etc., some on a wet weight basis, and others on dry weight).
- · Limits of detection in some cases are poor or are not reported.
- · Confirmation of each organic, if any was done, is not reported.

Because of these problems and a very limited data base, definitive statements concerning the prevalence of organic chemicals in sewage sludge can not be made. To rectify this situation, the following information and data are suggested as a minimum for reporting on the organic content of sewage sludge: (a) type of sludge, (b) percent dry solids, (c) number of samples analyzed, (d) number of "positive" samples above detection limit, and the following based on dry weight --maximum and minimum concentration (range), (e) detection limit, and (f) the median concentration of all samples tested. This information would provide a means of standardization for comparing data sets.

Residual levels of trace organic compounds found in sewage sludge analysis surveys are listed in Table 40. The majority of these data come from two sources (Burns and Roe, 1982; and Jacobs and Zabik, 1983) but are supplemented by several others. Studies reporting organic concentrations for fewer than 9 sludges were not included in this summary, except for one which provided data for dioxins and furans (Weerasinghe et al., 1985). Some limitations of the data reviewed (Table 40) are that detection limits were not reported, some data were reported on a wet weight basis without "% solids" values given, and median concentrations for all samples were not provided.

Compounds were listed under the following major groups based on similar physical-chemical characteristics:

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    phthalate esters
    monocyclic aromatics
    polynuclear aromatics (PAH's)
    halogenated aliphatics (short chain)
    triaryl phosphate esters
    aromatic and alkyl amines
    phenols
    dioxins and furans
    miscellaneous
    compounds
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These data show that sewage sludges can be highly contaminated with organic chemicals. Unusually high concentrations, such as the maximum levels shown for butylbenzylphthalate, bis(2-ethylhexyl) phthalate, toluene, methyl bromide, chloroethane, vinyl chloride, pentachlorophenol and others, suggest a high degree of industrial contamination. Sludges containing these "maximum concentrations" could have a significant impact on soil-plant systems, depending on the rate of sludge applied.

Concerns about organic chemicals in sludges must be kept in perspective, however. Of the 219 organic chemicals collectively measured in sludges, 70 (or 32%) were below detection limits (Table 41). About one-fourth (53) of these organics were present in >50% of the sludges (Table 41). The presence of "background concentrations" of many organics in purely domestic sewage sludges is not unexpected, given the wide variety of synthetic organic chemicals found in many household products (Hathaway, 1980). The fact that domestic septic tank effluents contain greater than 100 trace level organics provides additional evidence for their presence in household wastewaters (DeWalle et al., 1985; Tomson et al., 1984).

More important than the presence of an organic(s) in sewage sludge is the total amount which may get applied to the soil-plant system by application to land. Table 42 summarizes that part of the analysis data from Table 40 which had median concentration values. This summary suggests that about 90% of the organics in sludges will be present at concentrations less than 10 mg/kg. About 10% of the organics tested had median concentrations of 10-100 mg/kg, and only one organic had a median value of >100 mg/kg (Table 42).

To put potential organic chemical loadings into perspective, one can make a comparison with agricultural pesticides. Many pesticides used today are organic chemicals which are added to soil-plant systems at rates of 0.2-4.0 kg of active ingredient per hectare. Assuming an agronomic rate of sludge application of 10 mt/ha (dry weight basis) the organic chemical loadings expected for organic concentrations in sludges of 1, 10, and 100 mg/kg are 0.01, 0.1 and 1.0 kg/ha. At rates used to reclaim drastically disturbed land, 100 mt/ha, the organic loading for sludges containing 1, 10, and 100 mg/kg organic concentration would be 0.1, 1.0, and 10 kg/ha, respectively. For agronomic rates organic chemical concentrations of sludges approaching 100 mg/kg must be viewed as potentially having an impact on the soil-plant system, depending on the chemical/toxicological properties of that organic. At high sludge rates (e.g.,

100 mt/ha), concentrations approaching 10 mg/kg in sludge could be expected to add amounts of an organic comparable to quantities of pesticides added in agricultural operations.

Based on the prevalence of organics in sludges and potential loadings to soils, agronomic or environmental risk due to the application of domestic sewage sludge to agricultural soils appears to be minimal. In addition, many organics will be bound by soil organic matter and biologically degraded by soil microorganisms (Kaufman, 1983). However, persistent compounds like PCBs and the chlorinated pesticides could accumulate in soils from repeated sludge applications and can be a concern for food crop production.

## TRACE ORGANICS IN SOILS

Limited information is available regarding residual effects of sludge organics in soils. Monitoring for 22 persistent organics in unamended and sludge-amended soils (Baxter et al., 1983) showed trace levels of chlordane (<0.12 mg/kg), dieldrin, p,p'-DDE and PCBs present in untreated and treated soils. None of the other 22 organics were detected in any of the soil samples. The authors concluded that sludge applications had not measurably increased the level of persistent organics above the levels normally found.

In a Michigan study (Singh, 1983), sludge-treated and untreated soils were collected from 15 sites around the state and analyzed for 10 organic compounds: benzene, trichloroethylene, tetrachloromethane, PCBs, pentachloronitrobenzene, pentochlorophenol, chlorpyrifos, di-n-butylphthalate, bis(2-ethylhexyl) phthalate, and toxaphene. All results for soil analyses were reported as "none detected" except at two sites PCBs were found at 0.8 mg/kg for untreated and sludge-treated soils at one site, and 0.02 mg/kg pentachlorophenol was detected in sludge-treated but not in control soils at another site.

While these two studies suggest that sludge organic chemical loadings to soils will result in little or no residues in soils receiving sludges, additions of persistent organics can potentially be a concern. Two food processing companies were contacted to determine what level(s) of organic residues in soils they use to reject fields for use in growing vegetable crops. One company indicated that concentrations above 0.1 mg/kg of aldrin/dieldrin, chlordane, toxaphene, or lindane in mineral soils would be of concern. For muck soils, 1.6 mg/kg of aldrin/dieldrin or 0.5 mg/kg of chlordane, toxaphene or lindane could be tolerated. A second vegetable crop processing company provided the guideline information in Table 43.

To assess the potential impact of sludge organic loadings to agricultural soils, the theoretical residue levels can be determined. Using the highest median concentration for aldrin or dieldrin from Table 40 of about 1 mg/kg and assuming an agronomic rate of sludge is applied (10 mt/ha) for 10 years, the total amount of aldrin or dieldrin added to a soil

would be: 0.01 kg/ha x 10 (yr) = 0.1 kg/ha. To determine what the soil residue concentration would be, one can assume an average bulk density for soil of 1.3 g/cm<sup>3</sup> (1,300 kg/m<sup>3</sup>) and a 20 cm depth of mixing, so one hectare (10,000 m<sup>2</sup>) of soil would weigh 2,600,000 kg (10,000 m<sup>2</sup> x 0.2 m x 1,300 kg/m<sup>3</sup>). Assuming no loss of the organic chemical applied, the soil residue concentration would be: 0.1 kg (or 100,000 mg) of organic/ha  $\pm$  2,600,000 kg of soil/ha =  $\sim$ 0.04 mg/kg.

Under these conditions, some margin of safety would still be provided relative to the 0.1 mg/kg guideline level (Table 43) used for most sensitive root crops. However, if the sludge organic concentration was 10 mg/kg instead of 1 mg/kg, then the same sludge loading would give a soil residue level of 0.4 mg/kg and cause such a soil to be excluded for growing vegetable root crops. But a sludge with 10 mg/kg of aldrin or dieldrin would still be acceptable if 1 ton per hectare per year was applied for 10 years instead of 10 ton per hectare per year. Therefore, the total amount applied to a soil is the critical factor rather than the concentration in the sludge. Again, as noted above, these examples assume no loss of the organic chemical by volatilization, degradation, etc. from soil.

## EXTRACTION/LEACHING PROCEDURES

Under the Amendments to the Resource Conservation and Recovery Act of 1985 [Hazardous Waste Management System; Definition of Solid Waste; Final Rule (40 CFR Parts 260, 261, 264, 265 and 266), January 4, 1985], the U.S. EPA was directed to improve the ability to characterize hazardous waste. The Extraction Procedure Toxicity Characteristic (EPTC), or EP toxicity test, currently used entails a leaching test to measure the tendency of a waste to leach, coupled with extract concentrations above which the waste is to be regulated, and defined as a hazardous waste. This test was developed on the premise that a potentially hazardous industrial waste might be sent to a sanitary landfill, resulting in a high potential for groundwater contamination. The constituents currently included as part of this test were those for which National Interim Primary Drinking Water Standards have been established. These standards addressed 8 inorganics and 6 organic compounds (2,4-dichlorophenoxyacetic acid, endrin, lindane, methoxyclor, toxaphene, and 2,4,5-trichlorophenoxyacetic acid).

As part of the effort to improve the characterization of hazardous waste, EPA will be proposing a revised test (Friedman, 1985) that would expand the list of organic compounds tested to 44 and modify the procedure itself to the Toxicity Characteristic Leaching Procedure (TCLP). If the extractant concentration from the TCLP is above the maximum threshold limit for any of the 8 inorganic or 44 organic chemicals, the waste is defined as hazardous. Each municipality that produces sewage sludge must make the determination whether or not their sludge is hazardous. This determination can be based upon their knowledge of their sludge or they may choose to use the TCLP to help them make it. The EPA believes that the way to determine if a municipal sludge is hazardous is to determine whether or not its

extract concentrations exceed the maximum threshold limits. However, many have argued that a testing procedure based on the worst-case scenario in which large quantities of sludge are disposed of in a landfill has little relevance to assessing any potential hazard from recycling low rates of sludge to land.

The U.S. EPA is currently having eight sewage sludges tested with the new TCLP procedure. The sludges were selected to include purely domestic sewage sludge as well as sludges expected to have high concentrations of contaminants from industrial sources. While none of the sludges appear to have TCLP extract concentrations that exceed the threshold limits, results are too preliminary to know for sure. Therefore, the impact of changing from the EPTC to the TCLP on sludge application programs is too early to ascertain.

#### MUTAGENICITY TESTING OF SLUDGES

A number of studies have recently reported results of mutagenicity tests on extracts of sludge (Babish et al., 1983; Boyd et al., 1982; Hang et al., 1983; Hopke and Plewa, 1984; Hopke et al., 1982). While most sludge extracts tested by Babish et al.(1983) were mutagenic by the Ames test (Ames et al., 1975) many foods, drinking water, and other substances in our environment also test positive for mutagenic activity (Loper, 1980; Mast et al., 1984; Nagao et al., 1979; and Salmeen et al., 1985). Ames (1983) has also indicated that "the human diet contains a great variety of natural mutagens and carcinogens, as well as many natural antimutagens and anticarcinogens". Thus, one must use extreme care in interpreting mutagenic tests of sludge extracts to keep them in perspective with the presence of mutagenic constituents in all parts of our environment.

