Water

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An Exposure and Risk Assessment for Benzo[a]pyrene and Other Polycyclic Aromatic Hydrocarbons

Volume IV. Benzo[a]pyrene, Acenaphthylene, Benz[a]anthracene, Chrysene, Dibenz[a,h]anthracene, Benzo[b]fluoranthene, Benzo[k]fluoranthene, Benzo[g,h,i]perylene,

Indeno[1,2,3-c,d]pyrene



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This report assesses the risk of exposure to polycyclic aromatic hydrocarbons (PAHs). This is Volume IV of a four-volume report, analyzing 16 PAHs; it concerns nine of benzo[a]pyrene, acenaphthylene, benz[a]anthracene, benzo[b]fluoranthene, benzo[k]fluoranthene, benzo[g,h,i]perylene, chrysene, dibenz[a,h]anthracene, indeno[1,2,3-c,d]pyrene. This study is part of a program to identify the sources of and evaluate exposure to 129 priority pollutants. The analysis is based on available information from government, industry, and technical publications assembled in June of 1981.

The assessment includes an identification of releases to the environment during production, use, or disposal of the substances. In addition, the fate of PAHs in the environment is considered; ambient levels to which various populations of humans and aquatic life are exposed are reported. Exposure levels are estimated and available data on toxicity are presented and interpreted. Information concerning all of these topics is combined in an assessment of the risks of exposure to PAHs for various subpopulations.

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Exposure	Effluents	Polycyclic	Aromatic Hydrocarbons	Chrysene		
Risk	Waste Disposal	Indeno[1,2	3-c,d]pyrene	Acenaphthylene		
Water Pollution	Food Contamination	Benzo[g,h,]perylene	PAHs		
Air Pollution	Toxic Diseases	Benzo[b]flu	oranthene	Benzo[a]pyrene		
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AN EXPOSURE AND RISK ASSESSMENT FOR BENZO[a]PYRENE AND OTHER POLYCYCLIC AROMATIC HYDROCARBONS:

VOLUME IV. BENZO[a]PYRENE, ACENAPHTHYLENE, BENZ[a]ANTHRACENE, BENZO[b]FLUORANTHENE, BENZO[k]FLUORANTHENE, BENZO[g,h,i]PERYLENE. CHRYSENE.

DIBENZ[a,h]ANTHRACENE, AND
INDENO[1,2,3-c,d]PYRENE

Вy

Joanne Perwak, Melanie Byrne, Susan Coons, Muriel Goyer and Judith Harris Arthur D. Little, Inc.

U.S. EPA Contract 68-01-6160

Patricia Cruse, Robert DeRosier, Kenneth Moss and Stephen Wendt Acurex Corporation

U.S. EPA Contract 68-01-6017

John Segna and Michael Slimak Project Managers U.S. Environmental Protection Agency

Monitoring and Data Support Division (WH-553)
Office of Water Regulations and Standards
Washington, D.C. 20460

OFFICE OF WATER REGULATIONS AND STANDARDS
OFFICE OF WATER
U.S. ENVIRONMENTAL PROTECTION AGENCY
WASHINGTON, D.C. 20460

FOREWORD

Effective regulatory action for toxic chemicals requires an understanding of the human and environmental risks associated with the manufacture, use, and disposal of the chemical. Assessment of risk requires a scientific judgment about the probability of harm to the environment resulting from known or potential environmental concentrations. The risk assessment process integrates health effects data (e.g., carcinogenicity, teratogenicity) with information on exposure. The components of exposure include an evaluation of the sources of the chemical, exposure pathways, ambient levels, and an identification of exposed populations including humans and aquatic life.

This assessment was performed as part of a program to determine the environmental risks associated with current use and disposal patterns for 65 chemicals and classes of chemicals (expanded to 129 "priority pollutants") named in the 1977 Clean Water Act. It includes an assessment of risk for humans and aquatic life and is intended to serve as a technical basis for developing the most appropriate and effective strategy for mitigating these risks.

This document is a contractors' final report. It has been extensively reviewed by the individual contractors and by the EPA at several stages of completion. Each chapter of the draft was reviewed by members of the authoring contractor's senior technical staff (e.g., toxicologists, environmental scientists) who had not previously been directly involved in the work. These individuals were selected by management to be the technical peers of the chapter authors. The chapters were comprehensively checked for uniformity in quality and content by the contractor's editorial team, which also was responsible for the production of the final report. The contractor's senior project management subsequently reviewed the final report in its entirety.

At EPA a senior staff member was responsible for guiding the contractors, reviewing the manuscripts, and soliciting comments, where appropriate, from related programs within EPA (e.g., Office of Toxic Substances, Research and Development, Air Programs, Solid and Hazardous Waste, etc.). A complete draft was summarized by the assigned EPA staff member and reviewed for technical and policy implications with the Office Director (formerly the Deputy Assistant Administrator) of Water Regulations and Standards. Subsequent revisions were included in the final report.

Michael W. Slimak, Chief Exposure Assessment Section Monitoring & Data Support Division (WH-553) Office of Water Regulations and Standards

AN EXPOSURE AND RISK ASSESSMENT FOR BENZO[a]PYRENE AND OTHER POLYCYCLIC AROMATIC HYDROCARBONS

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The Materials Balance for the BaP group PAHs (Section 3.1) was produced by Acurex Corporation, under Contract No. 68-01-6017 to the Monitoring and Data Support Division (MDSD), Office of Water Regulations and Standards (OWRS), U. S. Environmental Protection Agency. Patricia Cruse was the task manager for Acurex, Inc.; other contributors include Robert DeRosier, Kenneth Moss and Stephen Wendt. Patricia Leslie was responsible for report production on behalf of Acurex, Inc.

John Segna and Michael Slimak were the EPA project managers for this assignment.

1.0 INTRODUCTION

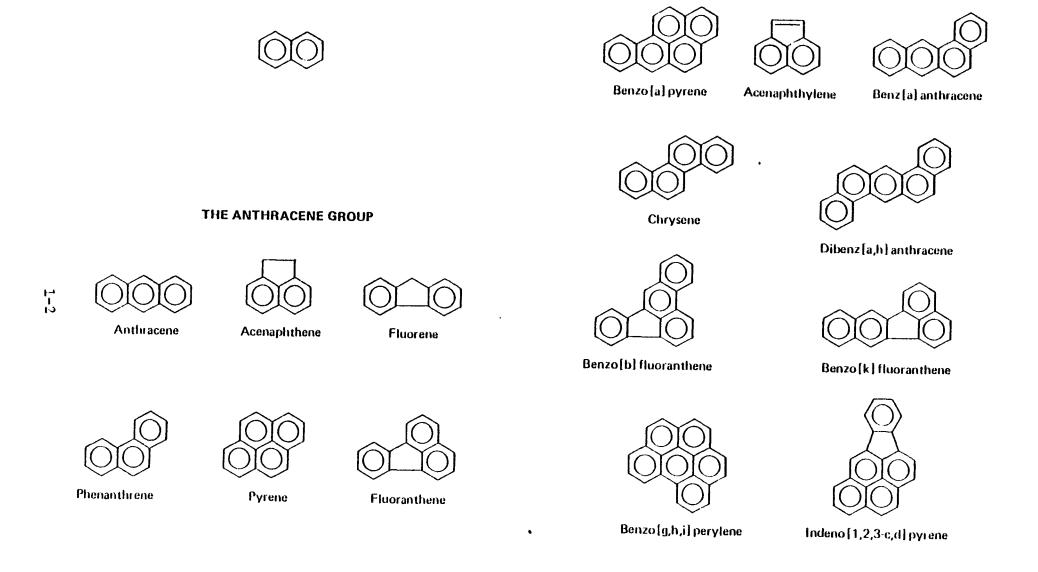
The Office of Water Regulations and Standards (OWRS), Monitoring and Data Support Division, of the U.S. Environmental Protection Agency is conducting a program to evaluate the exposure to and risk of 129 priority pollutants in the nation's environment. The risks to be evaluated include potential harm to human beings and deleterious effects on fish and other biota. The goals of the program under which this report has been prepared are to integrate information on cultural and environmental flows of specific priority pollutants, to estimate the likelihood of receptor exposure to these substances, and to evaluate the risk resulting from such exposures. The results are intended to serve as a basis for estimating the magnitude of the potential risk and developing a suitable regulatory strategy for reducing any such risk.

This report, comprised of four separate volumes, provides a summary of the available information concerning the releases, fate, distribution, effects, exposure, and potential risks of the 16 priority pollutants that are polycyclic aromatic hydrocarbons (PAHs). The chemical structures of these compounds are shown in Figure 1-1.

The number of chemicals considered in this exposure and risk assessment is appreciable. The possibility of preparing 16 separate exposure and risk assessment documents was considered and rejected because it would lead to considerable redundancy and because so little information was available on some of the individual PAHs. As an alternative, the 16 PAHs were organized at the onset of the work into three groups, as indicated in Figure 1-1.

The rationale for the organization into these three specific groups included considerations of materials balance, chemical properties related to fate and environmental pathways, and health effects, as described briefly below.

- Naphthalene is the only one of the 16 PAHs with substantial U.S. commercial production and with a significant potential for direct exposure to consumers of a commercial product (mothballs). It is significantly more volatile and more water soluble than any other PAH. It was not anticipated to have carcinogenic effects in humans.
- Anthracene, acenaphthene, fluorene, fluoranthene, phenanthrene and pyrene are all imported in rather small quantities for special commercial uses. These compounds are three- and four-ring PAHs, with moderately low volatility and water solubility. The question of their possible carcinogenicity was expected to require careful review. Most of the information pertaining to this group is specific to anthracene.



Benzo[a]pyrene (BaP) and the eight other PAHs in the third group have no commercial production or use, except as research laboratory standards. They are released to the environment inadvertently by combustion sources. With one exception (acenaphthylene), the chemicals in this group have very low vapor pressures and water solubilities. Several of the PAHs in the BaP group had been identified as carcinogens. Much of the information regarding this group of compounds is for BaP.

The exposure and risk assessment for each of the three groups of PAHs was treated in a separate chapter of a multivolume report; Chapter 3.0 (Volume II) concerns naphthalene; Chapter 4.0 (Volume III) concerns the anthracene group PAHs; and Chapter 5.0 (Volume IV) concerns the benzo[a]pyrene group PAHs. These chapters are bound separately.

Potential waterborne routes of exposure are the primary focus of these exposure and risk assessments because of the emphasis of OWRS on aquatic and water-related pathways. Inhalation exposures are also considered, however, in order to place the water-related exposures into perspective. Each chapter contains major sections discussing the following topics:

- Information on environmental releases of the subject PAHs, including the form and amounts released and the receiving medium at the point of entry into the environment (materials balance);
- Description of the fate processes that transform and/or transport the compounds from the point of release through environmental media until exposure of humans and other receptors occurs, and a summary of reported concentrations detected in the environment, with a particular emphasis on aquatic media;
- Discussion of the available data concerning adverse health effects of the subject PAHs on humans, including (where known) the doses eliciting those effects and an assessment of the likely pathways and levels of human exposure;
- Review of available data concerning adverse effects on aquatic biota and the levels of environmental exposure; and
- Discussion of risk considerations for various subpopulations of humans and other biota.

Two comments regarding the materials balance section are appropriate. First, these sections were based in large part on draft material prepared by Acurex Corporation, under EPA Contract 68-01-6017, and provided to Arthur D. Little, Inc. by EPA. Second, the phrase "materials balance" is somewhat inappropriate when applied to chemicals such as the PAHs that are produced primarily as byproducts of combustion

processes. Since most PAH production is inadvertent rather than deliberate commercial production, the conventional approach of trying to balance production versus use and environmental release is not strictly applicable to these chemicals. Therefore, the materials balance sections of these exposure and risk assessments are focused on estimates of releases from major sources such as combustion; considerable uncertainty is associated with most of these estimates.

After an initial review of the three exposure and risk assessments covering all 16 priority pollutant PAHs, it was determined that one chemical, benzo[a]pyrene, was of appreciably greater interest to OWRS than were the other 15 compounds studied. This interest reflects the more extensive data base available for assessment of environmental fate and exposure and also the existence of some, although limited, dose-response data to which various extrapolation models can be applied for estimation of potential human carcinogenic risk from ingestion of BaP. For the other PAHs considered, data on carcinogenic or other long-term effects were generally limited, nonquantitative, and/or did not indicate statistically positive results. Table 1-1 presents a summary of the hazard of the 16 priority pollutant PAHs in terms of carcinogenicity, based on qualitative review of available information.

For these reasons, the technical summary presented in Volume I is organized somewhat differently than the rest of the report (Chapter 3.0-5.0) (Volumes II-IV). The summary is focused on benzo[a]pyrene as the PAH of greatest interest. The estimated releases to the environment, environmental fate, monitoring data, human effects and exposure, biotic effects and exposure, and risk considerations concerning BaP are presented in expanded summary form. Abbreviated summaries are then provided for naphthalene, anthracene group PAHs, and the other PAHs considered.

Included in the summary volume are critical data and references to the literature so that this volume may be read and understood by itself without reference to the separately bound Chapters 3.0-5-0. The latter volumes contain more extensive compilations of data, more detailed discussions of the available information and of the interpretations drawn, and more complete documentation of the multiple literature sources that were reviewed in the course of this work.

TABLE 1-1. SUMMARY OF EVIDENCE FOR CARCINOGENICITY OF PRIORITY POLLUTANT PAHS

PAH*

Basis

Benzo[a]pyrene Positive oral carcinogen with

other positive carcinogenic

data.

Dibenz[a,h]anthracene Positive oral carcinogen with

other positive carcinogenic

data.

Benz[a]anthracene Positive oral carcinogen with

other positive carcinogenic

data.

Benzo[g,h,i]perylene Not tested orally, other posi-

tive carcinogenic or co-car-

cinogenic data.

Benzo[b]fluoranthene Not tested orally, other posi-

tive carcinogenic or co-car-

cinogenic data.

Chrysene Not tested orally, other posi-

tive carcinogenic or co-car-

cinogenic data.

Indeno[1,2,3-c,d]pyrene Co-carcinogen or initiator

with negative carcinogen or in

vivo mutagen.

Pyrene Co-carcinogen or initiator

with negative carcinogen or in

vivo mutagen.

Fluoranthene Co-carcinogen or initiator

with negative carcinogen or in

vivo mutagen.

Benzo[k]fluoranthene Negative in a single carcino

genic study.

Phenanthrene Several negative carcinogenic

and mutagenic studies but not

tested orally.

TABLE 1-1. SUMMARY OF EVIDENCE FOR CARCINOGENICTY OF PRIORITY POLLUTANT PAHs (Continued)

Anthracene Negative studies, tested

orally.

Naphthalene Negative studies, tested

orally.

^{*} No data for evaluation of carcinogenicity were available for acenaphthene, acenaphthylene, or fluorene.

5.0 BENZO[a]PYRENE, ACENAPHTHYLENE, BENZ[a]ANTHRACENE,
BENZO[b]FLUORANTHENE, BENZO[k]FLUORANTHENE,
BENZO[g,h,i]PERYLENE, CHRYSENE, DIBENZ[a,h]ANTHRACENE,
IDENO[1,2,3-c,d]PYRENE

5.1 MATERIALS BALANCE

5.1.1 <u>Introduction</u>

This section reviews both published and unpublished data concerning the production, use, and disposal of the benzo[a]pyrene (BaP) group PAHs in the United States. Information from the available literature has been reviewed to present an overview of major sources of environmental releases of these compounds. Annotated tables have been included to aid data evaluation.