In addition to the Ames <u>Salmonella</u> assay, plant test systems have been used to investigate the <u>mutagenic</u> activity of sewage sludges (Hopke and Plewa, 1984; Hopke et al., 1982). Mutagens present in sludge-amended soil can be transported into a crop plant and induce genetic damage in germ cells; however, no mutagenicity occurred in the kernels from corn grown on sludge-amended soil nor were mutagens transferred from the sludge to soil or surface waters.

These studies imply that the chemicals causing mutagenicity are trace organics, but the specific chemicals responsible for the mutagenic affects have not been identified. Another difficulty is interpreting these bioassay results, since agricultural soils can exhibit mutagenic response without sewage sludge amendments (Boyd et al., 1982; Brown et al., 1985; and Hopke et al., 1982). Therefore, results of these mutagenicity tests are not easy to put into perspective (Davis et al., 1984; Dean and Suess, 1985), and the data cited suggest that mutagen activity is greater for sludges generated by more industrialized municipalities.

While mutagens present in sludges were shown to degrade relatively rapidly (e.g., within 2-3 weeks) in one sludge-amended soil (Angle and

Baudler, 1984), recent work at Pennsylvania State University indicates that the loss of mutagenic activity may take as long as one growing season for other sewage sludge/soil mixtures (Baker et al., 1985). How well results from these laboratory incubations will duplicate under field conditions is untested and still unknown.

Due to the large number of organic chemicals which can be present in sewage sludge, a short-term bioassay offers the advantage of testing for potential biological toxicity inherent in a sludge (or other waste) containing a complex mixture of chemicals (Brown et al., 1982). For example, Peters (1985) used the Ames test to screen 38 Pennsylvania sewage sludges containing potentially harmful trace organics, and Brown et al. (1982) used the Ames test plus two other bioassays to examine the acute toxicity of ten hazardous wastes.

Using a bioassay test for identifying a sludge contaminated with an organic chemical(s) could provide an additional degree of safety in managing sewage sludge applications to agricultural/forest soils. To be useful, however, bioassay test results for sludges must be correlated to mutagenic activity or biological toxicity of soil/sludge mixtures in the field. Based on the generally low concentrations of trace organics in sludges and the low rates of sludge (e.g., agronomic) typically applied, the probability of any transfer of mutagenic activity to animals or humans as a result of sludge application to land is very low.

## FATE OF TRACE ORGNICS ADDED TO SOIL-PLANT SYSTEMS

Potential health hazards associated with organic chemical residues in sludge applied to land have been discussed in several review articles (Chaney, 1984; Dacre, 1980; Davis et al., 1984; Kowal, 1983; Kowal, 1985; Majeti and Clark, 1981; and Pahren et al., 1979).

Principal pathways by which organics could be transferred to humans from sludge-amended soils were listed by Dean and Suess (1985):

- 1. Uptake by plant roots in sludge-treated soil, transfer to edible portions of plants, consumption by humans;
- 2. Direct application to edible parts of plants as sludge, or as dust or mud after sludge is mixed with the soil, consumption by humans;
- 3. Uptake via plants used as feed or fodder for animals, transfer to animal food products, consumption by humans;
- 4. Direct ingestion of soil and sludge by grazing animals and transfer to animal food products, consumption by humans;
- Direct ingestion of sludge contaminated soil by children; an abnormal behavior called "pica".

Two other possibilities might be included with this list:

- Surface runoff/erosion to streams or rivers used as a source of drinking water downstream, and
- Leaching to a groundwater aquifer used as a source of drinking water.

These pathways have all been demonstrated but are not equally important. Indeed, Pathway 4, which does not go through plants, is the only one by which organic pollutants have been traced from sludge to animal products (Chaney, 1984).

While plant contamination can occur (as discussed later), soil residue levels necessary for this to happen are usually higher than would be anticipated from low application rates of non-industrialized sewage sludges. In addition, soil incorporation of organics, like PCBs for example, can greatly reduce plant "uptake" of the chemical (Harms and Sauerbeck, 1983). Lindsay (1983) also reported that several recalcitrant organics are so strongly bound to soil and sludge as to be almost totally unavailable for plant uptake.

Trace organics may biomagnify. For example, detritus eating insects were found to contain 1.3 x the soil concentration of PCB, which could lead to further bioconcentration in insect-eating birds (Davis et al., 1984). As with metals, trace organics may accumulate in animal food products following direct sludge ingestion during grazing. The problem is particularly important for dairy cows since milk is the animal product most likely to be influenced by organic contaminants in sludge applied to land (Dean and Suess, 1985), although management practices can significantly reduce this possibility.

The potential exists for direct ingestion of organics, especially by children through the phenomenon of pica (Pathway 5) if sludge was used to fertilize home gardens. Dean and Suess (1985) concluded, however, that this is likely to be a minor or insignificant route of exposure, as is inhalation of dust or vapors. As with Pathway 1, significant human consumption of sludge organics by human management (e.g., culinary procedures like cleaning and peeling of root crops that tend to accumulate lipophilic substances) would seem most unlikely (Naylor and Loehr, 1982b).

Bioaccumulation factors (i.e., ratio of an organic in plant or animal tissue to concentrations in soil) are available for very few compounds. For plants, the factor (when known) is almost always <1 and usually <0.1 (Overcash, 1985) and for animal products (e.g., milk) estimates of 0.7 for PCB (Fries, 1982) and 0.5 for dieldrin (Lindsay, 1983) have been made. Field data are largely non-existent, but Baxter et al. (1983) reported no plant uptake of 22 persistent organics from land amended with Denver Metro sludge. Also, no increases in persistent trace organics content of the fat tissue content of cattle grazing a sludge application site were observed.

If ingested, organics present in sludge or soil can be bioavailable (Chaney, 1984; McConnell et al., 1984). Jelinek and Braude (1977) found an increased content of PCBs in the milk fat of cattle fed green forages and roughages grown on sludge-treated land. This prompted the U.S. FDA to recommend a maximum permissible content of not more than 10 mg/kg PCBs in sludges used on agricultural lands (Braude et al., 1975). Therefore, apparently absorption of ingested sludge organics can only be prevented by limiting their concentration in sludges or avoiding direct ingestion until compounds have been degraded or dissipated.

# Assimilative Pathways Within the Soil-Plant System

Organic compounds may undergo a variety of chemical and biological processes when applied to a soil or soil-vegetation system. The various assimilative pathways have been discussed by several authors (Davis et al., 1984; Kaufman, 1983; Lue-Hing et al., 1985; Overcash, 1983). The pathways include:

- 1. adsorption onto soil and its constituents;
- 2. volatilization;
- 3. degradation (microbial, chemical, photolysis);
- 4. leaching to groundwaters and runoff/erosion to surface waters:
- 5. plant retention (contamination vs. uptake and translocation); and
- 6. macro- and micro-fauna uptake (bioaccumulation by insects, grazing animals).

While research on the fate of sludge organics in soils is limited, the behavior of organics in soil, has been extensively studied, particularly agricultural pesticides, (e.g., Guenzi, 1974; Goring and Hamaker, 1972) and with non-agricultural chemicals in the petroleum industry (API, 1983). In general, trace organics are strongly adsorbed to soils and its constituents, especially soil organic matter. Thus, leaching and plant uptake are usually very limited. Some runoff/erosion may occur for organics firmly bound to soil particles or debris, but this can be minimized by using good soil and water conservation practices at sludge application sites.

Some trace organics (notably, PCBs, lindane, dieldrin) are known to volatilize readily when surface applied, although soils and sludge itself can drastically reduce these volatizilation losses. Some organics are recalcitrant to microbial degradation, but most are expected to degrade. Of concern are the persistent organics and some of the readily degraded components that can break down to toxic metabolites. Pathways of most interest are plant uptake/contamination, degradation, volatilization, and leaching.

Chaney (1984) provided a good discussion about the uptake of organics by plants. Because this summary seemed more appropriate than others (Davis et al., 1984; Harms and Sauerbeck, 1983; Kaufman, 1983; Lue-Hing et al., 1985; and Overcash, 1983) and is not readily available, much of his discussion was used for this section, sometimes verbatim. Readers are encouraged to review these other references, however, for a more complete understanding of plant uptake/contamination.

Trace organics can enter edible parts of plants by two processes:

1) uptake from the soil solution, with translocation from roots to shoots, or 2) absorption by roots and shoots of volatile organics from the soil. Systemic pesticides are applied to the soil, then absorbed and translocated to the plant leaves. These kinds of compounds are quite water soluble and would probably not appear in wastewater treatment sludges at appreciable levels.

Lipophilic halogenated, organics represent the case for water insoluble compounds which are largely sorbed by plants from the soil air or the organic-enriched air near the soil surface. Beall and Nash (1971) developed a method to discriminate between movement of an organic through the plant vascular system (uptake-translocation) vs. vapor phase movement. They found soybean shoots were contaminated by soil-applied dieldrin, endrin, and heptachlor largely by uptake-translocation, while vapor transport predominated for DDT and was equal to uptake-translocation of endrin. Using this method, Fries and Marrow (1981) found PCBs reached shoots via vapor transport, while the less volatile PBBs did not contaminate plant shoots by either process (Chou et al., 1978; Jacobs et al., 1976).

Root crops are especially susceptible to contamination by the vaportransport route. Carrots have a lipid-rich epidermal layer (the "peel") which serves as a sink for volatile lipophilic organics. Depending on the water solubility and vapor pressure of the individual compound, it may reside near'y exclusively in the peel layer of carrots, or penetrate several millimeters into the storage root (Lichtenstein et al., 1964, 1965; Jacobs et al., 1976; Lichtenstein and Schulz, 1965; Iwata and Gunther, 1976; Iwata et al., 1974; Fox et al., 1964; Landrigan et al., 1978).

Carrot cultivars, however, were found to differ in uptake and in peel vs. pulp distribution of the chlorinated hydrocarbon pesticides endrin and heptachlor (Lichtenstein et al., 1965; Hermanson et al., 1970). Other root crops (sugar beet, onion, turnip, rutabaga) are much less effective in accumulating lipophilic organics in their edible roots, possibly because the surface of the peel is lower in lipids (Moza et al., 1979, 1976; Fox et al., 1964; Chou et al., 1978; Lichtenstein and Schulz, 1965).

Based on carrot accumulation of volatile chlorinated hydrocarbon pesticides, Iwata et al. (1974) evaluated PCB uptake by carrots from a low

organic matter (0.6%) sandy soil, which represents a worst-case surface soil, in the field (it should be recognized that 100 mg/kg is a very high concentration of PCBs in soil. For the environmentally persistent 5 and 6 chlorine isomers, unpeeled fresh carrots contained PCB at about 4.9% of the soil level. Peeling removed 14% of the carrot fresh weight and 97% of the PCB residue, so peeled fresh carrots contained PCB at only 0.16% of the soil PCB level.

The level of chlorinated hydrocarbon in carrots is also sharply reduced by increased organic matter in soil. The increased organic matter adsorbs lipophilic compounds and keeps them from being released to the soil solution or soil air (Filonow et al., 1976; Weber and Mrozek, 1979; Chou et al., 1978; Strek et al., 1981). Since added sewage sludge can increase the ability of soils to adsorb PCBs (Fairbanks and O'Connor, 1984), the increased sorption capacity may fully counteract the PCBs in sludge at low levels.

The residue of PCBs in waste materials such as municipal sludge can be depleted of the more volatile and more easily biodegraded lower chlorinated compounds. Because plant contamination (uptake via volatilization) by the higher chlorinated compounds is much less than for the more volatile lower chlorinated compounds at equal soil levels (Iwata and Gunther, 1976; Suzuki et al., 1977; Moza et al., 1976, 1979; and Fries and Marrow, 1981), the lack of plant contamination from sludge-applied PCBs is not unexpected. For example, in a study by Lee et al. (1980), a sludge containing 0.93 mg/kg PCBs was applied at a rate of 112 dry mt/ha, yet "no PCBs were detected in the sludge grown carrots". Since other root crops are not nearly as good PCB accumulators as carrot (Moza et al., 1979), remarkably low potential human PCB exposure would be predicted for recommended sludge utilization practices.

Other research efforts have centered on assessing risk from polycyclic aromatic hydrocarbons (PAHs), some of which are carcinogenic (e.g., benzo(a)pyrene). Researchers found that carrot roots (but not mushrooms) accumulated many PAHs from compost-amended soils (Müller, 1976; Linne and Martens, 1978; Wagner and Siddiqi, 1971; Siegfried, 1975; Siegfried and Muller, 1978; Ellwardt, 1977; Borneff et al., 1973). The level of 3,4-benzypyrene in carrot roots declined with successive cropping of compost amended soils. Harms and Sauerbeck (1983) also found PAH contamination of potato tubers, radish and carrots where direct contact with the soil allowed transfer of these organics. Concentrations in the aboveground parts of plants were, however, low.

Nitrosamines are another group of organics which have been found in sewage wastes (Yoneyama, 1981; Green et al., 1981). Although accumulated from nutrient solution and soil by plants (Brewer et al., 1980; Dean-Raymond and Alexander, 1976), nitrosamines appear to be rapidly degraded in soils and plants. Research on N-nitrosodimethylamine and N-nitrosodiethylamine found rapid degradation in soil; plant uptake could occur initially but these compounds were rapidly degraded (Dressel, 1976a,

1976b). While traces of nitrosamines are found in nitroanaline based herbicides, these compounds are rapidly degraded and no detectable nitrosamine was found in soybean shoots (Kearney et al., 1980b). An International Union for Physics and Chemistry (IUPAC) committee assessed the environmental consequences of these trace nitrosamines, and found no risk to the fuod chain (Kearney et al., 1980a).

Many other carcinogenic or toxic compounds could be present in sludges and contaminate the food chain through plant uptake or volatile contamination of crop roots or shoots. While information on these other organics is limited, two data bases are available which consider plant uptake of organic molecules. PHYTOTOX deals with the direct effect of exogenously supplied organic chemicals on the growth and development of terrestrial plants (Royce et al., 1984). As of July 1985, 9,800 papers had been included with data on 3,500 chemicals and 700 species (Rygiewicz, 1986, Personal communication). This data base is now available through a private service (Fein-Marquart Associates, 7215 York Rd., Baltimore, MD 21212). The second data base (UTAB) contains information pertaining to the Uptake, Transport, Accumulation and Biotransformation of organic compounds  $\overline{b}y$ vascular plants. This data base includes 3,900 papers, with information about 700 chemicals and 250 species and is available through the University of Oklahoma (John Fletcher, Dept. of Botany, Univ. of Oklahoma, Norman, OK 73019). These data bases offer the opportunity to evaluate basic research on the uptake of organics by plants which may help to understand the effects of sludge applied organics.

# Degradation

Degradation of organic chemicals in soil may occur by chemical, photochemical, or biological processes. Degradability of a compound depends on its chemical structure, some being rapidly decomposed while others are relatively recalcitrant to degradation. Biodegradation can occur in microbial cells, in the soil solution by chemical mechanisms, or by extracellular enzymes sorbed to soil particles (Kaufman, 1983).

Often, soil microbes capable of degrading a compound proliferate in soil, and the effective population may remain several years after the last treatment. Maintaining a supply of biodegradable organic matter in soils receiving wastes would likely provide a higher population of diverse microbes capable of degrading more kinds of trace organics. Microbes may utilize a particular organic as an energy source, or may cometabolize it with other normal metabolic processes. Although the kinds of organisms and even types of enzymes involved in biodegradation are known for some pesticides and other organics, little is known about most of the organics found in wastes like sewage sludges (Kaufman, 1983).

Microbiological as well as chemical reactions are usually acting simultaneously. Chemical reactions (abiotic routes) are a part of the overall measure of organic compound decomposition. Two typical reactions

are hydrolysis and neutralization of the parent organic species, but such reactions typically leave the bulk of the parent structure still intact (Overcash, 1983). Soil factors known to affect chemical degradation of organics include temperature, aeration, microbial populations, pH, organic matter, clay, cation exchange capacity, and moisture (Kaufman, 1983).

The action of sunlight may chemically alter and degrade organic chemicals in the environment. The importance of photochemical reactions to the degradation of waste organics applied to land will depend largely upon the mode of application and soil incorporation. Sludge organics can be subjected to some photolytic action during the time they are on exposed soil surfaces following a surface application (Kaufman, 1983). Under these conditions an organic compound may be degraded via photolytic mechanisms. Phenolics and polynuclear aromatics are two groups that readily undergo such reactions (Overcash, 1983). Photolytic degradation will be nonexistent, however, when sludges are incorporated into the soil since sunlight does not penetrate the soil surface (Kaufman, 1983).

Following an extensive literature search concerning the decomposition of specific organics in the terrestrial environment, Overcash (1983) concluded that very few organic compounds can be said to be non-degradable. Considering the long time periods typical in soil systems, only two classes of compounds were regarded as nondegradable based on available terrestrial research information: (1) synthetic polymers manufactured for stability, and (2) very insoluble large molecules, e.g., 5-10 chlorinated biphenyls (Overcash, 1983).

Other organics will have varying decomposition half-lives or persistence in soils. Overcash (1983) provided examples of half-life ranges for several organic chemicals (Table 44) and Kaufman (1983) listed the relative persistence for several organic chemical classes (Table 45). Tabak et al. (1981) compared the relative biodegradation of organic priority pollutants with a static culture flask procedure. While their decomposition results may not be directly extrapolated to degradation of organics in the soil, the relative degree of biodegradation may prove to be similar in soils. Significant biodegradation was found for phenolic compounds, phthalate esters, napthalenes, and nitrogenous organics; variable results were found for monocyclic aromatics, polycyclic aromatics, polychlorinated biphenyls, halogenated ethers, and halogenated aliphatics; and no significant biodegradation was found for organochlorine pesticides.

#### Volatilization

Vapor movement of organics (i.e., diffusion and volatilization) are important factors affecting the distribution and persistence of some organic chemicals in soil. An estimate of potential volatility can be obtained from the ratio of water solubility to vapor pressure, which indicates the proportion of an organic in the vapor phase. This ratio is only a guide.

however, since adsorption of the organic in the soil will decrease the amount present in the vapor phase (Kaufman, 1983).

An organic spread on the surface of or injected into a soil with sludge will partition between the gas and liquid phases to exert a vapor pressure. The conditions of the soil and the application technique used, as well as the inherent organic compound volatility, are important factors in quantifying how an organic compound might be lost through volatilization. The level of vapor pressure at which volatile losses are known to be significant is usually taken at  $5 \times 10^{-6}$  mm Hg at  $25^{\circ}$ C. However, vapor pressure alone may be misleading because highly volatile organics like toluene are prevalent in municipal sludges, even after opportunities have occurred for volatile loss during wastewater treatment (Overcash, 1983).

Volatilization losses were considered as significant processes of organic chemical removal when wastewaters are applied to land (Chang and Page, 1984; Jenkins et al., 1983). Jenkins et al. (1983) stated "as a rule the higher the vapor pressure the lower the water solubility, the higher the Henry's law constant and the higher the removal rate by volatilization." Once the organic reaches the soil, the actual volatilization loss of trace organics from the soil will depend on factors affecting the movement of the organics to the soil surface and its dispersion into the air (Chang and Page, 1984).

For soil-applied pesticides, the vapor density was found to be the main factor controlling volatilization (Farmer et al., 1972). Other factors which affect volatilization include soil pesticide concentration, temperature, rate of air movement over the soil surface, and soil water content (Farmer et al., 1972; Igue et al., 1972). More recently Jury et al. (1983, 1984a,b,c) have used benchmark properties of vapor density and solubility in water in a mathematical model to determine the relative volatility of a specific soil-applied organic.

As with earlier work done with pesticides, how well research results will predict volatilization losses for the same organics applied to soils as part of a sludge matrix is unknown. Research reported by Fairbanks and O'Connor (1984) indicate that sludge additions to soil can decrease volatilization losses of PCBs, so the sludge matrix could be expected to have some effect. Nevertheless, models and research data which apply to soilapplied pesticides provide a good "point of departure" for understanding potential volatilization losses of pesticides and other organics added by sludge applications.

## Leaching

The downward movement of an organic chemical is largely governed by sorption and biodegradation. At least two steps are involved in the leachability of an organic chemical in soil: (1) entrance of the compound

into solution, and (2) adsorption of the compound to soil surfaces (Kaufman, 1983). Partitioning between the adsorbed and soil solution phases may occur immediately upon application to the soil or may be delayed until the organic separates from the waste medium. At the same time, decomposition reactions can vary the actual amount of a particular organic compound that resides on the soil/waste phase or in the soil-water solution (Overcash, 1983).

Therefore, the inherent persistence of each chemical in soil will affect whether any mobile chemical might pollute groundwater. The half-life of many organic chemicals in soil is sufficiently short to make it highly unlikely that the chemical would ever reach the water table under ordinary field leaching conditions (Kaufman, 1983). Overcash (1983) also concluded that for most sludge application sites where normal application rates and management techniques are used, leaching of organics is probably negligible.

# Effects of Sludge Properties

Few studies have considered the effect of sludge on the assimilative pathways of adsorption, volatilization, and degradation. Since organics typically associate with the organic fraction of soils, one might expect even greater retention of trace organics in amended vs. unamended soils as was shown by Fairbanks and O'Connor (1984) for di-3-(ethylhexyl)phthalate (DEHP), PCBs, and two herbicides. The greater adsorption in sludge-amended soils should reduce contaminant mobility and plant availability, and data did show that volatilization of PCBs from sludge-amended soils was significantly reduced (Fairbanks and O'Connor, 1984).