The section is organized according to the major categories of the sources of releases to the environment. Section 5.1.2 describes the numerous combustion processes that release the BaP group PAHs; Section 5.1.3 discusses contained sources; and Section 5.1.4 describes the amounts of these compounds in the influents and effluents of Publicly Owned Treatment Works (POTWs). Section 5.1.5 is a summary of the materials balance.

5.1.2 Combustion Sources

Combustion is the major source of PAH releases to the environment. This section estimates releases of BaP group PAHs from residential heating, fireplaces, cigarettes, coal refuse piles, wildfire, carbon black manufacture, gasoline use, and utility boilers; the emission estimates are summarized in Table 5-1.

5.1.2.1 Residential Coal Combustion

Residential coal combustion is responsible for a significant fraction (5%) of PAH emissions for the compounds in this group. Since residential heating units are typically not controlled and are relatively inefficient, they produce large amounts of PAHs. However, the trend in energy utilization for home heating sources suggests a rapidly decreasing reliance on coal. Whereas coal was used for heating in 2.7% of residences in 1970, it was used in only 0.56% in 1977 (Census 1979). (This trend may have reversed in more recent years.) Figure 5-1 shows the estimated distribution of residential coal consumption by state. Emissions from residential coal combustion (see Table 5-1) were estimated from the emission factors in Appendix A, Table A-1 (EPA 1977a) and the 1978 coal consumption of 8688 x 10^3 kkg for residential heating use (DOE 1980).

Table 5-1. Estimated Air Emission of Benzo[a]pyrene-Group PAHs by Combustion Source, 1978 (kkg)

	Residential Coal Combustion	fireplaces	Wood	Auxiliary Residential Wood Heating	Cigarettes	Coal Refuse Piles	Prescribed Burning	Wildfire	Carbon Black	Yireb Wear	Agricultural Burning	Gasol Ine	ULITI Boll Coal		Incinerator	Utility and Industrial s IC Engines	011- fired
Acenaphthylene		10	400	500	2				1								
Benzo[a]antinracene	20 20 20	2	50	70	negŒ		100	60	neg		40				neg	•	
Benzo[b] f voranthene	20	2	40	50	•	neg	50	30	neg	neg	20				neg		
Benzo[1] Huoranthene	20	2	40	50		neg	50	30	neg	neg	20				neg		
Benzu(ghi)perylene	1	3	40	50	neg	•	90	50	neg	á	30	40	1	•	neg		
Benzo(a)pyrene	10	2	30	40	neg	neg	30	20	neg	ĭ	10	iŏ	i	i	neg		
Chrysene	10 20	2	50	70	-	•	100	60	ney	-	40		•	•	neg		
Dibenzo[a,h]anthracene	30	neg	5	6		neg					***				incy		
Indeno(a, 2, 3-cd)pyrene	30 20		neg	neg	neg	neg	60	30	neq	1	20				neg		
Total	1 40	43	655	836	-	•	480	280	i	10	180	50	2	2	wed		neg

a) Negligible defined <1 kkg/year. Blanks indicate data not available. Emissions factors are presented in Table Λ -1, Appendix Λ . The text and Appendix A describe derivations of estimates.

b) PAH strongly bound to particulate carbon blank.

Sources: See text.

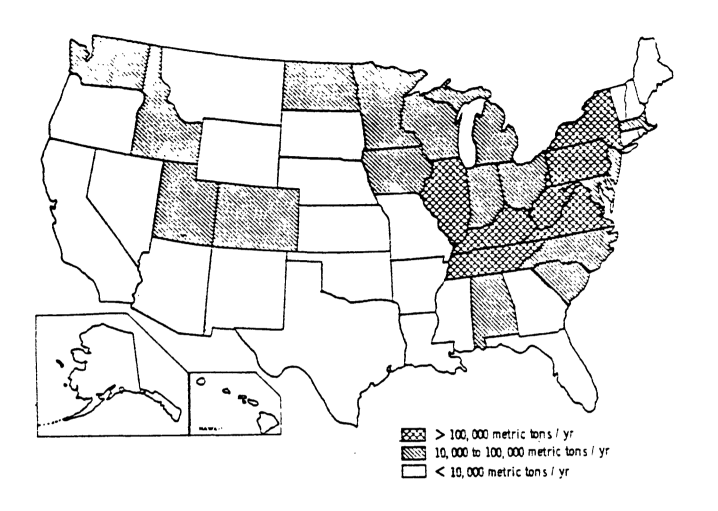


Figure 5-1. Estimated Residential Coal Consumption in 1974 by State (EPA 1977a)

5.1.2.2 Wood

Wood may be burned in homes in fireplaces or in stoves or furnaces for primary or for auxiliary heating. Estimated PAH emissions from these sources are listed in Table 5-1. Fireplaces produce smaller amounts of PAHs per kg of fuel than does wood combustion in primary heating units; in addition, fireplaces use less wood overall than is consumed for heating. The amounts of wood burned in fireplaces, for primary heating and for auxiliary heating in 1977, were 2.9 x 10^6 kkg, 6.9×10^6 kkg, and 9.2×10^6 kkg, respectively (see Appendix A, Note 1). Table 5-2 lists the estimated 1976 wood consumption for residential burning by state (EPA 1980a). The number of homes heated by wood has increased by more than 50%, from 794,000 in 1970 to 1,239,000 in 1977 (Census 1979).

Emissions of BaP group PAHs from these sources were calculated using the emission factors presented in Table A-1 (Appendix A) and 1977 wood consumption of 6.9 x 10^6 kkg and 9.2 x 10^6 kkg for primary and auxiliary heating, respectively. The emission factors presented in Table A-1 are averages of those for baffled and nonbaffled wood stoves (EPA 1980a). Total 1977 wood consumption for primary heating was obtained from the number of housing units that used wood for primary heat in 1976 and 1977 (912,000 and 1,239,000) and a proportional extrapolation of the estimated 5.1 x 10^6 kkg of wood burned in 1976 to 6.9 x 10^6 kkg for 1977 (EPA 1980a and Census 1979). Total 1977 wood consumption for auxiliary heating (9.2 x 10^6 kkg) was extrapolated from the 1976 estimate of 8.5 x 10^6 kkg (EPA 1980a) on the basis of the increase in the number of houses with fireplaces (Appendix A, Note 1). The estimated wood consumption for each state is shown in Table 5-2.

5.1.2.3 Cigarettes

Although cigarette smoking may be the source of a significant PAH exposure to individuals, it is a small contributor to national PAH emissions. Cigarette smoking contributes <1 kkg/yr of each of the BaP group PAHs to the atmosphere.

The emission factors for cigarette smoking, shown in Table A-1, were used, along with the number of cigarettes produced (616 x 10^9) (USDA 1979) in order to determine the amounts of BaP group PAHs emitted. The emission factors for the BaP group compounds are all from Neff (1979).

5.1.2.4 Coal Refuse Piles

Coal refuse piles, impoundments, abandoned mines, and outcrops may burn through the spontaneous combustion of coal. Because of the relatively poor air supply and uneven heat distribution, this combustion is inefficient and, therefore, produces relatively large amounts of PAHs. The estimated 52×10^6 kkg of coal in refuse piles (EPA 1978a) produce

Table 5-2. Estimated Residential Wood Consumption by State, 1976 (kkg)

			Auxiliary Wood
State	Primary heating	Fireplaces	Stoves
			252.000
Alabama	150,000	34,000	360,000
Alaska	50,000	5,000	32,000
Arizona	36,000	34,000	26,000
Arkansas	280,000	21,000	100,000
California	190,000	360,000	270,000
Colorado	18,000	40,000	61,000
Connecticut	22,000	29,000	140,000
Delaware	5,000	5,000	13,000
Florida	30,000	88,000	210,000
Georgia	340,000	46,000	220,000
Hawaii	1,000	12,000	9,000
Idaho	60,000	12,000	19,000
Illinois	29,000	120,000	260,000
Indiana	58,000	58,000	120,000
Iowa	16,000	32,000	68,000
Kansas	24,000	26,000	56,000
Kentucky	190,000	32,000	170,000
Louisiana	38,000	35,000	84,000
Maine	180,000	16,000	440,000
Maryland	47,000	39,000	94,000
Massachusetts	26,000	91,000	270,000
Michigan	76,000	97,000	410,000
Minnesota	110,000	43,000	360,000
Mississippi	200,000	21,000	110,000
Missouri	330,000	55,000	230,000
Montana	60,000	12,000	18,000
Nebraska	12,000	18,000	39,000
Nevada	15,000	10,000	8,000
New Hampshire	57,000	13,000	340,000
New Jersey	19,000	114,000	170,000
New Mexico	120,000	69,000	25,000
New York	150,000	290,000	870,000
North Carolina		51,000	120,000
North Dakota	3,000	7,000	14,000
Ohio	47,000	120,000	250,000
Oklahoma	78,000	28,000	34,000
Oregon	290,000	38,000	250,000
Pennsylvania	120,000	190,000	280,000
Rhode Island	5,000	15,000	22,000
SouthCarolina	190,000	26,000	62,000 16,000
South Dakota	20,000	7,000	
Tennessee	320,000	41,000	850,000 290,000
Texas	87,000	120,000 18,000	25,000
Utah	11,000	7,000	190,000
Vermont	33,000	48,000	110,000
Virginia Washington	280,000 210,000	58,000	390,000
Washington	36,000	18,000	43,000
West Virginia	91,000	50,000	210,000
Wisconsin Wyoming	10,000	6,000	9,000
# your ng	10,000		
Total	5,100,000	2,700,000	8,800,000

Source: EPA 1980a

the estimated PAH emissions shown in Table 5-1. The emissions do not account for burning refuse in impoundments or abandoned mines and outcrops since the quantity of burning coal is impossible to measure or estimate (EPA 1978a). It is, however, estimated that only 21% of the mass present in coal refuse piles is burning at any given time (EPA 1978a). Table 5-3 presents a distribution of burning coal refuse by state. It is worth noting that Pennsylvania and West Virginia account for 80% of these fires. Although the emissions were not estimated, the states of Montana, Wyoming, Colorado, and New Mexico account for 66% of the approximately 400 burning abandoned mines or outcrops.

From the composition of the particulate polycyclic organic matter (POM) shown in Table A-1, emissions were calculated based upon a refuse pile volume of 190 x 10^6 m³ (21% of which is estimated to be burning), a density of 1.5 kkg/m³, and a POM emission rate of 1.3 x 10^{-8} kg/kkg-hr (EPA 1978a). Only particulate emissions were analyzed here; and only preliminary sampling data are presented. Based upon these assumptions, the emissions are estimated to be less than 1 kkg. Alternatively, NAS (1972) gives an unsubstantiated estimate of 340 tons/year of BaP from coal refuse piles.

5.1.2.5 Forest Fires

Forest fires, either intentional or not (wildfire), are a major source of PAHs. Unfortunately, the estimates of PAHs produced are based on data that do not adequately cover the range of situations that may be termed wildfire. PAH production may be very crudely determined by using estimates of the amount of fuel burned and an emission factor. The amount of fuel burned (in terms of both fuel per unit area and total area burned) is for the most part based upon experienced judgments (EPA 1979a). Emissions factors are based upon laboratory simulations, which do not test the wide range of fuel type and configurations and may not appropriately simulate the fire intensity, weather conditions, fuel moisture, etc.

Prescribed burning is a controlled form of open burning used in land management to achieve specific objectives, which include fire hazard reduction, disease control, and silviculture, among others. An estimated 36 x 10⁶ kkg (dry weight) of fuel are burned annually, not including agricultural open burning (Pierovich 1978) (see Section 5.1.2.7). Table 5-4 presents a breakdown of this fuel usage by region; the estimated emissions for this source are presented in Table 5-1. These estimates were calculated from the amount of fuel burned and the emission factors for forest fires shown in Table A-1 (McMahon and Tsoukalas 1977).

Table 5-3. Coal Refuse Fires

State	Amount of 3 3 Coal Refuse (10 m)
Alabama	11,000
Colorado	12,000
Illinois	5,000
Kentucky	1,600
Maryland	23
Montana	230
Ohio	1,400
Pennsylvania	84,000
Utah	2,100
Virginia	2,800
Washington	2,300
West Virginia	67,000
Totala	190,000

a) Total does not add due to independent rounding

Source: EPA 1978a

Table 5-4. Estimated Net Total Weights of Fuel Consumed Annually in Prescription Burning on All Ownerships

	Net annual consumption, 10 ⁵ metric tons (dry weight)				
		Timber harvesting and land clearing residuesb			
_				Total	
Geographic ^a region	Piles or windrows	Broadcast (unpiled)	understory vegetation and litter	of all categories	
Alaska	0.5			0.5	
Eastern	0.1	0.3	0.1	0.5	
Intermountain	6.9	0.7	0.1	0.5 7.6	
Northern	82.7	6.3		89.0	
Pacific	02.7	0.5		09.0	
Northwest	50.4	37.0		87.4	
Pacific		U , • •		0/• T	
Southwest	20.2	8.2	0.5	28.9	
Rocky Mountain	4.9	1.0	0.2	6.1	
Southern	0.9	9.9	114.2	125.0	
Southwestern	12.3	8.3	0.3	20.9	
Total of all				2003	
regions	178.9	71.7	115.3	365.9	

a) See Figure 5-2 for states included in each region

Source: Pierovich 1978.

b) Includes precommercial thinning residues as follows: Pacific Southwest and Rocky Mountain, each <0.1 x 10^5 metric tons; Southwestern, 1.1 x 10^5 metric tons.

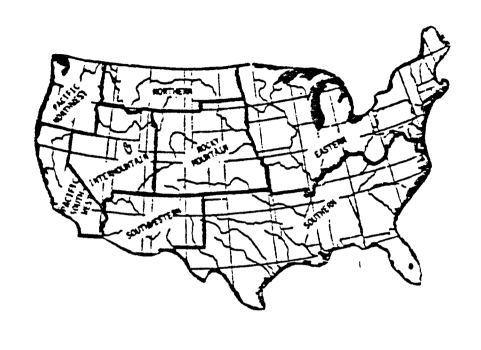


Figure 5-2. Geographic Regions used to Summarize Survey Data Within the Coterminous States. (Alaska Region also used in summarization not shown.)

Source: Pierovich 1978

Land burned by wildfire each year varies considerably in both the total area burned and the location of specific sites. On the average, $1 \times 10^{10} \text{ m}^2$ (3 x 10^6 acres) are burned each year, although in 1976, $2 \times 10^{10} \text{ m}^2$ (5 x 10^6 acres) were burned (Dahl 1980). The largest area is burned in the Southeastern section of the country, followed by the South, North Central, Pacific, and Rocky Mountain sections (Dahl 1980). Emissions estimates are shown in Table 5-1.

The emissions factors for prescribed burning and wildfire in Table A-1 (McMahon and Tsoukalas 1977) are averages obtained from six tests on pine needles, with differing fuel densities used for heading and backing fires; hence, these factors cannot provide a very accurate basis for nationwide emissions estimates. The total amount of fuel burned was estimated by assuming that $10^{10}~\rm m^2$ (3 x $10^6~\rm acres$) of land were burned on the average (Dahl 1980), and that the fuel loading was 2 kg consumed dry weight/m² (based upon estimates by EPA 1978b). Total quantities of BaP group PAHs emitted from this source are estimated in Table 5-1.

5.1.2.6 Carbon Black

Carbon black is produced primarily by the furnace process, in which a liquid hydrocarbon feedstock is incompletely pyrolyzed in a refractory-lined furnace by burning natural gas. The carbon black particles are recovered in a baghouse after being cooled by water sprays. The carbon black is then converted to a marketable product. The principal emission source from carbon black manufacture is baghouse exhaust gas from the main process vent. PAHs are present both in the vent gas and on the carbon black itself (see Appendix A, Note 2 and Tables A-1 and A-3). For the latter reason, carbon black was banned in food, drugs, and cosmetics in 1976.

Emissions of PAHs from carbon black manufacture are presented in Table 5-1; Table 5-5 lists the location and capacity of carbon black producers. For the most part, the emissions during production are negligible (i.e., <1 kkg/yr). Larger amounts, however, are contained in/on the carbon black. These are shown in Table A-3.

The primary use for carbon black is in the elastomers used in automobile tires (65%), while other elastomeric uses account for 29% (SRI 1979). The PAHs present on the carbon black are strongly bound to it and, therefore, are not likely to be easily separated from it, except by prolonged extraction (Locati et al. 1979). An estimate of PAH air emissions from tire wear, presumably associated with carbon black, is discussed in Note 2 and presented in Table 5-1 (see Appendix A). PAHs associated with carbon black that is transferred to road surfaces are presented in Table A-4. Although this release presents a possible source of PAHs in runoff, these PAHs are probably tightly bound to the carbon black particles.

Table 5-5. Locations and Capacities of Carbon Black Producers

Company and Location	Annual Capacity 10 ³ kkg
ASHLAND OIL, INC.	
Ashland Chemical Company, divisio	n
Carpon Black and Synthetic	••
Rubber Division	
Aransas Pass, Texas	74
Belpre, Ohio	45
Iberia, Louisiana	120
Mojave, California	30
Shamrock, Texas Total	<u>50</u>
local	320
CABOT CORPORATION	
Big Spring, Texas	110
Franklin, Louisiana	100
Rampa, Texas	24
Villa Platte, Louisiana	110
Waverly, West Virginia	74
Total	420
CITIES SERVICE COMPANY	
CITIES SERVICE COMPANY	
Chemicals Group Columbian Chemicals, division	
Conroe, Texas	AA
El Dorado, Arkansas	44
Eola, Louisiana	7 32
Mojave, California	24
Moundsville, West Virginia	71
North Bend, Louisiana	71
North Bend, Louisiana	25
Seagraves, Texas	40
Jlysses, Kansas	_23
Total	370
CONTINENTAL CARBON COMPANY	
owned by Continental Oil Company	
30% and witco Chemical Corporation,	
0%)	
Bakersfield, California	35
Phenix City, Alabama	23
Ponca City, Oklahoma	61
Sunray, Texas	43
Westlake, Louisiana Total	<u>55</u> 217
/ULB1	21/
.M. HUBER CORPORATION	
Baytown, Texas	117
Borger, Texas	62
Borger, Texas	19
Total	198
HILLIPS PETROLEUM COMPANY	
Rubber Chemicals Division	
Borger, Texas	130
Orange, Texas	61
Toledo, Ohio	_ 39
Total	230
ID RICHARDSON CARBON COMPANY	
Addis, Louisiana	46
Big Spring, Texas Total	<u>-5÷</u>
IUCAI	96 TOTAL 1 050
	TOTAL 1,850

Totals may not add due to rounding Source: SRI 1980.

5.1.2.7 Agricultural Open Burning

Agricultural open burning involves burning of crop residues for residue removal, field sanitation, or preparation of farmlands for cultivation. Wastes that may be burned include a wide variety of residues, including rice straw, orchard prunings, potato vines, and bagasse, among others. Although PAH formation is dependent upon fuel type, among the many other variables, the emissions in Table 5-1 were calculated based upon wood burning (forest fire; see Table A-1, Appendix A), and an estimated total of 13 x 10⁶ kkg dry weight of burned material (EPA 1977b). Table 5-6 lists the estimated distribution of open burning by state.

5.1.2.8 Motor Vehicles

Motor vehicles that burn gasoline are ubiquitous sources of PAHs, although emissions control devices for hydrocarbons in exhaust gas and from the crankcase reduce the amount emitted at efficiencies up to 99% (Gross 1972) PAHs released by motor vehicles are either emitted in the exhaust or dumped along with the used crankcase oil. Estimated PAH emissions from exhaust gas are presented in Table 5-1 (see Appendix A, Note 3). Although crankcase controls reduce hydrocarbon emissions from the vehicle, PAHs still reside in the crankcase oil and may be released to either land or water if the oil is disposed of haphazardly. Table 5-7 presents estimates of the PAHs contained in these releases. (See Appendix A, Note 3 and Table A-5.

5.1.2.9 <u>Incinerators</u>

PAH releases from municipal and commercial incinerators are estimated to be negligible (<1 kkg/yr). Releases were calculated for municipal incinerators using the release factors in Table A-6 (Davies 1976) for 104 plants with an average capacity of 385 kkg/day when operating at full capacity (EPA 1978b).

Releases for commercial incinerators were assumed to be similar to those from municipal incinerators. Even when the higher emission factors in Table A-6 are used, calculated emissions of any given PAH were negligible without controls. The incinerators were assumed to number 100,000 units, firing 3 hours/day, 260 days/year at an average capacity of 0.1 kkg/hr (EPA 1978b).

Table 5-6. Distribution of Agricultural Open Burning, 1973

	Area burned	Amount of crop residue burned
State	10 ⁷ m ² /yr	10 ³ kkg/yr
Alabama	36.	161
Arizona	4.5	20
Arkansas	26.	115
California	309.	2,075
Colorado	32.	143
Delaware	0.1	2
Florida	109.	1,716
Georgia	39.5	883
Hawaii	44.7	1,175
Idaho	4.8	22
Kansas	243.	544
Kentucky	13.	58
Louisiana	142.	1,904
Maine	15.	33
Maryland	0.6	7
Massachusetts	0.6	7 3 93
Michigan	21.	93
Minnesota	61.2	274
Mississippi	138.	617
Missouri	40.5	181
Montana	34.	152
Nebraska	67.6	303
Nevada	0.8	5
New Mexico	0.5	2
North Carolina	138.	619
North Dakota	97.	438
Ohio	32.	142
Oklahoma	36.	161
Oregon	107.	479
Pennsylvania	15.	69
South Carolina	8.	38
South Dakota	59.	265
Tennessee	14.	61
Virginia	9.5	43
Washington	57.	256
Wisconsin	32.	145
Wyoming	7.3	34
National Total	1995.	13,238

Source: EPA 1977b.

Table 5-7. BaP Group PAHs Discharged Annually in Used Crankcase Oila(kkg)

PAH	Discharge	
Benzo[a]anthracene Benzo[k]fluoranthene Benzo[ghi]perylene Benzo[a]pyrene Chrysene	2 3 3 neg ^b 2	

a) Releases go presumably to POTWs and landfills. No recycling is assumed.

b) Negligible is <1 kkg.

Source: Peake and Parker 1980 and Tanacredi 1977.

See Note 3, Appendix A. 1

5.1.2.10 Coal- and Oil-fired Utility Boilers

Table 5-1 summarizes PAH emissions from coal- and oil-fired utility boiler units. The estimates of total PAHs released to the environment are based upon the emission factors averaged from Table A -7 and a total coal and oil consumption for electricity generation of 4.8 x 10^8 and 7.8 x 10^7 kkg per year, respectively (Monthly Energy Review 1980).

These emissions are uncontrolled releases calculated from 1967 emissions factors, and are probably lower today with the use of baghouses or electrostatic precipitator units. Further, the emission factors for oil are for small- or intermediate-sized units, and thus serve as an upper limit of the PAH releases from higher capacity plants.

5.1.2.11 Utility and Industrial Internal Combustion Engines

Stationary internal combusion engines are used for electricity generation, oil and gas transmission, natural gas processing, and oil and gas production and exploration. These engines are either gas turbines or reciprocating engines and are fueled with either oil or gas. Table 5-1 presents the estimated amount of PAHs emitted from these sources in 1978 as estimated by EPA (1979d).

5.1.2.12 Gas- and Oil-fired Residential Heating

Table 5-1 lists PAH emissions from oil-fired residential heating units as estimated by EPA (1979e). Although a large fraction of residences provide space heat with gas, gas-fired units emit negligible amounts of PAHs. Figure 5-3 presents the geographic distribution of oil and gas used for residential heating in 1975 (EPA 1979e).

5.1.3 Contained Sources

This section presents information, primarily in tabular format, on sources of BaP group PAHs from which these compounds are not specifically isolated. Delineation of all such "inadvertent sources" of PAH releases is indeed a monumental task because of the omipresent nature of petroleum— and coal-derived oils, fuels or solvents that contain at least small amounts of PAHs. The major sources discussed here are summarized in two tables: Coal Tar Production and Distillation (Table 5-8) and Petroleum Sources (Table 5-9); supporting data and related information are presented in Tables 5-10 and 5-11 and Appendix A.

It is important to comment on the quality of available data in this section. Specifically, concentrations of the various PAHs in crude oils, coal, or coal and petroleum products are highly variable, depending upon the geographic source and method of processing. Furthermore, as subsequent calculations are based on these concentrations, they can be considered as estimates only, at the order of magnitude level of reliability.

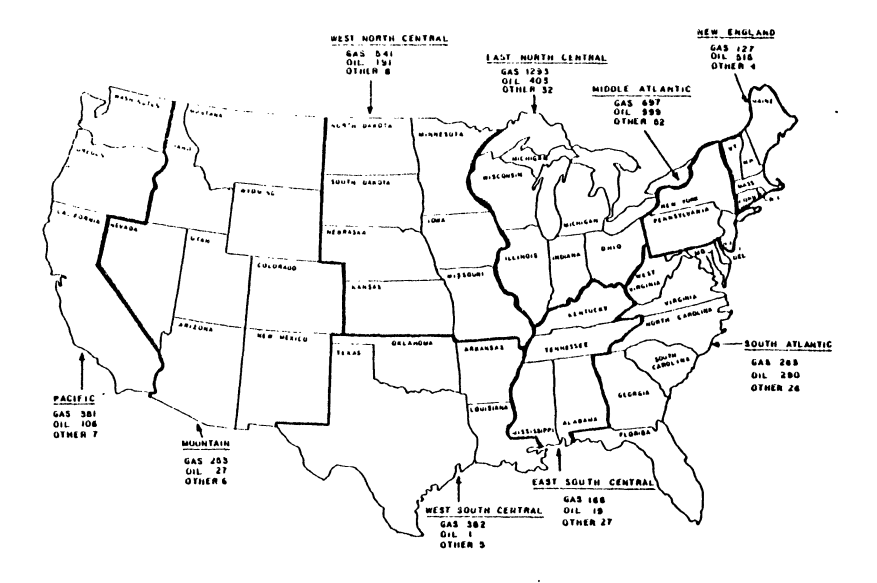


Figure 5-3. Geographical Distribution of Residential Fuel Consumption for Space Heating, 1975, 10¹⁵ J/yr (EPA 1979e)

Table 5-8. PAH Materials Balance: Coal Tar Production and Distillation, 1978 (kkg/yr)a

	Tar <u>Production</u> b	Used By Proc Refining/fuel Topping		Sold for Quantity	Refining b On Hand Dec. 31	Coal-Tar Pitch ^C	Coal-Tar Creosote		Envir f Land	1	tal Rele Water ^e Surface	eases E Total
Acenaphthylene. Benzo(a)anthracene Benzoperylenes		auaitable)				8	,	9 1 0.4	10	10	10	40 7 neq
Benzo(a)pyrene Chrysene. Indenopyrenes	77	18 13 2400 1700 available)	1.2 150	44 3700	6.6 860	8	urgg	2 3 0.4	0.6	0.7 2	0.9	4 10 neg
Total		_				Total		16	15	15	16	60

- a) All production values rounded to two significant figures, environmental releases to one figure; blank spaces = data not available. Totals may not add due to rounding.
- b) See Tables A-8, 9 for tar production/distribution totals and concentrations of PAHs, respectively. Density of Coal tar = 1.223 kg/l.
- c) See Table A-9 for PAH concentrations; total based on 790 x 10^3 kkg pitch produced in 1978 (USITC 1979). d) See Table A-9 for PAH concentrations; total based on 3.3 x 10^8 1 creosote oil produced in 1978 (USITC 1979). Density of creosote oil = 1.06 kg/l.
- e) See Table A-10 for coke plant effluent discharge factors. Distribution of discharge: 33% direct, 25% POTWs, 2% deep well, 40% quenching (20% - land, 20% - air) (EPA 1979d). All values rounded to one significant figure.
- f) Includes emissions from coke-oven doors (see Table 5-10).
- g) Less than Ikkq.

Table 5-9. PAlls in Contained Petroleum Sources, (kkg/yr)a

		h			Environmental Releases ^C					
PAH	Crude Oild	Input ^b Gasolinee	Diesel Fuelf	0il S Water	pills Land	<u>Gasoline</u> Water	Spills Land	Refi	Petroleu nery Was Water	
Acenaphthylene	340,000			10	neg 9			 		
Jenzo[a]anthracene	trace	250	3	neg	neg	neg	neg			
Benzo[b]fluoranthene	<4,200			neg	neg	-	_			
Benzo[k]fluoranthene	<4,200			neg	neg					
Benzo[ghi]perylene	17	170	0.9	neg	neg	neg	neg	5		
Benzo[a]pyrene	840	170	2	neg	neg	neg	neg	3		neg
Chrysene	<84,000	170	20	3	neg	neg	neg			
)ibenzo[a,h]anthracene					neg					
Ideno[1,2,3-cd]pyrene					neg					

a) Blanks indicate data not available.

a) Blanks indicate data not available.
b) See Table A--11 for PAH concentrations.
c) See Appendix A, Note 4, for derivations
d) Based on 8.4 x 10⁸ kkg of crude oil consumed (Guerin 1978).
e) Based on 7.4 x 10⁶ bbl/day consumed (Oil and Gas Journal 1979); 42 gal/bbl; 0.73 kg/l.
f) Based on 3.4 x 10¹⁰ l/yr; 0.865 kg/l.
g) Less than one kkg.
h) Based on data in Table A--13, 4.985 x 10⁶ bbl/day feed for catalytic cracking, 0.887 x 10⁶ bbl/day for catalytic hydrocracking (Oil and Gas Journal 1979), 42 gal/bbl.

Table 5-10. PAH Emissions: Coke-Oven Doors

	Emission Rate (mg/hr/oven) ^a	Y	/early Emission (kkg/yr) ^b
Chrysene Indenopyrene	16		1.0
Benzoperylene	6 6		0.4 0.4
Benzopyrene	16		1.0
		Total	3.0

a) EPA 1977e b) Based on 1300 kkg coke produced per typical coke oven battery of 58 ovens; 160,000 kkg/day typical capacity for total by-product cokemaking industry (resulting in a total of 123 batteries); 365 day, 24 hour operation; emission data in first column, EPA 1979f.