Sludge additions may also affect organic contaminant degradation. The increased microbial activity found in sludge-amended soils suggests that previous sludge applications cause a preconditioning with respect to microbes and/or enzymes which may increase organic degradation by cometabolism (Fairbanks and O'Connor, 1984). The degree and duration of sludge effects on trace organic behavior are influenced by type and concentration of compound, incubation time, sludge rate, and soil type.

Most experiments designed to determine the effects of sludge or organic behavior have used "spiked" systems in which the target organic is added as reagent grade chemical to soil or soil-sludge systems. Preequilibrium of the target organic with sludge has been minimal. Thus, most data generated to date are tainted by limitations similar to the early "mineral salt" work with metals. Organics indigenous to sludge may have drastically different properties with respect to their fate in soils compared to these same trace organics added to the soil alone or in combination with sludge. Research is needed with selected sludges to study the assimilative pathways of specific organics (indigenous to these sludges) compared to results for comparable amounts of the same organic added to the same soil in the absence of the sludge matrix. Correlations

between controls and sludge-treated soils could then be used to predict the fate of other sludge-applied organics when actual field data are unavailable.

## Utilizing Physical/Chemical Properties and Models

Due to the thousands of organics which can potentially be present in society's wastes such as sewage sludge, the task of researching each organic to determine its fate in the environment is impossible. A more realistic approach would be to utilize basic physical/chemical properties of organics and soils to compare the research results assessing the environmental fate of selected organics, representative of larger groups, with the fate predicted by mathematical models.

The more important physical/chemical characteristics for assessing the potential transport, persistence, and fate of substances in sludge land applications are: (a) water solubility, (b) soil adsorption-partitioning, (c) half-life in soil, and (d) vapor pressure. Laboratory measurements can be used to obtain values for all these characteristics except soil nalf-life, or they may be estimated by methods such as those discussed by Lyman et al. (1982).

The fates of greatest interest for sludge organics incorporated into the soil are volatilization, degradation, plant uptake, and leaching. The persistence, or ease of degradation, and volatilization of an organic are major characteristics which will affect the time during which an organic may be "available" for plant uptake or loss by leaching to groundwater. Adsorption to soil colloids (organic and inorganic) and water solubility of an organic are also important factors which help determine this availability. When plant uptake and leaching are not significant for an organic, potential for transfer back to man is reduced. Likewise, when an organic is completely degraded in soils, additional pathways (discussed earlier in this section) for transferring a sludge organic to humans are eliminated.

Examples of using benchmark properties in mathematical models or for estimating the behavior of organics applied to soils include Chang and Page (1984), Jury et al. (1983, 1984a,b,c), and Wilson et al. (1981). Based on calculations using the soil adsorption coefficient, water-air partition coefficient, and octanol-water partition coefficient, Chang and Page (1984) compared the environmental fate and transport in soils of several pesticides with several trace organics. Their conclusion regarding the addition of wastewater organics to soils was that their environmental impact was not expected to be very significant.

Using a simple mathematical model based on water solubility of an organic chemical and the organic carbon content of the soil, Wilson et al. (1981) were able to predict the retardation factors for 13 organic pollutants within a factor of three. They found retardation by soil with respect to water movement generally increased with decreasing water solubility.

Jury et al. (1983) developed a more complicated mathematical model for describing transport and loss of soil-applied organic chemicals. This screening model uses benchmark properties (organic C partition coefficient, vapor pressure, solubility, half-life) to determine the relative convective mobility, diffusive mobility, volatility, and persistence (Jury et al., 1984a,b). When this model was tested on published experimental data for volatilization, leaching, and persistence, experimental results and those predicted by the model agreed reasonably well (Jury et al., 1984c).

Although experiments under field conditions are the only reliable way to determine the fate of an organic applied to soils with sewage sludge, the expense and time required to test the large number of organic chemicals used in society and found in sewage sludges are prohibitive. Therefore, models like those mentioned above can and should play a significant role in assessing the environmental risk of applying sludge organics to soils.

# COMPARISON OF MUNICIPAL SLUDGE EXPOSURE/PISK ASSESSMENTS

Several independent evaluations have recently been published to assess the relative risk from specific organic compounds present in municipal sludge when applied to land (Metro, 1983; Munger, 1984; Connor, 1984; Naylor and Loehr, 1982a,b; EPA, 1985). All of the risk assessments cited above are published in non-peer reviewed journals. No in-depth scientific evaluation or analysis was performed on these individual exposures/risk assessments. Therefore, their results and conclusions should be viewed with this in mind. As far as the authors are aware, no risk assessments for sludge have appeared to date in a peer-reviewed journal.

Risk could be defined as a measurement of the probability of harm occurring to human health as a result of an organic chemical being present in land-applied sludge (Munger, 1984). To the extent possible a common concept was used to assess "acceptable risk", i.e., the ratio of daily intake required to stay below a risk level of  $10^{-6}$ . If this ratio is greater than 1.0, the resulting risk level is (numerically) greater than  $10^{-6}$  and when less than 1.0, the risk level is less than  $10^{-6}$ . This ratio is actually the inverse of the "hazard index" used in EPA environmental profiles (EPA, 1985) but was the method chosen to express risk levels in the other risk analyses.

A risk assessment concerning the health effects of land applying sludge was prepared by S. Munger for the Municipality of Metropolitan Seattle (Metro, 1983). This assessment was updated and expanded specifically for the "Municipal Wastewater Sludge Health Effects Research Planning Workshop" held in Cincinnati by EPA in January 1984 (Munger, 1984). That estimate of risk was based on the quantities of soil, water, or food obtained from a sludge application site which could be consumed without exceeding a risk level of  $10^{-5}$ .

The assumptions and values used by Munger (1984) for assessing the risk of two organics, polychlorinated biphenyls (PCBs) and benzo(a)pyrene [B(a)P], are shown in Table 46. The concentrations of PCBs and B(a)P which would occur in the various environmental compartments (i.e., soil, water, plant and animal tissues) were estimated. Using these concentrations (Munger, 1984) and the organic chemical consumption equivalent to  $10^{-6}$  risk level, the quantities of various materials that could be consumed without exceeding this risk were calculated (Tables 47 and 48). The quantities for materials from sludge-amended areas could then be compared to similar materials from control (untreated) areas to evaluate any increased risk due to sludge application to forest land. The author (Munger, 1984) concluded that for these two organics, PCBs and B(a)P, any increased risk would be minimal and could be controlled by proper site management.

The Naylor and Loehr efforts (1982a,b) considered several perspectives on risk, including the human intake routes by comparing the addition of sludge organics to soils with the application of agricultural chemicals to soil. Naylor and Loehr (1982a) begin by defining relative toxicity categories (Table 49) and then comparing the toxicities of common chemicals (Table 50), common pesticides (Table 51) and selected priority pollutant organics in 13 sludges (Table 52). In general, the agricultural chemicals are more toxic than the sludge organics. When comparing normal pesticide rates, the projected application rates for sludge organics are usually 10 fold less than for pesticides. In this perspective municipal sludge practices were judged to have no greater risk than using agricultural chemicals.

A second form of comparison was more similar to that used in the EPA Profiles for direct consumption of sludge against acute (LD $_{50}$  dose) and chronic (acceptable daily dose, or  $D_T$ ) concerns. From the group of 24 sludge organics listed, Naylor and Loehr (1982b) selected three for further evaluation: (1) hexachlorobutadiene (HCBD), a highly toxic chemical; (2) bis(2-ethyl-hexyl)phthalate, a chemical having a relatively low toxicity but present in high concentrations; and (3) 1,1,2-trichloroethane, a chemical of moderate concentration and moderate toxicity.

Based on the maximum sludge concentrations, the rat or a cow would have to eat an amount of sludge equivalent to more than 10 times it body weight to ingest a LD<sub>50</sub> dose of the most texic HCBD (Table 53). When considering a chronic exposure occurring by daily intake of sludge alone or soil treated with sludge (Table 54), a "pica" child would have to consume sludge for 41 years to consume a LD<sub>50</sub> dose of HCBD, the most toxic sludge organic Naylor and Loehr (1982b) considered. When considering the more logical case of a "pica" child consuming sludge treated soil rather than pure sludge, a safety factor of 45 to 450 was obtained (Table 54). Therefore, from a different perspective, 24 organics in sludge were judged to be a relatively low health hazard when sludges are land applied at agronomic rates.

Connor (1984) used the same sludge characteristics as Naylor and Loehr (1982a,b) but an independent risk/exposure assessment (Table 55). Assuming

a low (15 g) and high (139 g) amount of contaminated soil per day is ingested, safety factors were calculated based on soil concentrations expected from pesticide or sludge application. The safety margins were (1) greater for sludge organics than for common agricultural chemicals applied to soils and (2) greater than 1 for all chemicals except the polycyclic aromatic hydrocarbon (PAH) group which had a margin of safety of about 0.3 to 0.03.

While Naylor and Loehr (1982b) did not develop a dietary scenario for PAH chemicals, they cited references which indicated that soils may contain natural PAH concentrations of 0.05-0.14 mg/kg and manure can contain 0.15 to 1.21 mg PAH/kg. The addition of PAH assumed by Connor (1984) in his risk assessment (Table 55), i.e., 15 mt/ha of sludge containing 13 mg/kg of PAH, would be equivalent to a soil concentration of 0.08 mg PAH/kg soil Therefore, Connor's analysis for PAH would suggest that any soil containing the background levels given above would have a low safety factor.

The fourth risk assessment reviewed for sludge organics is the environmental profiles and hazard indices conducted on several sludge constituents (EPA, 1985). The methodology used to assess the risk of a particular organic included the use of 12 indices to evaluate different pathways by which the sludge-applied organic could be exposed to plants, animals, or humans. The index which most lends itself to a comparison of the Naylor and Loehr (1982a,b) and Connor (1984) assessments is index 12, the index of human cancer risk from soil ingestion by a "pica" child.

Only two organics included in Naylor and Loehr's or Connor's list of organics had an index 12 value calculated: Hexachlorobutadiene (HCBD) and B(a)P. By inverting the EPA hazard index value, essentially a safety factor value can be obtained. The safety factor for HCBD was calculated as 238 by EPA (1985) and 45 by Naylor and Loehr (1982b), and for B(a)P was 0.018 by EPA (1985) and 0.27 for total PAH by Connor (1984). While these results vary, there was agreement in terms of whether risk was greater than or less than  $10^{-6}$ .

Following a similar calculation as Naylor and Loehr (1982b) used for bis(2-ethylhaxyl)phthalate and HCBD, a safety factor was also calculated for methylene chloride (MeCl) and phenanthrene. Comparing the safety factors of Connor's and Naylor and Loehr's results for 15 g/day soil ingestion show:

	Naylor & Loehr	Connor
ВЕНР	415	4,700
MeC1	4.6	150,000
PAH	35 (phenanthrene)	0.27 (total PAH)

Results tended to be in the same direction from 1.0 except for PAHs, although the magnitude of differences were substantial.

Differences between the three risk assessments can be attributed to differences in acceptable daily intake values used, sludge concentrations assumed, sludge rates assumed, quantities of soil assumed to be ingested, etc. The large differences obtained by these authors and cited above indicate how important it is to have correct assumptions and realistic data for calculating safety factors. When the accepted daily intake values vary by  $10^3$  or more between assessments, large differences can be expected for the safety factors obtained.

Answers obtained using risk assessment methodology must provide realistic values to be useful. If nothing else, the methodology will indicate weaknesses in the assumptions or data used when unrealistic values are obtained with these models and help identify where further research data are needed. Overall, the general consensus of these risk assessments seems to indicate that organics applied to soil from sludge will not increase the health risk to animals and humans. However, the data base on which the previous statement is made is limited, and better risk assessment methodologies for land application of organics from sludge are urgently needed.

### CONCLUSIONS

- 1. Because sewage sludges can theoretically contain thousands of organic compounds, wastewater treatment plants should identify the organics being discharged by users, particularly industry. This information should guide the testing of sludges for appropriate organics to determine concentration levels.
- 2. Available surveys measuring trace organic concentrations in sludges indicate that sewage sludges can have unusually high concentrations (i.e., a few percent, dry weight). Most organics are present at concentrations less than 10 mg/kg and about 30% of the organics tested were below detection limits. Based on their prevalence and potential loading to soils using agronomic or low sludge rates, sludge organics appear to have minimal risk.
- 3. Mutagenicity tests have been used to evaluate the safety of sewage sludges from land application. While the test might provide an additional means of checking sewage sludges prior to land application, they are difficult to interpret and have not been correlated to mutagenic activity of soil/sludge mixtures in the field. Therefore, their value for helping manage land application programs is uncertain at this time.
- 4. Major assimilative pathways for organic chemicals applied to the soil-plant system include adsorption, volatilization, degradation, leaching, and plant uptake. Many organics are strongly adsorbed to soil organic matter and/or undergo degradation, reducing the potential for plant uptake or leaching.

- 5. Due to the thousands of organics which can potentially be present in sewage sludges, a realistic approach may be to utilize basic physical/chemical properties of an organic and mathematical models to predict the likely fate of that sludge organic in the soil. However, limited field research with selected sludge organics, which are representative of organic chemical groups, is needed to validate these models.
- 6. In general, risk assessments appear to suggest that most sludge organics will not increase the health risk to animals and humans, based on their relative toxicities and anticipated loadings to soil from agronomic or low sludge application rates.

TABLE 40. SUMMARY OF ORGANIC CHEMICALS FOUND IN SEWAGE SLUDGES.

	No. of sludges	*	Concentration Positive, i.e.	., > Detection	on Limits	
Chemical	tested	Occur- rence#	mg/kg (d Range	Median	ug/L Range	Ref. tt
		Phtha l	ate Esters			
Bis(2-ethylhexyl)phthalate	234	84	0.415 -58,300	168		1
	437	95			2-47,000	2
Butylbenzylphthalate	234	60	0.0469-12,800	59.1		1
	4 37	43			2-45,000	2
Diethylphthalate	234	63	0.0987-3.780	50.0		1
•	4 37	9			1-786	2
Dime thylohthalate	236	23	0.106-941	11.7		1
	437	5	,	****	3-650	2
Qi-n-butylphthalate	237	45	0.0776-3,210	17.3		1
	4 37	45			1-6,900	2
Di-n-octylphthalate	237	40	0.0222-2,610	4.9		1
	437	10	·		4-1,024	2
		Monocy	clic Aromatics			
Benzene	436	61			1-953	2
Chl orobenzene	158	6	2.06-846	10.2		1
	436	13			1-687	2
1-chloro-2,4-dinitrobenzene	0/238	0	All samples <	detection li	mit	1
1-chloro-2,6-dinitrabenzene	0/238	0	All samples (	detection li	mit	1
1-chloro-2-nitrobenzene	238	0 -	All samples <	detection ]i	mit	1
1-chloro-4-nitrobenzene	238	0	All samples <	detection li	mít	1
p-chlorotoluene	17/158	11	1.13-324	14.7		1

TABLE 40 (continued)

	No. of	ī	Concentrations Positive, i.e.	s for Sample:	Testing	
	s Judges	Occur-		ry wt.)	μ9/L	
Chemical	tested	rence	Range	Hedian	Range	Ref.
		Monocy	clic Aromatics (	ont 'd.)		
. 2-dichlorobenzene	101/215	47	0.0229-809	0.645		1
	71/437	16			3-1,319	2
,3-dichlorobenzene	117/215	54	0.0245-1,650	1.76		1
	40/437	9			14-1,900	2
.4-dichlorobenzene	141/215	66	C.0402-633	2.02		1
,	74/437	17		_,	2-12,000	2
-chloro-3,4-dinitrobenz	rene 0/238	0	All samples < o	detection lim	ni t	1
,4-dinitrotoluene	238	0	All samples < c	detection lin	nit	1
•	431	0	All samples Co	detection lim	ni t	2
,6-di ni troto luene	238	0	All samples < c	ietection lin	nit	1
•	431	0	All samples < o	ietection lia	ni t	2
t hy l benzene	220	6	1.22-65.5	19.8		1
•	4 3 6	63			1-4,200	2
exachlorobenzene	102/237	43	0.000188-26.2	0.018		1
	7/437	2			28-780	2
i trobenzene	0/238	0	All samples < c	detection lin	nit'	1
	0/431	0	All samples ( o	detection lim	nit	2
entach) oronitrobenzene	238	0	All samples ( o	detection lic	ni t	1
tyrene	219	10	1.53-5,850	26.6		1
-	4 37	13			2-8,300	2
oluene	434	94			1-427,000	2
,2,3-trichlorobenzene	215	37	0.00278-152	0.0667		1

TABLE 40 (continued)

	No. of	1	Concentrations ( Positive, i.e.,	> Detection	on Limits	
Chemical	sludges tested	Occur- rence	mg/kg (dry kange	/ wt.) Median	μg/L Range	Ref.
			clic Aromatics (con	nt'd.)		
,2,4-trichlorobenzene	217	57	0.00551-51.2	0.274		1
1,3,5-trichlarobenzeng	217	33	0.00502-39.7	0.0632		1
,2,3,4-tetrachlorobenzene	238	0	All samples < det	tection li	nit	1
,2,3,5-tetrachlorobenzene	738	0	All samples < det	tection lim	nit	1
,2,4,5-tetrachlorobenzene	238	0	All samples < det	tection lim	ni t	1
			iclear Aromatics (P)	AH)		
-chì oronaphthal ene	437	0.2			1,600	2
aphthalene	236	50	0.0554-6,610	30.3		1
	437	34			1-5,200	2
cenaphthene	437	5			6-4,600	2
luorene	437	6			1-1,300	2
henanthrene	437	53			1-10,100	2
nthracene	437	48			1-10,100	2
luoranthene	437	44			1-9,930	2
	12	100	0.34-11.4	2.06 <sup>†</sup>		10
yrene	437	53			1-1,700	2
hrysene	436	31			1-1,500	2
2,3-o-phenylenepyrene	437	2			17-102	2
• • •	12	100	0.06-6.86	0.88		10
lenzo(a)anthracene	437	27			1-1,500	2

TABLE 40 (continued)

	No. uf	<u> </u>		ns for Sample: e., > Detection		
Chenical	sludges tested	Occur- rence		(dry wt.) Median	ug/L Range	Ref.
		Polyni	clear Aromatics	(PAH) (cont'd	1.)	
Benzo (a) pyrene	437 12	5 100	0.12-9.14	0.88 <sup>†</sup>	1-490	2 10
3,4-t_nzofluoranthene	437 12	11 100	0.06-9.14	1.47	1-2,400	2 10
11,12-benzofluoranthene	438 12	8 100	0.06-4.57	0.49 <sup>†</sup>	1-379	2 10
Acenaphthylene	4 37	1			24-320	2
1,12-benzoperylene	437 12	2 100	0.06-9.14	0.65	12-133	2 10
1,2,5,6-dibenzanthracene	437	0.4			12-50	2
Isophorone	431	0	All samples <	detection lim	nit	2
		<u>Haloge</u>	nated Biphenyls			
PCBs (Arochlor 1248)	431	0	All samples <	detection lin	ni t	2
(Arochlor 1254)	107 431	39 0	0.0667-1,960 All samples (	5.35 detection lim	n <b>i t</b>	1 2
(Arochlor 1260)	111 431	58 0	0.0468-433 All samples <	4.18 detection lim	ni t	1 2
(decachlorobiphenyls) (ref.std.unspecified)	74 9	100 100	0.11-2.9 0.36-7.60	0.99 1.20		7 8
(Arochlor 1016, 1221, 1232, 1242)	431 ea.	ū	All samples <	detection lim	nit	2
PBB (polybrominated biphenyl)	210	0	All samples <	detection lim	ní t	1

TABLE 40 (continued)

Chenical	No. of sludges tested	% Occur- rence		ons for Sample: .e., > Detection (ary wt.) Median		Ref.
Chlorinated D	ibenzo-p-d	lioxins (CD)	Ds) and Chlori	nated Dibenzofu	rans (CDFs)§	
Tetra-CDDs	2 2	(1) (2)	0.138 and	7.2 µg/ 0.222 µg/kg	kg	9 11
Penta-CDOs	1	(0)	Sample < do	etection limit		9
Hexa-CDDs	2 2	(2) (2)	0.3 and 2. 1.41 and 1			9 11
Hepta-CDOs	2 2	(2) (2)	0.9 and 6. 9.4 and 7.			9 11
Octa-CDDs	2 2	(2) (2)	7.6 and 60 50 and 60			9 11
Tetra-CDFs	1	(0)	Sample < d	etection limit		9
Penta-CDFs	2	(0)	Sample < d	etection limit		9
Hexa-CDFs	2	(0)	Sample < d	etection limit		9
Hepta-CDFs	2	(0)	Sample < d	etection limit		9
2,3,7,8-TCDD -	431	0	All sample	s < detection l	imit	2
	На	logenated A	liphatics (Sho	rt Chain)		
Acrolein	431	0	All sampl	es < detection	limit	2
Acrylonitrile	155 436	61 1	0.0363-82	1.04	5-290	1 2
bis(2-chloroethoxy)methane	431	0	All sampl	es < detection	limit	2

TABLE 40 (continued)

Ch eni ca l	No. of	% Occur-	Concentrations for Samples > Detection Limits mg/kg (dry wt.)			ting
	sludges tested	rence	Range	Medi an	Range	Ref
	Halogena	ited Aliphat	ics (Short Chain)	(cont'd.)		
bis(chloromethyl)ether	431	0	All samples < d	etection 1	imi t	2
bis(2-chloramethyl)ether	431	0	u			2
bis(2-chloroisopropyl)ether	431	0	All samples ( de	etection 1	imit	2
Carbon tetrachloride	436	4			5-3,030	2
Chlorodibramomethane	435	2			10-75	2
Chloroethane	436	7			5-71,000	2
2-chloroethyl vinyl ether	431	0	W			2
Chlorofona	436	24			1-366	2
Dichlorobromomethane	436	6			3-260	2
Dichlorodifluoromethane	436	9			2-4,300	2
1,1-dichloroethane	436	34			1-2,880	2
1,2-dichloroethane	436	10			1-10,000	2
1,1-dichloroethylene	436	3			1-14,000	2
l,2-dichloropropane	157 435	70 6	0.00243-66.0	0.464	1-103	1 2
1,3-dichloropropane	158	26	0.209-309	3.08		1
1,3-dichloropropene	157	80	0.00203-1,230	3.42		1