Table 5-11. Water Discharges of Benzo[a]pyrene Group PAHs: Timber Products, 1978 (kkg/yr)^a

		Raw Disc	harge	Treated Discharge		
РАН	Input	Concentration (mg/x)	Quantity ^b	Concentration (mg/1)	Quantity ^b	
enaphthylene	6,900	0,68	5	0.05		
enzo[a]anthracene	neg	0.85	5 6	0.05 0.43	neq 3	
enzo[b]fluoranthene	neg	0.39	3	0.43	2	
enzo[k]fluoranthene	neg	0.45	3	0.03	ney	
enzo[ghi]perylene	neg	0.05	neg	0.01	neg	
enzo[a]pyrene	neg	0.36	2	0.04	neg	
hrysene	0.4	0.69	5	0.90	neg	
ibenzo[a,h]anthracene	neg	0.007	neg	ND	neg	
ndeno[1,2,3-cd]pyrene	neg	0.18	1	0.02	neg	

a) See Table 5-8, coal-tar creosote.

Source: EPA, 1979d.

b) Based on 56% of 476 total plants using creosote or mixture thereof, 350 day/yr, 75,500 liters/day/plant, assumed to go to POTW.

5.1.3.1 Coal Tar

Coal tar is the heavy distillate fraction from the destructive distillation (coking) of coal. The distribution of coke-oven tar production in the U.S., PAH concentrations, and environmental releases (Tables 5-10. A-8, and A-9) have been combined to provide the information presented in Table 5-8. Naphthalene (see Section 3.1), present in the largest concentration in coal tar, is the only PAH compound warranting recovery and isolation in large amounts. Anthracene is also isolated, but to a lesser extent and usually in crystals which form during creosote oil recovery (see Section 4.1). Creosote oil is used in the wood preserving industry. Information on PAH concentrations in creosote oil and wastewater discharges during use are presented in the previously mentioned tables and Table 5-11, respectively. Of a total of 224 wood preserving plants responding to EPA's data collection protocol, two reported direct discharge, 47 reported discharge to POTWs and the remainder reported self-contained no-discharge operations (mostly evaporation with some oil irrigation or treated effluent recycle; EPA 1979g).

Besides creosote, other principal tar products containing PAHs are pitch and refined tar, used in a variety of applications ranging from road materials and electrodes to shampoos. Coal tar and tar products are also used as fuel (see Table 5-8), either by producers (e.g., iron and steel plants) or distillers.

5.1.3.2 Petroleum Sources

The other fossil fuel source containing appreciable amounts of PAHs is petroleum. The concentration and emissions data in Appendix Tables A-11, A-12 and A-13 have been combined for use in the summary Table 5-9.

Only spills and petroleum refinery environmental releases are presented here; gasoline combustion has been covered in Section 5.2.1 with other combustion sources. As mentioned previously, the type of crude feedstock determines its chemical composition and, therefore, the composition of specific waste streams. Other variables include pollution controls, age of the processes used (level of technology), and operational practices and control. It has been estimated that a total of 0.1 kkg benzo[a]pyrene is released in solid waste streams industry-wide from petroleum refining (EPA 1976). The air emissions also listed in Table 5-9 are specifically from petroleum catalytic cracking, which accounts for over 50% of the annual oil feed to refineries (0il and Gas Journal 1979); the remainder is used for catalytic reforming, for which emissions factors were not available.

5.1.4 Publicly-Owned Treatment Works

Input of PAHs to POTWs is largely dependent upon variations in industrial discharges feeding the POTWs and the types of industry in a particular municipality. A recent EPA study of 20 urban POTW facilities with secondary treatment and varying feed conditions produced a materials balance of PAHs shown in Table 5-12.

Table 5-12. Materials Balance of Benzo[a]pyrene Group PAHs: Municipal POTWs (kkg/yr)a

		Enviro	Environmental Releas		
РАН	Input ^b	Air ^C	Water ^d	Land ^e	
Chrysene Indeno[1,2,3-c,d]pyrene Benzo[a]pyrene Benzo[b]fluoranthene Benzo[k]fluoranthene Benzo[ghi]perylene Dibenzo[a,h]anthracene Benzo[a]anthracene Acenaphthylene	7.3 3.7 3.7 3.7 3.7 11 3.7 7.3 ND	NA ^f 3.7 3.7 0.7 3.7 11 3.7 NA	3.7 ND ND ND ND ND ND ND 3.7 3.7	9.1 h neg neg neg 9.1 neg	

a) All values rounded to two significant figures.

c) Difference between input and water and land.

Based on influent concentrations shown in Table 5-13, 10¹¹ liters per day total POTW flow.

Based on secondary effluent concentrations shown in Table 5-13, 10^{11} 1/day total POTW flow.

Based on wet sludge concentrations shown in Table 5-13, 6x10⁶ metric tons dry sludge generated/yr, wet sludge 95% water (by weight).

Not available. Not detected. f)

<1 kkg. h)

Table. 5-13. PAH Concentrations in Municipal POTWs ($\mu g/l$)

PAH	Influent	2° Effluent	Raw Sludge
Acenaphthylene	ND	0.1	3
Benzo[a]anthracene	0.2	0.1	76
Benzo[b]fluoranthene	0.1	ND	25
Benzo[k]fluoranthene	0.1	ND	2.5
Benzo[ghi]perylene	0.3	ND	0.6
Benzo[a]pyrene	0.1	ND	3.9
Chrysene	0.2	0.1	76
Dibenzo[a,h]anthracene	0.1	ND	0.2
<pre>Indeno[1,2,3-c,d]pyrene</pre>	0.1	ND	0.2

a) Average values.

Source: EPA 1980d

The materials balance in Table 5-12 was constructed using a total POTW flow of approximately 10^{11} 1/day (EPA 1978c) and the average concentrations of the various PAHs in influent, effluent and sludge, presented in Table 5-13. For purposes of these calculations, influent and effluent flow rates are assumed to be equal, i.e., water losses from sludge removal and evaporation are small compared with influent flows. With these assumptions, an estimated 70 kkg of PAHs were discharged to water from POTWs in 1978, while there was an input of 590 kkg in influent. Of aquatic releases from POTWs, 11 kkg are attributed to BaP group PAHs.

PAHs discharged in sludge can be estimated from the PAH concentrations in sludge and the quantity of dry sludge produced annually: $6.0 \times 10^6 \ \mathrm{kkg}$ (EPA 1979i). Wet sludge is assumed to be 95% water by weight. As ocean dumping of sludge is mandated to cease by 1981, and if one assumes that more stringent air quality standards curb incinerator use (EPA 1979j), the 25 kkg of the BaP group of PAHs contained in sludge are assumed to be discharged to land.

PAHs released to the atmosphere were estimated by the difference from the above calculations (influent loading - effluent and sludge loading). On this basis, an estimated 30 kkg of these PAHs were released to the atmosphere from POTWs in 1978.

5.1.5 Summary

As shown in Table 5-14, the largest amount, 96% (2700 kkg) of the total releases of BaP group PAHs (2800 kkg), is emitted to the atmosphere. Of the atmospheric emissions, combustion (Table 5-1) is the source of 99% of the BaP group PAHs released. Of the various compounds, releases of acenaphthylene are much greater than the other PAHs in this group. Aquatic discharges that have been identified (37 kkg total) are chiefly the result of oil/gas spills and POTWs (Tables 5-9 and 5-12, respectively). Releases to land (50 kkg) are again mainly due to coal tar production and POTWs.

Table 5-14. Estimated Environmental Releases of BaP Group PAHs (kkg/yr) (Continued)

Chrysene	Λir	Land	<u>Wate</u> Surface	POTW	Total
combustion ^a crankcase oil disposal ^b coal tar production contained petroleum sources ^e timber products	340 3	l 2 neg	3 3	l 2 neg	340 2 10 3 neg
tire wear POTW Total:	340	9 10	4 10	3	1 <u>3</u> 370
Dibenzo[a,h]anthracene combustion crankcase oil disposal coal tar production	40				40
contained petroleum sources timber products tire wear POIW	4	neg	nea	neg	neg neg
Total:	40	neg neg	<u>neg</u> neg	neg	$\frac{4}{40}$
Indeno[1,2,3-c,d]pyrene					
combustion crankcase oil disposal	130		•		130
coal tar production contained petroleum sources timber products	neg	neg		neg	neg neg neg
tire wear POTW Total:	$\frac{\frac{1}{4}}{130}$	2 <u>neg</u> 1	neg neg	neg	2 130

5.2 FATE AND DISTRIBUTION IN THE ENVIRONMENT

5.2.1 <u>Introduction</u>

This section describes the fate processes that determine the ultimate distribution of BaP and the other related PAHs in the aquatic environment and, therefore, the opportunities for water-borne exposure to humans and other biota. Much of the information presented pertains specifically to BaP; however, since the properties and fate characteristics of this compound are similar to those of the other compounds in the group, the behavior of BaP in the environment is believed to be a good model. In addition, some information concerning benz[a]anthracene is presented from a major study of the fate of organics in the environment (Smith et al. 1978).

Section 5.2.2 presents an overview of the environmental loading of BaP to the aquatic media, both direct releases to surface water and deposition from the atmosphere. In Section 5.2.3, physical/chemical properties of the BaP group PAHs are summarized in order to identify the processes that transform and transport the chemical upon its release to the environment (Section 5.2.3.1). Section 5.2.3.2 discusses the interplay of fate processes that determines the major pathways of BaP in aquatic environmental media.

Modelling efforts were undertaken based upon environmental loadings estimated in Section 5.1, in order to characterize the fate and distribution of BaP in specific environmental scenarios; these are discussed in Section 5.2.3.3. Monitoring data from STORET and a limited number of other surveys are summarized in Section 5.2.4 to provide indications of concentrations of BaP group PAHs actually detected in aquatic media. Finally, Section 5.2.5 summarizes those aspects of the fate and ultimate environmental distribution of BaP having the greatest significance for the water-born exposure of humans and other biota.

5.2.2 Inputs to Aquatic Media

5.2.2.1 Atmospheric Deposition

Data presented in Section 5.1 indicate that direct releases of BaP to surface waters are negligible; atmospheric emissions of BaP from combustion sources were estimated to be approximately 170 kkg in 1978 (Section 5.1). Atmospheric BaP may be transported to the aquatic environment indirectly via wet and/or dry deposition. These physical removal mechanisms are expected to be significant for BaP; Cupitt (1980) suggests an atmospheric residence time (time required for BaP to be reduced to 1/e of its original value) of approximately 8 days.

The air-to-surface transport pathway has been evaluated for BaP and is detailed in Appendix B; the results of that analysis are summarized in Table 5-15. Under ambient conditions in either rural or urban areas, virtually all of the BaP is adsorbed onto aerosols. The deposition

Table 5-14. Estimated Environmental Releases of BaP Group PAHs (kkg/yr)

			Wat	er	
Acenaphthylene	Air	Land	Surface	POTW	Total
combustion ^a	930	_c			930
crankcase oil disposal ^b coal tar production ^d contained petroleum sources ^e timber products tire wear	9	10 neg	10 10	10 neg	40 10
POTW Total:	940	<u>neg</u> 10	$-\frac{4}{20}$	10	<u>4</u> 980
Benzo[a]anthracene					
combustion crankcase oil disposal	340	1		1	340
coal tar production contained petroleum sources timber products tire wear	1	2 neg	2 neg	3	2 7 neg 3
POTW Total:	340	$-\frac{9}{10}$	$-\frac{4}{6}$	6	$\frac{13}{360}$
Benzo[b]fluoranthene					
combustion crankcase oil disposal coal tar production	210				210
contained petroleum sources timber products tire wear		neg	neg	2	2
POTW Total:	$\frac{1}{210}$	$\cdot \frac{3}{3}$	<u>neg</u> neg	2	$\frac{4}{220}$
 a) See Table 5-1. b) See Table 5-7. c) Blanks mean data not available; neg means <1. d) See Table 5-8. 	f) See g) See	Table 5-9. Table 5-11. Table A-4, air Table 5-12.	releases	included	in combustion.

Table 5-14. Estimated Environmental Releases of BaP Group PAHs (kkg/yr) (Continued)

	· 				
Benzo[k]fluoranthene	Air	Land	Wate Surface	POTW	Total
combustion ^a crankcase oil disposal ^b coal tar production	210	1.5	• .	1.5	210 3
contained petroleum sources ^e timber products tire wear		neg	neg	neg	neg neg
POTW Total:	4 210	neg 2	neg neg		$\frac{4}{220}$
Benzo[ghi]perylene					
combustion crankcase oil disposal coal tar production contained petroleum sources	310 neg 5	1.5 neg	neg	1.5	310 3 neg 5
timber products tire wear POTW Total:	$\frac{8}{11}$	7 <u>neg</u> 9	neg neg	neg	neg 15 <u>11</u> 340
Benzo[a]pyrene					
combustion crankcase oil disposal coal tar production	160 2	neg 0.6	0.9	neg 0.7	160 neg 4
contained petroleum sources timber products tire wear	3 1	neg 1	neg	neg	4 3 neg 2
POTW Total:	<u>4</u> 170	neg 2	neg 1		2 4 170

TABLE 5-15. EVALUATION OF AIR-TO-SURFACE PATHWAY FOR BENZO[a]PYRENE

	<u>Rural</u>	Urban
Adsorbed fraction of airborne mass	0.99	1
Dry deposition Velocity (cm/sec)	1	1
Precipitation scavenging ratio:	6x10 ⁴	6x10 ⁴ -1.2x10 ⁵ a
Percent of atmospheric emissions deposited		
dry depositionwet depositiontotal	22 4 26	19 4-7 23-26

 $^{{\}tt a}\,{\tt Depends}$ on distance from combustion source.

parameters for BaP vary only slightly from rural to urban environments, since they are highly dependent upon vapor/aerosol partitioning. As shown in Table 5-15, approximately 26% of atmospheric BaP will be deposited on the surface. The analysis indicates that dry deposition of BaP adsorbed onto atmospheric aerosols accounts for most of the removal; wet deposition is less significant by a factor of from three to five, depending upon the extent of equilibrium achieved in the combustion plume.

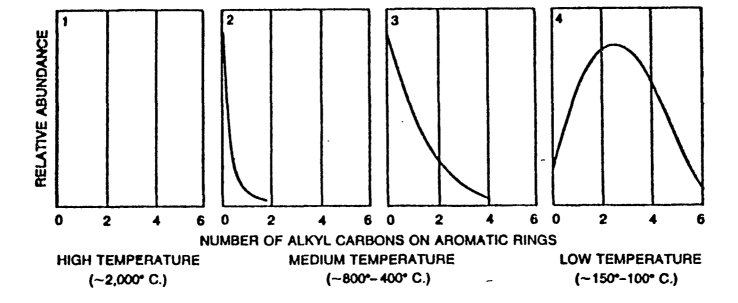
On the basis of total annual atmospheric emissions of 170 kkg for 1978 (Section 5.1), approximately 44 kkg may have been deposited on the surface of the U.S. Some of the fallout will land on surface waters. If approximately 2% of the total area of the continental U.S. is surface water (U.S. Bureau of the Census 1980), direct deposition onto aquatic systems would have amounted to less than 1 kkg.

A fraction of the remaining fallout deposited on land would be transported ultimately to aquatic systems via surface runoff. Data cited by Neff (1979) suggest that 500 kkg/yr of BaP are deposited on surface waters worldwide (including oceans) and that 118 kkg/yr end up in aquatic systems via surface runoff. These ratios indicate that surface runoff would deposit much less than 1 kkg of BaP in U.S. waters each year. The local effects of surface runoff, especially near combustion sources, could still be significant. The BaP adsorbed to surface particles is tightly bound and would probably be carried along with these particles in runoff. However, there are insufficient data to estimate the importance of this pathway.

5.2.2.2 Source Identification by PAH Composition

Sediments have an integrating effect on patterns of PAH input over time and supply geographical information when current patterns, sediment origins, and settling rates are known (Dunn and Stich 1976). It should, however, be mentioned that the composition in sediment does not always directly reflect the original PAH composition at the source. Partitioning between the sediment and aqueous phases may be different for parent PAHs and alkylated PAHs within an aromatic series, since solubility in water decreases as the number of alkyl carbons increases (Armstrong et al. 1977).

Information on whether the origin of PAHs in water is atmospheric deposition from combustion sources or direct discharge can be gained from studying the PAH composition in the sediments. The extent of alkylation of PAH mixtures has been shown to be highly dependent upon the combustion temperatures (Blumer 1976). The number of alkyl carbons on PAHs formed during combustion of fuel (400°C-2000°C) is generally 0-2; at the low temperatures at which petroleum was formed over geologic time (100°C-150°C), PAHs with 2-4 alkyl carbons are most common (Figure 5-4).



Source: Blumer (1976)

FIGURE 5-4 EFFECT OF COMBUSTION TEMPERATURE ON RELATIVE ABUNDANCE OF ALKYL CARBON ATOMS ON PRODUCED POLYNUCLEAR AROMATIC HYDROCARBONS

Furthermore, it has been reported that the distribution of homologs in PAHs measured in soils and marine sediments varies little over a wide range of depositional environments. Homologs extending to at least seven alkyl carbons occurred in all samples and all PAH series. However, within a PAH series, the unsubstituted hydrocarbons were the most abundant with the intensities of isomeric homologs, decreasing nearly twofold with each additional carbon atom (Blumer and Youngblood 1975). one study of PAH composition of sediment, a semilog plot of abundance vs. number of alkyl carbons was found to produce a straight line with a highly negative slope (Hase and Hites 1977). These data are presented in Figure 5-5, along with similar measurements for river water and air particulates. The authors postulated that the steep slope characteristic of urban air particulates was changed to the more gradual slope of the sediments due to the differential solubilities and partitionings of parent PAHs and their alkyl homologs. After airborne particulates are deposited on soil or water, the lower homologs continuously fractionate into the water phase, thereby increasing the relative abundance of the alkylated species in the sediment. The slope of the airborne particulate plot is intermediate between those of water and sediment, and this led the authors to the conclusion that the main source of the aquatic load of PAHs is anthropogenic, airborne PAHs produced during combustion.

It should be noted that industrial effluents containing PAHs would undergo similar fractionation (Lewis 1975, Armstrong et al. 1977). PAH profiles from petroleum sources show that parent compounds of each series are less abundant than the alkylated forms. The fact that the unsubstituted PAHs are the most abundant homolog (despite higher aqueous solubility) in sediment from non-industrial areas further indicates that the major source of PAHs in sediment is generally a combustion source rather than direct contamination by fossil fuels. Sediment known to be contaminated with fossil fuels exhibited an elevated concentration of four- and five-ring PAHs (due to lower water solubility) similar to that in the uncontaminated sediments, but also contained a much higher proportion of alkylated compounds (Youngblood and Blumer 1975).

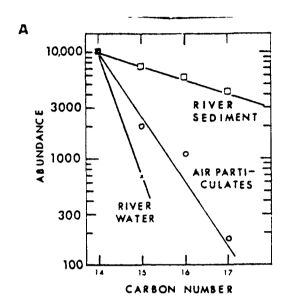
5.2.3 Environmental Fate

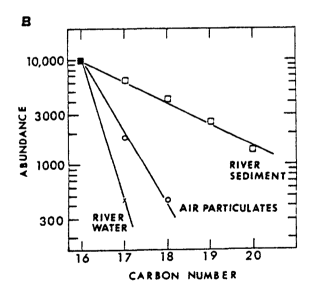
5.2.3.1 Basic Physical/chemical Properties

The physical/chemical properties that are relevant to the fate and distribution of the BaP group PAHs are summarized in Table 5-16. The properties of all of these PAHs, exclusive of acenaphthylene, are quite similar; the properties of acenaphthylene more closely resemble the compounds discussed in Chapters 3.0 and 4.0 of this report.

5.2.3.2 Pathways in the Aquatic Environment

This section examines the details of the fate pathways of BaP in the aquatic environment. The major processes for removal from the water





Source: Hase and Hites (1977).

Note: Data are normalized such that the unsubstituted species is 10,000 units in each case; A-pyrene, B-phenanthrene.

FIGURE 5–5 DISTRIBUTION OF ALKYL HOMOLOGS IN THE PHENANTHRENE-TYPE SERIES AND PYRENE-TYPE SERIES IN RIVER WATER, AND AIR — CHARLES RIVER, BOSTON, MA

TABLE 5-16. BASIC PHYSICAL/CHEMICAL PROPERTIES OF BENZO[#] PYREJE GROUP PARS

	Benzo[a]pyrene	Acenaphthylene	Benz{a] - anthracene	Benzo[b] - Cluoranthene	Benzo[k]- fluoranthene	Benzo[g,h,i]- perylene	Chrysene	Dibenz[a,h]- anthracene	Indeno[1,2,3-c,d]- pyrene
Abbreviation	B⊲P	-	BaA	₩bF	BkP	Bg.h.iPr	Chr	Adabb	11,2,3-c,df
Formula	c ₂₀ 112	€ ₁₂ #8	C18 H12	C20 H12	c ₂₀ w ₁₂	C22 H12	C18 H12		
MW	252. 32 ^{4.}	152. 21 °	228.28	252.32 ^c	252. 32°	276.34°	228.28°	278. 36 ^C	C ₂₂ H ₁₂ 276. 34 ^e
Helting Pt. (°C)	179 ^b	92 ^b	155-157 ^h	167-168 ^b	217 ^b	2 <i>22</i> b	256b	270 ^b	162.5-164 ^h
Boiling Pt. (°C)								-70	10%.7-104
Vapor Pressure (torr)	5.6x10 ⁻⁹ ±(25°C) ^a 5x10 ⁻⁹ (20°C) ^b	0.030(25°C) ^a 16" ³ 10 ² (20°C) ^b	2. 2x10 ⁻⁸ (25°C) ^A 5x10 ⁻⁹ (20°C) ^h	5x10 ⁻⁷ (25°C) ^a 10 ⁻¹¹ -10 ⁻⁶ (20°C) ^b	5x10 ⁻⁷ (25°C) ^a 9. 6x10 ⁻¹¹ (20°C) ^b	ix10 ⁻¹⁰ (25°C)*	6.3×10 ⁻⁹ (25°C) ⁸ 10 ⁻¹¹ -10 ⁻⁶ (20°C) ^b	1.0x10 ⁻¹⁰ (25°C) ^a	1x10 ⁻¹⁰ (25°C) ⁴
Water Solubility mg/t at 25°C	0,0038 ^a	3. y.jh	0.0057(20°) ^d 0.014b 0.009b	0.001	0.000554	0.0003 ^d 0.00026 ^b	0.0018 ⁶ 0.002 ^b	0.0005 b	0.0005 ^a
Log K _{OM} , 25°C	6.08 ^a 6.04 ^b	3. 72 ^a 4. 07 ^b	5.61 a.b	6.ก _{มี} ล 6.57 ^b	6.08 ² 6.84 ^b	6.51 ^d 7.23 ^b	5.61 ^{a,b}	6. 84 ⁴⁸ 5. 97 ^b	6 51 ^a 7.66 ^b
Log K _{Oc} , 25°C		3. 46ª	5. J4 ^a				5.34ª		• •
Henry's law constant at 25°C (atm m ³ /mole)	4.89×10 ⁻⁷	1.52×10 ⁻³	7. 34×10 ⁻⁷	1.66x10 ⁻⁴	3.02x10 ⁻⁴	1.21x10 ⁻⁷	1.05×10 ⁻⁶	7. 3#10 ⁻⁸	bx10 ⁻¹⁰

^d584 (1980). bCallabao et <u>a</u>l. (1979)

Weast (1974),

column are reviewed first, i.e., volatilization and sedimentation; the transformation and degradation pathways for BaP in solution are then described. Finally, biodegradation and its role in determining the ultimate fate of PAHs, especially in sediment, are considered.

Volatilization

Benzo[a]pyrene has a vapor pressure of 5×10^{-9} torr at 25°C and a Henry's law constant of $4.8 \times 10^{-7} \, \frac{\text{m}^3 \cdot \text{atm}}{\text{mol}}$, and these properties suggest that volatilization will not be an important pathway for this compound in the environment. When the role of volatilization in removing BaP and benzo[a]anthracene from the aquatic environment was studied, these two high-molecular-weight PAHs were not rapidly volatilized. With maximum wind (4 m/sec) and current velocity (1 m/sec), half-lives due to volatilization were 150 hours for benzo[a]anthracene and 430 hours for BaP, compared with 3.2 hours for naphthalene and 16 hours for anthracene (Southworth 1979). The variability of volatilization was also examined; for both of these PAHs, a ten-fold increase in current velocity roughly doubled volatilization, while a ten-fold increase in wind velocity increased volatilization five-fold. These larger PAHs appear to be much less sensitive to changes in current velocity and slightly more sensitive to wind velocity changes when compared with lower-molecular weight PAHs.

In a separate study of the environmental fate of these PAHs, the volatilization half-lives were determined to be 89 hours for benz[a]anthracene and 22 hours for BaP under conditions of rapid stirring (Smith et al. 1978). In the same study, volatilization half-lives were calculated using a one-compartment model:

	Volatilization Half-life (hours)						
	River	· •		Oligotrophic			
		Pond	<u>Lake</u>	Lake			
benzo[a]pyrene	140	350	700	700			
benz[a]anthracene	1000	1000	1000	1000			

For these compounds, volatilization is a slow process in comparison with the degradation pathways, such as photolysis, discussed below. Furthermore, most of these larger PAHs will be found sorbed onto sediment, and volatilization of sorbed material is presumed to be very slow (Smith et al. 1978). Another study of fate and transport of radio-labeled BaP, conducted in a laboratory model ecosystem (Lu et al. 1977) failed to detect any radioactivity in traps; this supports the premise that volatilization is not a significant fate process for BaP.

As shown in Section 5.2.3.1, most of the other PAHs in this group exhibit vapor pressures and Henry's law constants that are fairly close to those for BaP and benzo[a]anthracene and would be expected to act similarly with respect to volatilization. Acenaphthylene, on the other hand, will volatilize more rapidly as indicated by its vapor pressure and Henry's law constant.

Adsorption, Sedimentation, and Solubilization

Partition coefficient and solubility data suggest that BaP in the aquatic environment is likely to be adsorbed onto sediment and biota. Concentrations (in rivers, ponds and lakes) predicted from a one-compartment model show that the expected concentrations of BaP and benzo[a]anthracene on sediments and suspended solids are greater than the dissolved concentrations by a factor of more than 10⁴ (Smith et al. 1978). This is expected since the partition coefficients for all of the PAHs in this group range from 5.61 to 7.66, with the exception of acenaphthylene at 3.72.

The patterns of buildup and decline of BaP in pond, river, and lake simulations were also analyzed by Smith et al. (1978). These predictions are shown in Figure 5-6 for a river system. In all aquatic systems modeled, the concentrations of dissolved BaP rapidly approached steady-state before and after the discharge ceased; sediment concentrations changed very slowly (see Figure 5-6). Cessation of the discharge was shown to cause a 100-1000-fold decrease in dissolved BaP concentrations (solution). Continuous desorption from sediments is sufficient to maintain a low concentration of BaP in the water column, roughly equal to the background level in groundwaters (Suess 1976, Andelman and Snodgrass 1974). The authors obtained similar results in modeling of benzo[a]anthracene (Smith et al. 1978).

BaP sorption onto sediments was shown to be strongly correlated with the organic carbon content of the sediment and not related to the cation exchange capacity (Smith et al. 1978). Since calcium montmorillonite clay (0.06% carbon) displayed some affinity for BaP, it seems likely that BaP sorption involves weak interaction with solid surfaces, as well as strong binding to organic matter. Sorption onto bacterial cells was also investigated, and very high sorption coefficients were obtained. These results are in agreement with other data indicating that PAHs will preferentially sorb onto organic and biological material (Southworth 1977). Sorbent data and partition coefficients for BaP are presented in Table 5-17.

It appears that very little of the BaP in aquatic systems will be found in solution, and this compound can be expected to accumulate in the sediments of placid lakes and reservoirs. River-borne BaP, however, will be transported to the ocean, where inshore and alongshore currents combine to restrict suspended matter to continental areas and spread the PAHs almost uniformly along the coastal regions. In these areas, continuous resuspension and transport via wave action and currents have

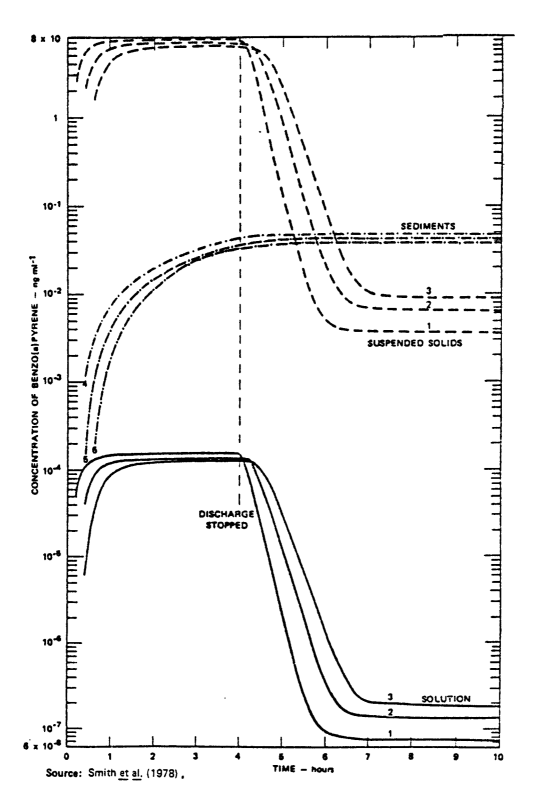


FIGURE 5—6 RESULTS OF MODELING OF THE EFFECT OF SUSPENDED SOLIDS ON THE CONCENTRATION OF BENZO[a] PYRENE IN A PARTIALLY MIXED RIVER SYSTEM

TABLE 5-17. BENZO[a]PYRENE PARTITION COEFFICIENTS FOR VARIOUS SORBENTS

Sorbent	Total Organic Carbon (%)	Carbon Exchange Capacity (meq/100g)	Partition Coefficient Kp
Ca-montmorillonite Clay	0.06	69	1.7x10 ⁴
Des Moines River Sediment	0.6	10.5	3.5x10 ⁴
Coyote Creek Sediment	1.4	13.5	7.6x10 ⁴
Searsville Pond Sediment	3.8	34.5	1.5x10 ⁵
Mixed Bacteria (dry wt. equivalent to 97 mg/1)			3-4×10 ⁴

Source: Smith et al. 1978.