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TABLE 40 (continued)

Chemi cal	No. of	% Occur-		Concentrations for Samples Te-  > Detection Limits  mg/kg (dry wt.) ug/L			
Circiii Cu i	sludges tested	rence	Range	Median	μg/L Range	t Ref.	
	Halogena		ics (Short Chain				
Hexachloro-1,3-butadiene	217 437	47 0.2	<10 <sup>-4</sup> -3.74	0.0355	2,700	1 2	
Hexachl oroethane	216 431	61 0	0.00036-61.5 All samples <	0.0199 detection lim	ni t	1 2	
Methyl bromide	431	0			33-30,000	2	
Methyl chloride	4 36	6			12-6,100	2	
Methylene chloride	436	0.4			5-19	2	
Pentachloroethane	199	28	0.00025-9.22	0.030		1	
1,1,2,2-tetrachloroethane	434	15			1-3,040	2	
Tetrachloroethylene	128 436	73 40	<10 <sup>-5</sup> -0.122	0.00052	1-2,800	1 2	
Tribromomethane	436	0.2			5	2	
1,1,1-trichloroethane	436	19			1-10,900	. 2	
1,1,2-trichlorocthane	434	4			1-2,100	2	
1,2-trans-dichloroethylene	436	60			1-96,000	2	
1,2,3-trichloropropane	141 ·	48	0.00459-19.5	0.352		1	
1,2,3-trichleropropene	137	48	/ <10 <sup>-4</sup> -167	1.14		1	
Trichloroethylene	. 432	. 54			1-32,700	2	
Trichlorofluoromethane	436	5			2-113	2	
vinyl chloride	435	8			8-62,000	2	

TABLE 40 (continued)

Chemi ca l	No. of	1 Occur-		r Samples T Limits μg/L	s resumy	
	sludges tested	rence	Range	Median	Range	Ref
		Triaryl Ph	osphate Esters			
Cresyldiphenyl phosphate.	238	3	0.607-179	18.9		1
Tricresyl phosphate	235	69	0.069-1,650	6.85		1
Trixylyl phosphate	236	68	0.0273-2,420	7.11		1
		Aromatic a	nd Alkyl Amines S			
Be nz idi ne	238	0.4		12.7		1
	431	0	All samples < d	etection lim	if t	2
3,4-dichloroaniline	238	0	All samples < d	etection lim	if t	1
3,3'-dichlorobenzidine	238	0	All samples < d	etection lim	iit	1
	431	0	All samples ( d	etection lim	ii t	2
p-nitroaniline	238	0	Ali samples < d	etection lim	it	1
N-nitrosodimethylamine	431	0	All samples < d	etection lim	ı <b>i</b> t	2
	15 11	93 82	0.6-53 µg/kg	5.3 µg/		3
	11	92	0.5-93 µg/kg	2.5 µg/	Kg,	4
N-nitrosodiethylamine	15	27	0.6-3.8 µg/kg	0.9 µg/	-	3
	11	54	0.9-12 µg/kg	1.7 µg/	Kg	4
N-nitrosodibutylamine	15	0	All samples <			3
	11	0	All samples <	detection li	mit	4
N-nitros op iperidi ne	15	0	All samples (	detection li	mít	3
••	11	9	-	2.9 µg/	kg	. 4
N-nitrosopyrrolidine	- 15 11	13	1.3 and 4.2 µg 1.7 and 2.8 µg			3

TABLE 40 (continued)

Chemi cal	No. of	% Occur-		> Detection	r Samples Te Limits μg/L	sting
	sludges tested	renc e	Range	Median	Range	Ref
		Aromatic and	d Alkyl Amines (	cont'd.)		
N-nitros omo rpholine	15 11	20 54	2.0-9.2 µg/kg 1.3-2.9 µg/kg	2.9 μg/ 1.8 μg/		3 4
N-nitrosodiphenylamine	431	0	All samples <	detection li	mit	2
N-nitrosodi-n-propylamine	431	0	All samples <	detection li	mit	2
			Phenols			
p-chloro-m-cresol	438	1			12-35	2
o-ch I orophenol	231 4 38	9 2	0.0766-90.0	3.6	11-72	1 2
m-chlorophenol	231	7	0.123-93.3	0.891		ı
p-chlorophenol	231	9	0.0277-90.0	3.28		1
o-cresol	231	6	0.177-183	2.05		1
2,4-dichlorophenol	230 438	7 2	0.209-203	4.75	14-298	1 2
2,4-dimethylphenol	231 431	18 0	0.0899-86.7 All samples <	2.19 detection li	mit	1 2
4,6-dinitro-o-cresol	228 431	9 .0 -	0.202-187 All samples <	2.34 detection li	mit	1 2
2,4-dinitrophenol	228 431	30 0	0.153~500 All samples (	4.62 detection li	mit	1 2
Nydroquinone	229	27	0.138-223	2.55		1

TABLE 40 (continued)

	No. of	ĭ	Conc	entrations for > Detection t		ting
Chemical	No. of sludges	Occur-	mg/kg (dry		μg/L	
	tested	rence	Range	Median	Range	Ref
			Phenols (cont'd	).		
2-nitrophenol	431	0	All samples <	detection lim	nit	2
4-nitrophenol	431	0	All samples <	detection lim	ni t	2
Pentachlorophenol	223	70	0.172-8,490	5.00		1
,	4 78	14	•		10-10,500	2
Phenal	229	78	0.0166-288	2.00		1
	4 38	50		-	5-17,000	2
2,4,6-trichlorophenol	223	30	0.195-1,330	4.81		1
	438	0.4			11-16	2
	Chlori	nated Pesti	cides and Hydroc	arbons		
Aldrin	223	0	All samples <	detection lim	nit	1
	431	0		detection lim	nit	2
	74	100	0.05-0.64	0.08		7
Chlordane*	431	0	All samples (	detection lis	ni t	2
	74	100	0.46-12	2.75		7
Dieldrin	221	28	0.000377-64.7			1
	431	0		detection lim	nit	2
	40	?	(0.01-1.26	0.26		2 5 6
	14	93	0.04-2.2	0.16		
	74	100	0.05-0.81	0.11		7
Endrin	223	0	All samples	< detection li	imit	1
	431	0				2
	74	100	0.11-0.17	0.14		7

<u>ي</u>

TABLE 40 (continued)

Cheni cal	No. of Occur-		Concentrations for Samples To  > Detection Limits  mg/kg (dry wt.)	esting
	sludges tested	rence	Range Median Range	Ref.
	Chlori	nated Pestic	ides and Hydrocarbons (contid.)	
Endrin aldehyde	431	0	All samples < detection limit	2
p,p'-000	221	48	0.00114~84.1 0.363	1
	431	0	All samples < detection limit	2
p,p'-DDE	219	92	0.00118-564 1.14	1
	443	0.2	10,000	2
	40	100	0.01-0.49 0.02	5
	74	0	All samples < 0.05	7
p,p'-DDT	219	95	<10 <sup>-4</sup> -135 0.211	1
	4 3 1	0	All samples < detection limit	2
	74	100	0.06-0.14 0.09	7
Heptach) or	431	0	All samples < detection limit	2
Heptachlor epoxide	74	100	0.05-0.55 0.13	7
•	431	0	All samples < detection limit	2
Lindane(y-BHC)	221	17	0.00059-12.5 0.0746	1
•	431	0	All samples < detection limit	2
	40	?	<0.01-0.93 0.18	5
	74	100	0.05-0.22 0.11	7
Methoxychlor	223	0	All samples < detection limit	1
2,4-0	223	25	0.000554-7:34 0.122	1
Hexachlorocyclopentadian	e 431	0	All samples < detection limit	2
Toxaphene	431	0	All samples < detection limit	2

TABLE 40 (continued)

		1	Concentrations for Samples > Detection Limits	Testing
Chemi cal	No. of sludges	Occur-	mg/kg (dry wt.) μή/l	
	tested	rence	Range Median Range	Ref.
	<u>Chlori</u>	nated Pestic	ides and Hydrocarbons (cont'd.)	
a-endosulfan	431	0	All samples < detection limit	2
β-endosulfan	431	0	All samples < detection limit	2
Engosulfan sulfate	431	0	All samples < detection limit	2
a-BKC	431	0	All samples < detection limit	2
в-вис	431	0	All samples < detection limit	2
6-BHC	431	0	All samples < detection limit	2
		Miscellane	ous Compounds	
1,2-diphenylhydrazine	431	0	All samples < detection limit	2
4-chlorophenyl phenyl ether	431	0	All samples < detection limit	2
4-branophenyl phenyl ather	431	0	All samples < detection limit	2
Mercaptobenzothiazole	238	0	All samples < detection limit	1
Biphenyl	236	33	0.0437-1,730 8.61	1

<sup>†</sup>A concentration range was given in Reference 10 for each of the 12 sludges tested. The concentration for each PAH organic was obtained by taking the average of the high and low values to get an average concentration for each of the 12 sludges. These 12 averages were then used to report the range and median value in Table 40.

\*"% occurrence" times the total "no. of sludges tested" equals the number of samples testing positive, i.e., having a concentration greater than the detection limit.

\*Under "% occurrence", Values given in parentheses are the number of samples which had a detectable concentration rather than a "percent" value.

§Note: Concentrations (on dry weight basis) for dioxins and furans and several alkyl amines are in µg/kg rather than mg/kg as for all other organics.

## ttReferences:

- 1. Jacobs and Zabik, 1983. (Various sludges from 204 Michigan WWTPs)
- 2. Burns and Roe, 1982. (Primary, secondary, and combined sludges from 40 POTWs)
- 3. Mumma et al., 1934. (Various sludges from 31 American cities)
- 4. Mumma et al., 1983. (Various sludges from 24 New York communities)
- 5. McIntyre and Lester, 1982. (Various sludges from 40 WWTPs in England)
- 6. Furr et al., 1976. (Various sludges from 14 American cities)
- 7. Clevenger et al., 1983. (/arious sludges from 74 Missouri WWTPs)
- 8. Diercxsers and Tarradellas, 1983. (Various sludges from 9 Switzerland WWTPs)
- 9. Weerasinghe et al., 1985. (Two sludges from Syracuse, NY and Sodus, NY)
- 10. McIntyre et al., 1981. (Various sludges from 12 United Kingdom WWTPs)
- 11. Lamparski et al., 1984. (Two samples of Milorganite, one each produced in 1981 and 1982)

TABLE 41. SUMMARY COMPARING THE NUMBER OF ORGANIC CHEMICALS TESTED TO THE NUMBER OF ORGANICS NOT DETECTED IN SEWAGE SLUDGES OR FOUND IN 10, 50 OR 90% OF THE SLUDGES.

	No. of No. of organics organic undetected in chemicals all samples			No. of organic chemicals w occurrence:			
Reference <sup>†</sup>	tested	tested tested		>50%	> 90%		
			Phthalate Ester	rs			
1 (204 WWTPs) 2 (40 POTWs)	6 6	0	6	3	0		
2 (40 70185)		•	Monocyclic Arm	natics	•		
		•					
1 (204 WWTPs) 2 (40 POTWs)	23 12	12 3	8 7	3 3	0 1		
2 (40 POINS)	12	3	,	3	1		
			Polynuclear Arc	omatics (PAH	1		
1 (204 WVTPs)	1	0	1	1	0		
2 (40 POTWs)	18	1	9	2	0		
10 (12 WWTPs)	6	0	6	6	6		
			Halogenated Bir	henyls			
1 (204 WWTPs)	3	1	2	1	0		
2 (40 POTWs)	3	3	0	Ö	0		
3 (31 WWTPs)	1	0	1	1	1		
4 (24 WVTPs)	1	0 0	1	į,	1		
5 (40 NWTPs) 6 (14 WWTPs)	1	0	1	1	1		
6 (14 WWIPS) 7 (74 WWTPS)	1	0	i	i	1		
8 (9 WWTPs)	ī	ō	1	ī	i		
			Dioxins and Fur	rans			
2 (40 POTWs)	1	1	0	0	0		
1 (2 WHTPS) 11 (2 samples of	-	enough samples	tested to sugge	est % occurr	-		
11 (2 samples of Miloryanite)	•	enough samples			•		

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TABLE 41 (continued) No. of No. of organics No. of organic chemicals with organic undetected in occurrence: chemicals all samples Reference<sup>†</sup> tested tested >90% >10% >50% Halogenated Aliphatics 1 (204 WWTPs) 2 (40 POTWs) 10 0 0 32 7 0 Triaryl Phosphate Esters 2 1 (204 WWTPs) 3 0 0 Aromatic and Alkyl Amines 1 (204 WWTPs) 3 5 0 2 (40 POTWs) 5 0 2 3 (15 WMPs) 6 1 4 (11 WWTPs) 0 Phenols 1 (204 WWTPs) 2 (40 POTWs) 12 0 5 0 11 0 Chlorinated Pesticides and Hydrocarbons 1 (204 WWTPs) 9 2 (40 POTWs) 19 18 0 3 0 5 (40 NWTPs) 6 (14 WWIPS) 1 0 7 7 (74 HITPS) 1 Miscellaneous Compounds 1 (204 HHTPs) 2 0 2 (40 POTHS) 3 3 0 TOTALS: 219 70 102 53 26

<sup>†</sup>Reference number used refers to the same references as used in Table 40. Number of wastewater treatment plants tested are given in parentheses.

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TABLE 42. SUMMARY SHOWING THE DISTRIBUTION OF MEDIAN DRY MATTER CONCENTRATIONS FOR DATA REPORTED IN TABLE 40.\*

	No. of organic	No. of organic chemicals tested having median concentrations in sludges (mg/kg, dry wt. basis):					
henical group	chemicals tested	ND <sup>†</sup>	α `	1-10	10-100	>100	
Phthalate esters	6	0	0	1	4	1	
Monocyclic aromatics	23	12	5	2	4	0	
Polynuclear aromatics (PAH)	7 %	0	4	2	1	0	
Halogenated biphenyls	9	1	3	5	0	0	
Dioxins and furans					ions reporte mg/kg ur 10		
Halogenated aliphatics	10	0	6	4	0	0	
Triaryl phosphate esters	3	0	0	2	1	0	
Arcmatic & alkyl amines	· 16	6	9	0	1	0	
Phenols	12	0	1	11	0	0	
Chlorinated pesticides and hydrocarbons	21	4	14	3	0	ó	
Mi scel laneou s	2	1	0	1	0	0	
TOTALS:	110	24	42	31	11	1	

<sup>\*</sup>Summary does not include data from Burns and Roe (1982) which was reported on a wet basis without median values provided. Also note that median values used are only for those samples having detectable concentrations and are not true median values, which would be lower if all "ND" samples were included as zeroes. Waste Water Treatment Plant (WWTP); Publicly Owned Treatment Plant (POTW).

<sup>\*</sup>ND = organic was "not detected" in any sludge samples tested

TABLE 43. GUIDELINES USED BY ONE FOOD PROCESSING COMPANY FOR INTERPRETING THE SIGNIFICANCE OF RESIDUES IN SOILS BEING CONSIDERED FOR GROWING ROOT CROPS.

Significance	Range o	f soil residues (mg/) DDT	(g) Diuron*
Suitable for plantingt	0-0.1	0.75	0.3
May be planted, but crop must be analyzed before acceptance	0.1-0.2	0.75-1.5	0.3-0.5
Do not plant	over 0.2	over 1.5	over 0.5

<sup>\*</sup>Regardless of residues present, beets and carrots must not be planted in soil which has received (a) an improper application of diuron or (b) an application of diuron for which the minimum treatment-toplanting interval has not expired.

tPlant carrots in least-contaminated soil.

TABLE 44. ILLUSTRATIVE RANGE OF DECOMPOSITION HALF-LIFE FOR OFGANIC COMPOUNDS.\*

Compound	Approximate half-life				
Aminoanthroquinone dyes	100-2,200 days				
Anthracene	110-180 days				
Benzo(a)pyrene	60-420 days				
Di-n-butylphthalate ester	80-180 days				
Nonionic surfactants	300-600 days				
2,4-methyaniline	1.5 days				
n-Nitrosodiethylamine	40 days				
Pheno!	1.3 days				
Pyrocatechin	12 hours				
Cellulose	35 days				
Acetic acid	5-8 days				
Hydroquinone	12 hours				

<sup>\*</sup>From Overcash (1983), p. 211.

TABLE 45. RELATIVE PERSISTENCE AND INITIAL DEGRADATIVE REACTIONS OF NINE MAJOR ORGANIC CHEMICAL CLASSES.\*

Chemical class	Persistence	Initial degradative process
Carbamates	2-8 weeks	Ester hydrolysis
Aliphatic acids	3-10 weeks	Dehalogenation
Nitriles	4 months	Reduction
Phenoxyalkanoates	1-5 months	Dealkylation, ring hydroxylation or oxidation
Toluidine	6 months	Dealkylation (aerobic) or reduction (anaerobic)
Amides	2-10 months	Dealkylation
Benzoic acids	3-12 months	Dehalogenation or decarboxylation
Ureas	4-10 months	Dealkylation
Triazines	3-18 months	Dealkylation or dehalogenation

<sup>\*</sup>From Kaufman (1983), p. 119.

## Table 46. ASSUMPTIONS/VALUES USED FOR METRO ANALYSIS (MUNGER, 1984).

Sludge contains: 1.1 mg/kg DW PCBs (Metro sludge)

2.6 mg/kg DW B(a)P (Metro sludge)

Application rate: 45 mt/ha for silviculture

Estimated soil

concentration: Calculated assuming even mixing in top 15 cm

Risk level used

as benchmark: 10<sup>-5</sup> (Values from Munger, 1984 were divided by

10 to give a  $10^{-6}$  risk level for values in

Tables 45 and 46.)

Normal daily dietary intake: 8,700 ng PCBs/day

160 - 1,600 ng B(a)P/day

Consumption equivalent to a

lifetime cancer risk of 10<sup>-5</sup>: 204 ng PCBs/day

61 ng B(a)P/day

TABLE 47. METRO ASSESSMENT OF LIFETIME CANCER RISK FOR PCB.\*

Estimated quantities of environmental compartments which can be consumed on a daily basis without exceeding a lifetime cancer risk of  $10^{-6}$  based on PCB concentrations

	Months after application					
Environmental compartment	0	3	6	12	24	
Sludge-soil (g/day)	0.8	0.8	0.9	1.0	1.0	
Control soil (g/day)			<u>&lt;</u> 2.0 -			
Surface water (liters/day)			<u>&lt;</u> 2.0 -			
Control water (liters/day)			<u>&lt;</u> 2.0 -			
Edible plants (g/day)	•		20	-100		
Control plants (g/day)			<u>&lt;</u> 200 -			
Deer fat (g/day)	•			2		
Control deer fat (g/day)		U	nknown			
Groundwater (liters/day)			<u> </u>			

<sup>\*</sup>Munger (1985). See Table 46 for assumptions used.

TABLE 48. METRO ASSESSMENT OF LIFETIME CANCER RISK FOR B(a)P.\*

Estimated quantities of environmental compartments which can be consumed on a daily basis without exceeding a lifetime cancer risk of 10<sup>-6</sup> based on B(a)P concentrations

	Months after application					
Environmental compartment	0	3	6	12	24	
Sludge-soil (g/day)	0.1	0.2	0.6	0.6-6	0.6-6	
Control soil (g/day)	y)					
Edible plants (g/day)	1-10	2-20	30-300	30-300	30-300	
Control plants (g/day)			30-300			
Animal tissue			unknown			
Groundwater and control groundwater (liters/day)			- <u>&lt;</u> 0.6 -			

<sup>\*</sup>Munger (1985). See Table 46 for assumptions used.

TABLE 49. DEFINITIONS FOR "RELATIVE TOXICITY" CATEGORIES AS USED BY NAYLOR AND LOEHR (1982a).

Ratings	Acute oral LD <sub>50</sub> mg/kg	Relative toxicity				
Supertoxic	<5	6	a taste to 7 drops			
Extremely toxic	5-50	5	7 drops to a teaspoon			
Very toxic	50-500	4	1 teaspoon to 1 ounce			
Moderately toxic	500-5,000	3	1 ounce to 1 pint (1 pound)			
Slightly toxic	5,000-15,000	2	1 point to 1 quart (2 pounds			
Practically non-toxic	>15,000	1	more than 1 quart			

TABLE 50. EXAMPLES OF CHEMICALS COMMONLY CONSUMED OR USED AND THEIR TOXICITY RATINGS (NAYLOR AND LOEHR, 1982a).

	Acute oral LD50 *	
Chemical	for rats, mg/kg	Toxicity rating'
Sodium chloride	3,000	3
Sugar	25,800	1
Aspirin	1,000	3
Nicotine	-,	4
Oxalic acid (present in chard, spinach,		
rhubarb leaves, etc.)	375	4
Caffeine	192	4
Ethyl alcohol Safrole (80%) of oil	14,000	2
of sassafras)	-	5
Gasoline, kerosene	-	3
Antifreeze	-	3
Strychnine	-	3 3 6 3
Cayenne pepper	-	3
Laundry bleaches	-	3-4
Aftershave lotions	-	3 1
Vanilla and lemon extract	-	1
Bouncing putty	-	3

<sup>\*</sup>Lewis and Tatkin (1979)

<sup>†</sup>For interpretation, see Table 49

TABLE 51. TOXICITIES AND APPLICATION RATES FOR SEVERAL PESTICIDES (NAYLOR AND LOEHR, 1982a).