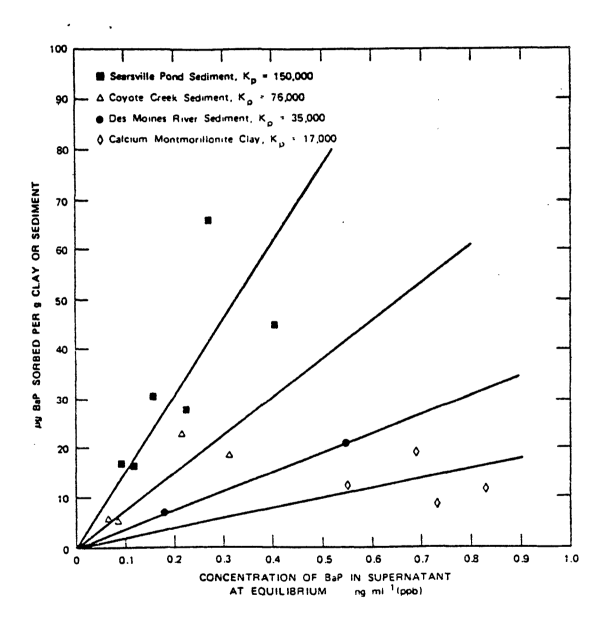
been observed (Gross 1970). PAHs have been identified in coastal waters adjacent to or distant from developed areas (Mallet 1961); coastal plankton have also been found to contain significant amounts of BaP, whereas plankton from the open ocean were uncontaminated (DeLima-Zanghi 1968). Both of these observations indicate that physical transport along the coast is an important fate process for adsorbed PAHs.

The fate of adsorbed PAHs in the water column is influenced by a number of factors; duration of PAH exposure to sunlight will largely determine the extent of photo-oxidation (discussed below). The duration of exposure is partially controlled by particulate sedimentation and resuspension. It has been estimated that estuarine sediment a few millimeters deep is recycled through the water column daily, and that the top 2 cm are recycled annually. Particulates less than 0.5 mm in diameter may reside in the water column for 200-600 years. It has also been postulated that it will take 500-1000 years to bury a single layer of particulate. These data are relevant to determining the fate of adsorbed PAHs since much of this material will ultimately find its way to the coastal regions (Gross 1970).

The low solubility and high $K_{\rm OW}$ for BaP do not necessarily mean that all BaP is sorbed; many natural waters and industrial effluents also contain appreciable amounts of organic material that may increase the solubility of BaP. PAHs could be solubilized by incorporation into micelles if a critical micellar concentration is reached. The concentration of detergents, such as linear alkylbenzene sulfonate, must reach 10-50 mg/l before significant solubilization of BaP occurs; detergent concentrations of 0.1-1.0 mg/l, which are probably closer to environmental concentrations, have no effect on BaP (Bohm-Gasol and Kruger 1965, II'nitskil and Ershova 1970, II'nitskil et al. 1971).

Another mechanism for solubilization of PAHs in water exists as a result of the presence of other organic compounds not associated with colloid or micelle formation. With some specific organic mediators, such as butyric and lactic acids, organic solvents, and nitrogencontaining organics, the extent of solubilization has been reported to be proportional to the concentration of the mediator or the square of its concentration (Neff 1979). However, naturally occurring dissolved organic matter (humic acids, fulvic acids, etc.) in seawater appeared to have no effect on solubility of phenanthrene and anthracene (Boehm and Quinn 1973) and may have little effect on PAH solubility, in general.

Sorption of PAHs and other chemicals to natural sediments and biogenous materials has been shown to be a reversible equilibrium process (Smith et al. 1978). Elevated particulate PAH concentrations are generally accompanied by elevated concentrations of dissolved PAHs as shown in Figure 5-7 for BaP. These data do suggest an exchange between adsorbed and dissolved states, and further indicate that there may be a significant amount of BaP in solution, particularly in heavily contaminated systems.



Source: Smith et al. (1978).

FIGURE 5-7 SORPTION ISOTHEMS FOR BENZO[a] PYRENE

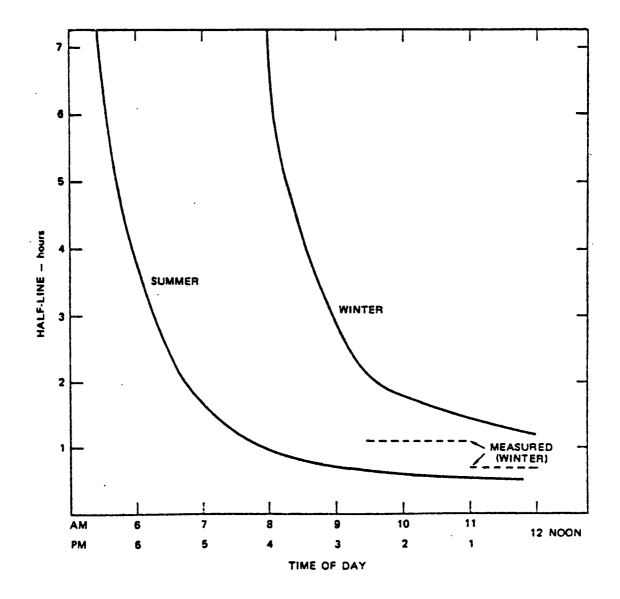
The larger-molecular-weight PAHs discussed in this section, particularly BaP, will tend to adsorb onto particulate matter and will ultimately accumulate in the sediment due to sedimentation; half-lives in sediments are expected to be on the order of a few years (White and Vanderslice 1980). Chemical degradation pathways for BaP dissolved in aquatic environments are discussed below; however, these are generally less important as fate processes than sedimentation.

Chemical Degradation

In oxygenated water, photolysis of BaP by light with wavelengths in the solar region gives a mixture of three quinones, as shown below.

Several authors (Andelman and Suess 1970, Smith et al. 1978, NAS 1972, Stevens and Algar 1968, Neff 1979) have suggested that direct photo-oxidation of PAHs is a major oxidation pathway in water. This reaction is postulated to be mediated by singlet oxygen formed by energy transfer from the electronically excited aromatic molecule in its triplet state. However, contrasting evidence has recently been reported (Zepp and Scholtzhauer 1979), suggesting that neither molecular oxygen nor singlet oxygen is the exclusive mediator in any of the major photochemical reactions of the PAHs.

The half-life for direct photolysis of BaP in sumlight, as a function of the time of day was calculated by the procedure of Zepp and Cline (1977) using a quantum yield of 8.9×10^{-4} and the measured UV spectrum of BaP (Smith <u>et al.</u> 1978). Their results are presented in Figure 5-8. (These data were obtained from BaP in a solution of 20% acetonitrile in pure water.) The calculated half-life of 1.2 hours for midday photolysis in winter is in close agreement with the measured results of 1.1 hours and 0.7 hours. Similar experiments with benz[a]—anthracene yielded a half-life of 2 hours for that compound.



Source: Smith et al. (1978).

FIGURE 5-8 CALCULATED SEASONAL AND DAILY VARIATION OF PHOTOLYSIS HALF-LIFE OF BENZO[a] PYRENE

Benzo[a]pyrene in natural waters or pure water containing humic acid exhibited slower rates of photolysis than in similar experiments conducted in pure water (Smith et al. 1978). The rate of photolysis of BaP in the presence of humic acid was five times slower than photolysis in pure water; natural water caused an intermediate, but definite retardation of BaP photolysis. A light-screening effect is probably not entirely responsible since absorbance of natural waters at 366 nm is less than 0.02: the 0.11 absorbance of the humic acid solution should only reduce the photolysis rate by 13%. The authors speculated that substances present in natural waters may quench a BaP excited state, singlet oxygen, or any excited state complex occurring in the reaction. Another possible cause is formation of a complex of BaP with natural organic or inorganic substances in solution, which may alter the reactivity of ground state BaP itself. Evidence for this mechanism was found in that only 50% of the 270 ng/ml of BaP in solution with 30 $\mu g/ml$ of humic acid could be recovered from aqueous solution by hexane extraction (Smith et al. 1978).

Inhibition of photolysis was also observed when BaP was adsorbed onto Kaolinite clay (McGinnis and Snoeyink 1974); products formed during photolysis are thought to be responsible. Mechanisms by which such effects occur may include competitive reactions of the oxidizing agent(s) formed with the organic matrix. Similar results were obtained in studies of BaP dissolved in acetone and adsorbed onto calcium carbonate (Andelman and Suess 1970).

In another study, BaP adsorbed onto calcite particles in water was exposed to illumination roughly equivalent to one-fourth of the winter solar radiation at the earth's surface (Suess 1972). Half-lives of BaP were 15 hours when the water sample was exposed to atmospheric air, 11 hours under oxygen atmosphere, and 35 hours under helium atmosphere. These results indicate a dependence of BaP degradation on oxygen concentration in water; no oxidation was observed in the absence of sunlight. Again, all of these half-lives are significantly longer than the photolysis half-life of 1-2 hours predicted above.

Benzo[a]pyrene suspended in the water column as a condensed particulate was shown to decompose rapidly under normal daylight conditions (McGinnis and Snoeyink 1974), the rate of decomposition being governed by particle size. For particulates 1.5 mm in diameter, the reaction exhibited first-order kinetics until 55-65% of total BaP was decomposed; at this point, a residual was left that was not affected by an increase in radiant energy. Apparently the decomposition products formed a protective barrier, preventing further reaction of the residual BaP. The authors postulated that in particulate of greater than 0.4-mm diameter a residual would remain. Benzo[a]anthracene did not exhibit this effect; after a threshold of sunlight had been attained, the reaction went to completion, presumably due to the solubility of the decomposition products.

The free radical oxidation of BaP was studied under experimental laboratory conditions at 50°C in order to obtain a first-order rate constant of 5.7×10^{-5} sec⁻¹ (Smith et al. 1978). Extrapolating these data to 25° C, this rate corresponds to a second-order rate constant of 1.86×10^{-3} M⁻¹ sec⁻¹; using Smith's estimate of 10^{-10} M RO2 • the half-life was calculated to be 4.3 days (~ 100 hours). This half-life is considerably shorter than that estimated by Radding et al. (1976). However, the fact that both are relatively long suggests free radical oxidation of BaP is not competitive with photolysis and adsorption under environmental conditions.

A number of authors have studied the oxidation of PAHs by chlorine (Perry and Harrison 1977) and ozone (Il'nitskii et al. 1968) and the data have been summarized by Radding et al. (1976). The data in Table 5-18 indicate that oxidation by chlorine and ozone may be significant fate processes when these oxidants are available in sufficient quantities, such as in water treatment plants. The products of aqueous chlorination of PAH solutions are not fully known.

Predictions of the fate of BaP in natural waters, based on studies performed under laboratory conditions using pure water or organic solvents, must be made with caution since there may be numerous factors affecting the photo-oxidation rate. The half-lives for individual transformation and removal processes were calculated for various water systems using a one-compartment model (Smith et al., 1978), and are shown in Table 5-19. Chemical degradation due to photolysis is clearly the major transformation pathway for BaP in solution, since the half-lives are at least an order of magnitude smaller than the half-lives for other pathways. Sorption rates were not measured, but were postulated to be at least 100 times faster than photolysis rates; therefore, it is probable that most of the BaP is removed to the sediments.

Biological Fate

Analyzing the exposure and bioavailability of PAHs to aquatic organisms requires an examination of the disposition of these compounds in the biological compartments of the environment. The considerations discussed here include bioaccumulation in aquatic organisms, both as seen in the laboratory and in the field, biotransformation, and biodegradation.

Bioaccumulation

The uptake and bioaccumulation of BaP have been investigated by several authors, using both laboratory model ecosystem studies and predictions based on the octanol:water partition coefficient ($K_{\text{OW}} ~ \% ~ 6$) and water solubility (0.0038 mg/l of BaP. A wide range of bioconcentration data has resulted; some of the data are presented in Table 5-20. As shown, there are significant differences in bio-

TABLE 5-18. RELATIVE HALF-LIVES OF BENZO[a] PYRENE GROUP PAHS IN REACTION WITH MAJOR OXIDANTS

	Half-life (hours) in Reaction w				th	
Compound	10 ⁻¹⁰ M	Singlet Oxygen	Ozone (water) 10 ⁻⁴ M	Ozone (air) 2x10 ⁻⁹ M	c1 ₂	110*
Benzo[a]pyrene	2.4 x 10 ⁵	5	1.05	870		
Benz[a]anthracene		10	0.45	370	t 1/2 ∿ 0.5hr for	t _{1/2 ∿ 10 hr.}
Dibenzanthracene ^a			0.42	340	all PAHs	all PAHs

Source: Radding et al. (1976).

a Isomer not specified.

TABLE 5-19. PREDICTED HALF-LIVES FOR BENZO[a]PYRENE TRANSFORMATION AND REMOVAL PROCESSES IN GENERALIZED AQUATIC SYSTEMS

Process	Half-life (hours)				
	River	Eutrophic Pond	Eutrophic Lake	Oligotrophic Lake	
Photolysis	3.0	7.5	7.5	1.5	
Oxidation	>340	>340	>340	>340	
Volatilization	140	350	700	700	
Biodegradation	>104	>104	>104	>104	
Hydrolysis	NA	NA	NA	NA	

Source: Smith et al. (1978).

TABLE 5-20. BIOCONCENTRATION OF BENZO[a]PYRENE IN FRESH-WATER AND SALTWATER SPECIES

Species	Duration	Bioconcen- tration Fact	or Reference		
	Freshwat	er Species			
Alga, Oedogonlum cardiacum	3 days	5,258 ^a	Lu <u>et al</u> . (1977)		
Snail, Physa sp.	3 days	82,231 ^a	Lu <u>et al</u> . (1977)		
Cladoceran, <u>Daphnia</u> <u>pulex</u>	3 days	134,248 ^a	Lu <u>et al</u> . (1977)		
Mosquito, <u>Culex pipiens</u> <u>quinquefasciatus</u>	3 days	11,536 ^a	Lu et al. (1977)		
Mosquitofish, Gambusia affinis	3 days	930 ^a	Lu <u>et al</u> . (1977)		
Saltwater Species					
Clam, Rangia cuneata	24 hours	8.66	Neff <u>et al</u> . (1976a)		
Clam, Rangia cuneata	24 hours	236	Neff <u>et al</u> . (1976b)		
Eastern oyster, Crassostrea virginica	14 days	242	Couch <u>et al</u> . (in press)		
Mudsucker Gillichthys mirabills	96 hours	0.048	Lee <u>et al</u> . (1972)		
Tidepool sculpin, Oligocottus maculosus	1 hour	0.13	Lee <u>et al</u> . (1972)		
Sand dab, Citharichthys stigmacus	1 hour	0.02	Lee <u>et al</u> . (1972)		

a Model ecosystem concentration factor.

concentration factors (BCFs) among species and over time, BaP accumulated to high levels (BCFs of 900-134,000) over 3 days in freshwater organisms, but to much lower levels $(0.02-242\ BCF)$ in marine biota in less than 3 days.

The rates of BaP uptake and depuration have been investigated in a microcosm study using freshwater insect larvae (midges), Daphnia, periphyton (diatoms), and bluegill sunfish. Organisms were exposed to C^{14} -labeled BaP at a concentration of 1.0 μ g/l for 8 hours, and the rate of depuration was determined. In periphyton, accumulation was linear over time, surface-area dependent, and diffusion-limited. Daphnia had high uptake and depuration rates, perhaps due to their large surface area/volume ratio; very little biotransformation occurred in Daphnia, however. In the midges (Chironomus riparius), greater than 50% of the BaP was transformed into polar compounds in 1 hour, and 10% of the accumulated BaP was associated with the exo-skeleton. Bioconcentration factors derived for these insects were 950 (steady state) and 150 (based on BaP analysis). Half-lives of BaP in the four species tested ranged from 5.3-8.5 days (Leversee et al. 1980). From these and several other studies, there is evidence that considerable variation exists in uptake and depuration rates and in the levels found in various types of tissues, even with the same species and compound.

Several investigations of the accumulation of BaP by biota from sediment and water in the natural environment have been reported. Experiments on the uptake of hydrocarbons, including BaP, from contaminated sediments by marine organisms showed that deposit feeders (the clam Macoma inquinata and sipunculid Phascolosoma agassizii) generally accumulated the compound to a greater extent than the suspension feeder (the clam Protothaca staminea). Experiments with the deposit-feeding clam M. inquinata, however, indicate that compounds directly associated with particulate matter in sediment were less available for uptake than those released from sediment in the surrounding seawater. Concentration factors for uptake from sediment were ≤ 0.2 , while those from seawater were 10-1349. This work indicated that the PAHs present in interstitial water may be a prime source of contamination for benthic infauna and probably accounted for much of the uptake by the sediment-feeding organisms (Roesijadi et al. 1978).

Numerous studies have examined the PAH content of aquatic organisms in relation to the environment in which the organisms reside. Data are presented in Table 5-21 that show the range of BaP levels in various species from different locations. In general, but not always, organisms sampled from coastal regions near major industrial or domestic point sources of PAH (e.g., Norfolk, VA harbor) contained higher BaP concentrations than organisms from more remote areas (Neff 1979). Benzo[a]pyrene concentrations in various aquatic organisms from several locations in the U.S. and southern Canada ranged from approximately <0.1 µg/kg (dry weight) to as high as 5000 µg/kg. In most cases, the organisms contained low (0-20 µg/kg range)

TABLE 5-21. CONCENTRATIONS OF BENZO[a]PYRENE IN TISSUES OF AQUATIC ORGANISMS

Organism	Location	BaP (µg/kg dry wt)	Reference
Cod (Gadus sp.)	Holsteinborg, Greenland Atlantic, 40 km off Toms River	15	Mallet <u>et al</u> . (1963)
	New Jersey, USA	< 10	Pancirov and Brown (1977)
Menhaden Anchovy and smelt	Raritan Bay, New Jersey, USA San Diego Bay, California, US	6.0 SA up to 5000	Pancirov and Brown (1977) Lee <u>et al</u> . (1972)
Flounder (unidentified)	Long Branch, New Jersey, USA South of Long Island, New Yor	< 8 •k	Pancirov and Brown (1977)
	USA, 40°27'N 73°06'W	< 4	Pancirov and Brown (1977)
Shrimp (Penaeus aztecus)	Palacios, Texas, USA	< 4	Pancirov and Brown (1977)
Clam (unidentified)	Chincoteague, Virginia, USA Darien, Connecticut, USA	1.2 <0.4	Pancirov and Brown (1977) Pancirov and Brown (1977)
Crab (unidentified) Crab (<u>Cellinectes sapidus</u>)	Raritan Bay, New Jersey, USA Chesapeake Bay, Virginia, USA	12 <2.0	Pancirov and Brown (1977) Pancirov and Brown (1977)
Soft-shell clam (Mya arenaria)	Tillamook Bay, Oregon, USA Alsea Bay, Oregon, USA	1.2	Mix et al. (1977) Mix et al. (1977)
Butter clam (Saxidomas glganteus)	Coos Bay, Oregon, USA Coos Bay, Oregon, USA	1.32-26.64 1.16- 4.20	Mix <u>et al</u> . (1977) Mix <u>et al</u> . (1977)
Oyster (<u>Crassostrea virginica</u>)	Norfolk Harbor, Virginia, USA	20-60	Cahnmann and Kuratsune (1957)
Crassostrea gigas Gaper clam (Tresus capax)	Long Island Sound, USA Chincoteague, Virginia, USA Tillamook Bay, Oregon, USA Tillamook Bay, Oregon, USA Netarts Bay, Oregon, USA Yaquina Bay, Oregon, USA Alsea Bay, Oregon, USA Coos Bay, Oregon, USA	8.0 1.2 <0.4 0.64-12.8 <0.4 1.0-2.24 0.4 0.56-2.04	Pancirov and Brown (1977) Pancirov and Brown (1977) Mix et al. (1977)

TABLE 5-21. CONCENTRATIONS OF BENZO[a]PYRENE IN TISSUES OF AQUATIC ORGANISMS (Continued)

		BaP			
Organism	Location	(µg/kg dry wt)	Reference		
Mussel					
Mytilus edulis	Falmouth, Massachusetts US	SA			
The state of the s	Little Sippewissett	<2	Pancirov and Brown (1977)		
	Wild Harbor	2	Pancirov and Brown (1977)		
	Tillamook Bay, Oregon, USA	-	Mix et al. (1977)		
	Yaquina Bay, Oregon, USA	0.48-120.8	Mix et $\frac{1}{a1}$. (1977)		
	Alsea Bay, Oregon, USA	<0.4	Mix et al. (1977)		
	Vancouver, B.C., Canada	• • • • • • • • • • • • • • • • • • • •	Dunn and Stich (1975)		
	outer harbor	8 + 1.2	baill and Selen (1773)		
	wharf, marina and dock are	rana			
	inner harbor	168 + 24			
	Vancouver marina area				
	(May 174)	7.6 + 0.4			
	(Sept '74)	7.2 + 0.4			
	wharf area on rock and cables				
	(May '74)	52 + 21.2			
	(Sept '74)	156 + 29.2			
	creosoted pilings				
	(May '74)	272 + 52			
	(Sept 174)	532 + 76			
M. edulis and M. californeanus	25 stations between				
	Bodega Head and San Diego,				
	California, USA	<0.4-32.8	Dunn and Young (1976)		
M. californeanus	West coast of Vancouver		Sam and Touris (1770)		
	Island, British Columbia, Canada	0.4 ± 0.4	Dunn and Stich (1975)		

BaP concentrations, except for animals collected from severely polluted areas or from the immediate vicinity of marina (creosoted pilings) or fish processing facilities (Mix et al. 1979). Fish from San Diego Bay, a large shipping port, contained up to 5000 μ g/kg BaP (dry weight) (Lee et al. 1972).

Several species of bivalve mollusks (clams) from Oregon estuaries were sampled for levels of BaP. The maximum concentrations found were in the range of 15-30 ng/g (ppb). However, specimens from Coos Bay, the most heavily industrialized bay on the Oregon Coast, did not contain significant levels of BaP overall, except for the softshell calm $\underline{\text{Mya}}$ arenaria (Mix et al. 1977).

One report of BaP accumulation by mussels following an oil spill in a coastal area indicated that after 11 days, $\underline{\text{Mytilus}}$ edulis contained approximately 55 µg/kg BaP (Bories et al. 1976).

Biotransformation

The process of biotransformation and the resultant metabolites of BaP and other PAHs in aquatic organisms have been studied fairly extensively. It is known that enzymatic activity in the excretory systems -i.e. hepatopancreas, intestine, etc -- of these organisms, facilitates the metabolizing of PAHs and excretion of water-soluble products. The extent to which animals are able to remove PAHs enzymatically from their tissues varies among species. Some marine organisms, primarily the invertebrates including starfish, sea anemones, oysters, and ctenophores, have been found to have very low or non-existent enzyme activity related to hydrocarbon metabolism. One explanation may be that organisms that pass large volumes of water across their tissues have not developed the enzyme mechanisms important in the toxification/detoxification system (Lee et al. 1977). Most of the teleost fish studied so far (including flounder, mackerel, trout, salmon, cunner, and sheepshead) and some members of the Phyla Arthropoda and Annelida are able to metabolize PAHs to polar compounds (Neff 1979). Such metabolic activity has not been detected in several species of marine algae (Payne 1977).

Microbial Biotransformation

Some of the members of the BaP group of PAHs are subject to microbial breakdown; on the whole, though, biodegradation is slower and not as extensive in this group as in the lower-molecular-weight PAHs. In a comparative study of the relative biodegradability of approximately 42 different polynuclear aromatic hydrocarbon compounds, McKenna and Heath (1976) found no significant biodegradation of compounds with more than three rings.

Within the BaP group, biodegradation rate data were available for only BaP, benz[a]anthracene, chrysene, and dibenz[a,h]anthracene. Within this group, the rate and extent of degradation is variable. Since structural factors such as ring number and molecular geometry influence biodegradation, there is danger in extrapolating from even relatively similar compounds to compounds for which no data are available. Other influences include environmental fate characteristics of the compounds, such as adsorption, solubility, vaporization, and other competitive transformation reactions (if any). Most of the biodegradation studies on PAHs are laboratory investigations, commonly conducted in simplified systems with the goal of eliciting biodegradation. Under environmental conditions, persistence may be longer than that measured in laboratory studies due to influences of external factors usually controlled for in the laboratory. As noted in Section 3.2.3.4, these environmental factors may include availability of oxygen, soil solution pH, and the presence of natural humic polymers.

Microorganisms act on PAHs by removing one cyclic unit at a time (Alexander 1977). This is supported by an observed inverse relationship between ring number and degradation rate. Presumably a process analogous to that described for anthracene (Section 4.2.3.4) (Alexander 1977) occurs in the breakdown of the BaP group.

Presented in Table 5-22 are reported biodegradation products derived from the breakdown of two members of the group: BaP and benz[a]anthracene. The products are primarily intermediates in the multistep pathway of reactions leading to eventual complete mineralization. Gibson et al. (1975) isolated a mutant strain of the bacterium, Beijerinckia, which converted benz[a]anthracene to four isomeric dihydrodiols. (However, this study had limitations. For example, it was not possible to confirm whether the first step in biodegradation was removal of one ring from the molecule.) No other data were available on other group members.

It must be mentioned, however, that other factors such as substituent type and size, interfere with this relationship. Malaney did not find a direct relationship between ring number and biodegradability (Malaney et al. 1967).

TABLE 5-22. BIODEGRADATION PRODUCTS REPORTED FOR THE BENZO[a]PYRENE GROUP PAHS

PAH

Benzo[a]pyrene^a

Benz[a]anthracenea

Degradation Products

cis-9,10-dihydroxy-9,10-dihydrobenzo[a]pyrene^b

cis-1,2-dihydroxyl-1,2-dihydrobenzo[a]anthraceneb

a Fungi.

 $^{\mathrm{b}}$ Tentative identification.

Source: Gibson (1976).

The rate of biodegradation is quite variable for the BaP group of PAHs. Table 5-23 presents quantified rates of biodegradation reported for the BaP group in soil, freshwater, and estuarine systems. Due to the variety of test methods, analytical techniques, microbial species, and data analyses used in biodegradation testing, it is difficult to compare the results from the different tests reported in Table 5-23.

A total of nine studies was available on BaP; five of these studies found no degradation, two found minimal degradation, and two found relatively rapid degradation, as much as 86% in 5 days. Degradation of acclimated populations was generally faster than that of unacclimated ones. The addition of naphthalene, or especially phenanthrene, significantly increased biodegradation (McKenna and Heath 1976), apparently due to cometabolism. The results of five studies on benz[a]anthracene are similar to those found for BaP; in some cases no degradation occurred, but in one study as much as 41% was lost in 1 week. The addition of other PAHs again resulted in an increase in degradation. The one study available on chrysene found a moderate rate of decay, as much as 59% in 1 week. The sole study on dibenz[a]anthracene found negligible degradation for the compound alone, but a significantly higher rate in the presence of phenanthrene (McKenna and Heath 1976). No biodegradation data were available for acenaphthylene, benzo[b]fluoranthene, benzo[k]fluoranthene, benzo[g,h,i]perylene, or ideno[1,2,3-c,d]pyrene.

A study on biodegradation of BaP and benz[a]anthracene in estuarine sediment populations found that the presence of a polychaete worm species increased the rate of degradation, possibly due to its role in sediment mixing or metabolism of the substances. Degradation was greatest in populations associated with large-grain-size sediment and higher in the surface than in subsurface sediment layers. Approximately 50% of the initial benz[a]anthracene concentration was degraded after 30 weeks in fine sand without the polychaete worm. No long-term measurements were reported for BaP (Gardner et al. 1979).

There is other evidence that BaP and benz[a]anthracene are not readily degradable by microbial populations. Biological treatment in wastewater treatment is reported to be ineffective at removing PAHs from waste streams (Andelman and Snodgrass 1974). Smith $\underline{\text{et}}$ al. (1978) were unable to isolate enrichment microbial cultures capable of degrading either compound; however, the authors allowed that under environmental conditions such microbes may exist.

The turnover time and transformation rate of PAHs in both acclimated and unacclimated stream populations (Schwall and Herbes 1978) provide a good estimation of the environmental persistence of these compounds, as well as the importance of microbial adaptation (see Table 5-24). The turnover time in acclimated populations ranged from 417 days for benzo[a]anthracene to more than 3.5 years for BaP. Turnover times in non-acclimated populations were significantly greater by at least an

TABLE 5-23. BIODEGRADATION RATES OF THE BENZO[a]PYRENL GROUP PAHs: INDIVIDUAL COMPOUND STUDIES

Test Type/Population Origin	Compound Tested	Results	Reference
Static flask (wastewater population)	Benz[a]anthracene	Inconsistent degradation over month period of acclimation from 0% degraded to 41% degraded in one week at 5 mg/l	Quave <u>et al</u> . (1980)
Static flask (wastewater population)	Chrysene	59% lost at 5 mg/l and 38% at 10 mg/l at one week in acclimated culture	Quave <u>et al</u> . (1980)
Freshwater populations - enrichment shake flask, also using naphthalene in culture	Benzo[a]pyrene	No degradation observed in 6-week period	Colwell and Sayler (1978)
	Benz[a]anthracene	No degradation observed in 6-week period	Colwell and Sayler (1978)
Adapted soil populations of Pseudomonas aeruginosa and Escherischia coli	Benzo[a]pyrene	90% taken up from medium, 10-26% metabolized	Lorbacher <u>et al</u> . (1971)
Salmonella typhimurium, Aerobacter aerogenes, Escherischia coli, Saccharomyces cerevisiae	Benzo[a]pyrene	Species accumulate compound but little metabolized. Can take up as much as 1 to 2 x 10^{-10} µg/cell (E. coli).	Moore and Harrison (1965)
Mycobacterium flavum M. rubrum, M. lacticolum, M. smeginatis, Bacillus megaterium, Bacillus sphaericus	Benzo[a]pyrene	M. rubrum and M. flavum metabolized 50% of compound in 4 days. Other species accumulated the compound (no mention of biodegradation)	Poglazova, et al. (1966, 1976a,b)

TABLE 5-23. BIODEGRADATION RATES OF THE BENZO[a]PYRENE GROUP PAHs: INDIVIDUAL COMPOUND STUDIES (Continued)