Pesticide	Acute oral LD50 for rats* mg/kg	Relative toxicity <sup>†</sup>		Recommended appli- cation rate of active ingredient to soil, tbs/acre
Methyl parathion	6	5	Oat and	0.25
Parathion	2	6	wheat	0.25 to 0.38
Malathion	885	3	insect	1.0
Diazinon	76	4	control	1.0
Dilox	945	3 4		0.5 to 1.0
2,4-D	375	4	Forages,	1.0
Methoxychlor + Malathion	5,000	2	alfalfa	1.5 + 1.5
Diazinon	76	4	Corn borer	1.0 to 2.0
Sevin	250	4	Corn borer	1.0 to 2.0
Disyston	5	6	Corn root	1.0
Dasinit	2 3	6	worm	0.75 to 1.0
Dyfonate '	3	6	WOLIL	0.75 to 1.0
Lorsban	145	4	worm	0.75 to 1.0
Phosdrin	4	6	Corn leaf	0.125
Maneb 80	6,750	2	aphid Herbicide:	
Sencor	2,200	3 3	tomato, potat	o up to 2.0
Systox	1,700	3	Herbicide: soybeans, Systemic insecticide	0.38

<sup>\*</sup>Lewis and Tatkin (1979)

<sup>†</sup>See Table 49 for explanation

<sup>‡</sup>New York State Coll. of Agric. & Life Sci. (1982a,b)

TABLE 52. TOXICITIES, SLUDGE CONCENTRATIONS, AND PROJECTED APPLICATION LOADINGS FOR SELECTED PRIORITY POLLUTANT ORGANICS (NAYLOR AND LOEHR, 1982a).

			No.times				sludges	Projected	
	Acute oral		detected in combined	<u></u>	l, wet	mg/	kg, dry	application rate9 kg/ha, dry	
Chemical	rating mg/kg*	Toxicty ratingt	sludge	median	range	median	range	median	ranye
bis-2-ethylexyl phthalate	31000	1	13	3806	157-11257	109	4.1-273	1.2	0.053-2.1
chloroethane	volatile	-	2	1259	517- 2000	19	14.5- 24	0.17	0.16-0.17
1,2-trans-dichloroethylene	volatile	•	11	744	42-54993	21	0.72-865	0.24	0.009-8.4
toluene	5000	2	12	722	54-26857	15	1.4-705	0.16	0.018-1.3
butylbenzyl phthalate	3160	3	11	577	1-17725	15	0.52-210	0.11	0.0063-1.4
2-chloronaphthalene	2078	3	1	400	400	4.7	4.7	0.03	0.03
hexachlorobutadiene	90	4	2	338	10- 675	4.3	0.52-8	0.03	0.0063-0.05
phenanthrene	700	3	12	278	34- 1565	7.4	0.89-44	0.05	0.009-0.53
carbon tetrachloride	2800	3	1	270	2/0	4.2	4.2	0.041	0.041
vinyl chloride	500	3	3	250	145- 3292	5.7	3-110	0.064	0.02-1.3
dibenzo (a.h) anthracene	-	-	1	250	25	13	13	0.16	0.16
1.1.2-trichloroethane	1140	3	2	222	3- 441	3.5	0.036-6.9	0.034	0.0002-0.068
anthracene	-	-	13	272	34- 1565	7.6	0.89-44	0.050	0.009-0.53
naphthalene	1780	3	9	238	23- 3100	7.5	.9-70	0.070	0.01-0.59
ethylbenzene	3500	3	12	248	45- 2100	5.5	1.0-51	0.063	0.013-0.38
di-n-butylphthalate	1200	3	12	184	10- 1045	3.5	0.32-17	0.047	0.003-0.21
phenol	414	4	11	123	27- 4310	4.2	0.9-113	0.032	0.0011-1.5
methylene chloride	167	4	10	89	<b>5- 1</b> 055	2.5	0.06-30	0.022	0.0004-0.97
pyrene	-	•	12	125	10- 734	2.5	0.33-18	0.024	0.04-0.22
chrysene	_	-	9	85	15- 750	2.0	0.25-13	0.022	0.0024-0.16
fluoranthene	2000	3	10	90	10- 600	1.8	0.35-7.1	0.016	0.0024-0.05
benzene	1400	3	11	16	2- 401	0.32	0.053-11.3	0.0027	0.0007-0.13
tetrachloroethylene	8100	2	11	14	1- 1601	6.38	0.024-42	0.0035	0.0002-0.54
trichloroethylene	4920	3	10	57	2- 1927	0.98	0.048-44	0.0125	0.00036-0.52

<sup>\*</sup>National Academy of Sciences (1972)

§Feiler (19d0)

<sup>†</sup>See Table 49 for interpretation

TABLE 53. TIMES AND AMOUNTS OF SLUDGE WHICH MUST BE INGESTED BY THE RAT OR COW TO REACH LD<sub>50</sub> DOSES OF THREE SLUDGE ORGANICS (NAYLOR AND LOEHR, 1982a).

Priority pollutant	Max.concn. in sludges mg/kg*	LD <sub>50</sub> dose, mg/kg*	Toxicity rating*	Example animal	Typical animal wt, kg	LD <sub>50</sub> dose	Amount of sludge equal to LD <sub>50</sub> dose of chemical, kg	Time to consume LD50 dose of chemical, yrs
hexachloro- butadiene	8	90	4	rat cow	0.5 500	45 mg 45 g	5.6 5600	7.7 (2800 days) 6.2
dis-2- ethylhexyl phthalate	273	31000	1	rat	0.5	15.5 g	57	78
1,1,2-tri- chloroethane	6.9	1140	3	cow rat cow	500 0.5 500	15.5 kg 570 mg 570 g	57000 83 83000	62 113 91

<sup>\*</sup>From Table 50.

Daily food intake: rat = 20 g/day (6); cow = kg/day, with sewage sludge (dry basis) intake equivalent to 10 percent by weight of total diet

Because of the length of exposure period to consume an LD50 dose of chemical, health effects observed are not necessarily equivalent to those observed where the dosage is ingested within more conventional LD50 test exposure periods of several days or less.

Life expectancy of rat = 700 to 800 days (2 to 2.5 years) and of a lactating cow = 5 to 10 years

TABLE 54. EVALUATION OF POTENTIAL INTAKE OF THREE SLUDGE ORGANICS DUE TO SLUDGE OR SOIL WITH SLUDGE INGESTED BY A "PICA" CHILD OR A COW.

		LD <sub>50</sub> dose	Soil con- sump- tion g/d	Time to consume LD <sub>50</sub> dose of priority pollutant, yrst		(D <sub>2</sub> )	(D <sub>T</sub> )	C-5-4 :
	Weight, kg			Soil w/sludge	_	Daily intake from soil, g	daily dose, g§	Safety factor <sup>5</sup> D <sub>T</sub> /D <sub>2</sub>
Child Cow	20 500	1.8 g 45 g		1 x 10§ 1 x 10‡	41 10	4 x 10 <sup>-7</sup> 4 x 10 <sup>-5</sup>	1.8 x 10 <sup>-5</sup>	45
	Ma	aximum c				LD <sub>50</sub> = 31000 m soil = 1.0 mg/k		273 mg/k
Child Cow	20 ( 500 :	0.62 kg 15.5 kg	15 1500	1 x 10 <sup>5</sup> 1 x 10 <sup>4</sup>	415 104	1.5 x 10 <sup>-5</sup> 1.5 x 10 <sup>-3</sup>	6.2 x 10 <sup>-3</sup>	415
	Maximur	l n concen	.,1,2-tr itration	richloroetha of chemica	ane LD50 = 1 al in soil =	.140 mg/kg = 0.034 mg/kg,	in sludge = 6.	9 mg/kg
Child Cow				1 x 10 <sup>5</sup>	604 151	5.1 x 10 <sup>-7</sup> 5.1 x 10 <sup>-5</sup>	2.3 x 10 <sup>-4</sup>	450

<sup>\*</sup>Adapted from Naylor and Loehr (1982a)

tFrom Table 53

<sup>#</sup>Estimated LD<sub>50</sub> dose = LD<sub>50</sub> mg/kg x body wt. kg §For humans, acceptable daily dose of toxic pollutants (D<sub>T</sub>) =  $10^{-5}$  x LD<sub>50</sub> (safety factor of  $10^{5}$ ).

TABLE 55. SAFETY FACTORS FOR INGESTING SOIL CONTAINING PESTICIDE AND SLUDGE ORGANICS (CONNOR, 1984).

	Application rate (kg/ha)	Soil concn. (µg/g)	ADI (µg/day)	Safety Low diet	factor High diet
Pesticide					
2,4-D Diazinon Malathion Methoxychlor Parathion Methyl parathion	1.1 1.1 1.7 .2944	0.5 0.5 0.5 0.9 .1522	21,000 140 1,400 7,000 350 70	2,800 19 190 420 110 33	300 2 20 56 14 4
Sludge Bis-2-ethyl-	Concn. (ppm dry)				
hexyphthalate Toluene Ethyl benzene Di-n-butylphthalate Phenol Methylene chloride Total PAH*	109 15 5.5 3.5 4.2 2.5	0.6 0.08 0.032 0.024 0.016 0.011	42,000 29,500 1,600 88,000 100 25,000 0.4	4,700 25,000 3,300 240,000 420 150,000 0.27	500 2,700 360 26,000 45 16,000 0.03
			Potency (kg day/mg)	10 <sup>-6</sup> risk	10 <sup>-6</sup> risk
Hexachlorobutadiene Carbon tetrachloride Vinyl chloride 1,1,2-trichloroethane Benzene Tetrachloroethylene Trichloroethylene Total PAH	4.3 4.2 5.7 3.5 0.32 0.38 0.98 13	0.015 0.020 0.032 0.017 0.0014 0.0018 0.006 0.10	0.495 0.083 0.017 0.057 0.052 0.040 0.012 11.5	0.16 0.36 0.12 0.21 0.016 0.015 0.015 250	1.5 3.3 1.9 1.9 0.14 0.14 0.14 2300

Safety factors (ADI divided by daily consumption) and  $10^6$  risk calculated assuming 70 kg person ingesting equivalent of 15 g (low diet) or 139 g (high diet) or contaminated soil per day. See text for further explanation

PAH includes acenaphthene, fluoranthene, benzanthracene/chrysene, anthracene/phenanthrene, and pyrene. ADI calculated from WHO drinking water standard of 0.2  $\mu g/liter$  and assuming consumption of 2 liters of water per day.

Soil concentration calculated assuming an average sludge application rate of 15 tons/ha.

## SECTION 7

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