Test Type/Population Origin	Compound Tested	Results		Reference
Coastal estuary sediment populations (3 types) with	Benzo[a]pyrene Benz[a]anthracene		% removed in 1 week	Gardner <u>et al</u> . (1979)
and without presence of polychaete worm, Capitella		Experiment	BaP BaA	
capitata		Fine sand	1.2 1.5	
		Fine sand & <u>C. capitata</u>	2.4 2.7	
		Med. sand	1.4 1.8	
		Med. sand & <u>C</u> . <u>capitata</u>	3.0 3.0	
		Marsh sed.	0.84 1.4	
		Marsh sed.		
		& C. capitata	1.98 1.8	
Soil bacteria from benzo- pyrene contaminated area and from non-contaminated area	Benzo[a]pyrene	Acclimated population (75-86% of compound in non-acclimated populatin same period	n 5 days;	Shabad (1978) Shabad (1971a,b) Shabad <u>et al</u> . (1971b)
Bacteria in power plant and coke over wastewater	Benzo[a]pyrene	Metabolized <15% of c	ompound	Poglazova <u>et al</u> . (1972)
¹⁴ CO ₂ evolution with sea water population from treated area	benz[a]anthracene henzo[a]pyrene	Not degraded Not degraded		Lee <u>et al</u> . (1978)

TABLE 5-23. BIODEGRADATION RATES OF THE BENZO[a]PYRENE GROUP PAHs: INDIVIDUAL COMPOUND STUDIES (Continued)

Test Type/Population Origin	Compound Tested	Results	Reference
CO ₂ evolution from contaminated stream sediment population	$[C^{14}]$ benz[a]anthracene $[C^{14}]$ benz[a]pyrene	10 ⁻⁴ /h No measurable transformation in 26 days	Schwall and Herbes (1978)
Shake flasks with natural water populations		Percent main compound (column 2) remaining at 4 weeks	McKenna and Heath (1976)
		+ naphthalene + phenanthrene	
	Benzo[a]pyrene	83.5 38.3	
	Benz[a]anthracene	58.3 33.8	
	Dibenz[a,h]anthracene	92.7 32.9	
		Negligible degradation was observe for each compound alone.	ed

TABLE 5-24. KINETIC PARAMETERS OF BIOTRANSFORMATION OF BENZ[a]ANTHRACENE AND BENZO[a]PYRENE

	Rate Constant k (1/h)		Turnover Time ^a		Transformation Rate (µg/g/hr)	
Compound	Contaminated	Uncontaminated	Contaminated	Uncontaminated	Contaminated	Uncontaminated
Benz[a]anthracene	1.0×10^{-4}	4.0×10^{-6}	417 days	28.5 years	1.2×10^{-5}	$<4 \times 10^{-8}$
Benzo[a]pyrene	$<3 \times 10^{-5}$	$<3 \times 10^{-5}$	3.5 years	57 years	$<2 \times 10^{-6}$	$<3 \times 10^{-7}$

a_{Turnover rate} = 1/k

Source: Schwall and Herbes (1978).

 $b_{Transformation rate = k \times concentration}$

order of magnitude: an estimated 28.5 years and 57 years, respectively. How applicable these aquatic turnover times are for soil systems is unknown. However, the turnover time for the acclimated population could be comparable to that of a well-acclimated soil population.

In another small-scale biodegradation study using acclimated stream populations, the degradation time of PAHs increased 30-100 times for each additional ring from naphthalene to benz[a]anthracene (Schwall and Herbes 1978). In a static flask study of several PAHs (Quave et al. 1980), a considerable variation was observed over a four-ring range. Acclimation was required for all but benz[a]anthracene; for some compounds 10 mg/l of the compound was inhibitory (Quave et al. 1980).

Thus the benzo[a]pyrene group of PAHs is relatively resistant, as a class, to biodegradation, especially in comparison with lower-ringed aromatics. Most biodegradation studies are too short-term to quantify the half-lives for persistence of most of these PAHs. In some cases, half-lives on the order of weeks were reported; however, in general, PAHs with three rings or more were presistent on the order of months or longer. No information at all was available for acenaphthylene, benzo[b]fluoranthene, benzo[k]fluoranthene, benzo[g,h,i]perylene, and ideno[1,2,3-c,d]pyrene.

5.2.3.3 Modeling of Environmental Distribution

Mackay Model

The Mackay equilibrium partitioning model (Mackay 1979) described in Section 3.2, was utilized for BaP. The values for the input parameters used in the Mackay model and the predicted distribution of BaP are given in Tables 5-25 and 5-26. Benzo[a]pyrene has a large adsorption coefficient for soil or sediment and a very low water solubility and vapor pressure. These characteristics suggest very little partitioning into the air or water with most of the BaP concentrated in the sediments. The differences in partitioning among the solid phases are a result of the relative amounts of each in the model ecosystem. The concentrations of material in the air and water compartments are significantly lower than those in the solid phases, while the concentration in the aquatic biota is somewhat higher. The number of moles predicted to reside in each compartment is directly proportional to the total number of moles in the whole system.

The Mackay model predicts that 99.9% of the BaP load to the system will reside in the sediment compartment. Less than 0.1% will be present in the water column, and less than 0.001% will partition to the atmosphere. Dibenz[a,h]anthracene, benz[a]anthracene, benzo[g,h,i]-perylene, indeno[1,2,3-c,d]pyrene all have physical properties similar to

TABLE 5-25. VALUES OF PARAMETERS USED FOR CALCULATING THE EQUILIBRIUM DISTRIBUTION OF BENZO[a]PYRENE PREDICTED BY THE MACKAY FUGACITY MODEL

Chemical-Specific Parameters (25°C):

Henry's Law Constant (atm m ³ /mole) Adsorption Coefficients:	4.8×10^{-7}
Suspended Solids Sediment Biota (0.2 x K _{OW})	1.26×10^{4} 1.26×10^{4} 2.2×10^{5}
Total Amount in System (kg)	11 ^a
Compartment-Specific Parameter (25°C):	
Air	
area depth volume	$1 \times 10^4 \text{ m}^2$ $3 \times 10^3 \text{ m}$ $3 \times 10^2 \text{ m}^3$
Water	
area depth volume biomass content suspended solids	$1 \times 10^4 \text{ m}^2$ 2 m $2 \times 10^4 \text{ m}^3$ 129 mg/1

Sediment

suspended solids

30 mg/1

^aThe load to the MacKay model was assumed to be equal to the total pond accumulation predicted by EXAMS.

TABLE 5-26. EQUILIBRIUM PARTITIONING OF BENZO[a]PYRENE CALCULATED USING MACKAY'S FUGACITY MODEL

	Partitioning at Equilibrium				
Compartment	Moles	Concentration	Percent		
Air	2.95×10^{-4}	$2.48 \times 10^{-4} \mu \text{g/m}^3$	0.007		
Water	1.02×10^{-2}	0.13 µg/l	0.0235		
Suspended Solids	3.88×10^{-3}	1.628 mg/kg	0.0089		
Sediment	43.65	1.628 mg/kg	99.89		
Aquatic Biota	2.91×10^{-2}	28.38 mg/kg	0.0666		
Biotic Sediment	2.82×10^{-3}	28.38 mg/kg	0.0065		

those of benzo[a]pyrene, and would be expected to be partitioned into the various environmental compartments in the same manner. The benzo-fluoranthenes have larger vapor pressures and Henry's law constants, suggesting they might partition into air to a somewhat greater extent than the other compounds in this group. Acenaphthylene has physical properties more closely resembling naphthalene and anthracene, and atmospheric partitioning for this compound would be expected to be greater than for BaP.

EXAMS

The U.S. EPA EXAMS (Exposure Analysis Modeling System) model (U.S. EPA 1890b), described in Section 3.1 was run for BaP. All six EXAMS environments (pond, eutrophic lake, oligotrophic lake, river, coastal plain river and turbid river) were modelled using the ecosystem parameters that are provided with the model. The input parameters for BaP are given in Table 5-27. Any variables not in the table were set to zero, since these variables (such as hydrolysis) were generally not considered important for BaP. Photolysis is an important fate process for these PAHs, thus the absorption spectrum was included as input (Table 5-28).

A loading rate of 0.1 kg/hour was specified for the BaP calculations. However, in some environments this loading would cause a concentration of benzo[a]pyrene in the input stream that was greater than the maximum water solubility of the compound. When this type of overloading occurs, the EXAMS model automatically decreases the loading to a physically allowable level. Once the concentration of BaP is below the maximum solubility level, the maximum concentrations and accumulations are proportional to the loading, and the self-purification times and disposal rates remain unaffected. As shown in Table 5-29, the loading rates for the relatively static aquatic systems (ponds and lakes) were reduced by the model to 3.6×10^{-2} kg/day or lower. The load to the rivers was allowed to remain at 2.4 kg/day due to the fact that rapid dilution prevented BaP concentrations above solubility levels. The materials balance (Section 5.1) suggests that a maximum loading would be approximately $3x10^{-2}$ kg/day. Therefore, the concentration estimates for the pond and lakes are reasonable, but the actual river concentrations would be expected to be lower than the EXAMS estimates, shown in the following tables, by a factor of 80.

Table 5-29 summarizes the BaP concentrations predicted for the simulated environments under steady-state conditions. Concentrations in water are generally 10^2-10^4 times lower than concentrations in sediment and biota. The steady-state accumulation of BaP is highest in those environments with a higher biomass, i.e., coastal river, eutrophic lake and pond. The distribution and fate of BaP in the aquatic systems are presented in Table 5-30. In the oligotrophic lake, about 81% of the BaP load remains in the sediment, while more than 94% of the load in the rest of the environments resides in the sediment.

TABLE 5-27. INPUT PARAMETERS FOR EXAMS MODELING OF THE FATE OF BENZO[a] PYRENE IN GENERALIZED AQUATIC SYSTEMS

Explanation	Value
Molecular weight	252.0
Ratio volitilization: re-aeration rate	0.3760
Henry's Law constant	4.8×10^{-7}
Quantum yield in water(at 313 nm)	8.9×10^{-4}
Absorption spectra	See Table 5-28
Aqueous solubility(mg/l)	3.8×10^{-3}
Partition coefficient for biomass:H ₂ 0	1.89×10^5
Octanol:H ₂ O partition coefficient	1.1×10^{6}
Second-order biolysis rate constant: in sediment and in the water column	3.0×10^{-12}
Increase in biolysis rate constant for 10°C increase in temperature	2.0

Source: SRI (1980).

TABLE 5-28. MOLAR ABSORBTIVITY FOR BENZO[a]PYRENE AS A FUNCTION OF WAVELENGTHS

Interval	Center of Interval (nm)	Molar Absorbtivity (lcm/mole/
1	297.5	46,600
2	300.0	27,700
3	302.5	13,900
4	305.0	6,670
5	307.5	4,840
6	310.0	3,970
7	312.5	3,870
8	315.0	3,650
9	317.0	3,730
10	320.0	3,570
11	323.1	3,650
12	330.0	5,400
13	340.0	8,330
14	350.0	12,300
15	360.0	18,100
16	370.0	19,680
17	380.0	21,910
18	390.0	15,160
19	400.0	2,100

Source: Smith et al. (1978).

TABLE 5-29. STEADY-STATE CONCENTRATIONS OF BENZO[a]PYRENE IN VARIOUS GENERALIZED AQUATIC SYSTEMS RESULTING FROM CONTINUOUS DISCHARGE^a

			Max	kimum Concer	trations			
System	Loading (kg/hr)	Water Dissolved (mg/1)	Water Total (mg/l)	Maximum in Sediment Deposits (mg/kg)	Plankton (µg/g)	Benthos (µg/g)	Total Steady-State Accumulation (kg)	Total Daily Load (kg/day)
Pond	6.2×10^{-5}	2.2×10^{-4}	1.2×10^{-3}	16	43	43	11	1.5x10 ⁻³
Eutrophic Lake	1.5×10^{-3}	1.6×10^{-4}	1.7x10 ⁻³	4.3	30	30	59	3.6×10^{-2}
Oligotrophic Lake	1.5x10 ⁻³	3.2×10^{-5}	3.4×10^{-4}	0.11	6.1	0.3	0.047	3.5×10^{-2}
River	0.1	4.1×10^{-5}	1.5×10^{-4}	0.33	7.8	1.4	6.2	2.4
Turbid River	0.1	1.3x10 ⁻⁵	1.2x10 ⁻⁴	0.09	2.4	1.2	1.9	2.4
Coastal River	0.1	7.4×10 ⁻⁴	1.9x10 ⁻³	19	140	37	290	2.4

^aAll data simulated by EXAMS (U.S. EPA 1980b) model (see text for further information).

TABLE 5-30. THE FATE OF BENZO[a]PYRENE IN VARIOUS GENERALIZED AQUATIC SYSTEMS a

	Percent Distribution Percent Lost by Various Processes						
System	Residing in Water at Steady-State	Residing in Sediment at Steady-State	Transformed by Chemical Process	Transformed by Biological Process	Volatilized	Lost by Other Processes	Time for System Self- Purification ^C
Pond	0.24	99.76	68.79	0	0.16	31.03	69 years
Eutrophic Lake	1.85	98.15	95.10	0.05	0.21	4.64	25 years
Oligotrophic La	nke 18.75	81.25	99.97	0	0.02	0.01	393 days
River	2.17	97.83	0.04	0	0	99.96	95 days
Turbid River	5.63	94.37	0	0	0	100	89 days
Coastal River	0.56	99.44	2.89	0	0.02	97.08	376 days

^aAll data simulated by the EXAMS (U.S. EPA 1980b) model. See text for further information.

 $^{^{\}mathrm{b}}$ Including loss through physical transport beyond system boundaries.

^CEstimate for removal of ca. 97% of the toxicant accumulated in system. Estimated from the results of the half-lives for the toxicant in bottom sediment and water columns, with overall cleansing time weighted according to the pollutant's initial distribution.

The ultimate fate of BaP in these systems is controlled primarily by chemical processes in the static systems, and physical transport beyond the system boundaries for the river systems. Volatilization and biological degradation are negligible processes for BaP.

The EXAMS estimates for the persistence of BaP upon cessation of loading in the aquatic environments are given in Table 5-31. The fastest system to purify itself of the BaP load is the turbid river: 8.93% is lost within one day: Table 5-30 indicates a self-purification time of 89 days due to physical transport. The other river systems show similar purification characteristics, although somewhat slower. Of the pond and lake systems, the oligotrophic lake exhibits the lowest persistence of BaP. Within 24 days 30% of the accumulated BaP will be lost. Chemical degradation processes control the fate of BaP in the oligotrophic lake, probably due to the high rate of photolysis in the clear waters and the fact that almost 20% of the BaP is residing in the water column and available for absorption of sunlight. The pond and eutrophic lake require much longer time periods for removal of BaP, and have self-purification times of 69 years and 25 years. respectively. Chemical transformation appears to account for the ultimate fate of BaP in these systems. However, the waters of the pond and eutrophic lake are not clear and photolysis will be slower than in the oligotrophic lake. Furthermore, more than 98% of the BaP resides in the sediment of these systems, where light penetration is extremely low.

Comparison of Mackay and EXAMS Models

The Mackay model used is simply a partitioning model, whereas EXAMS utilizes kinetic data to predict the fate of the compound after partitioning. Therefore, the models are not directly comparable. The EXAMS pond environment was taken to be the best system to compare with the Mackay model, since there is very little transport across system boundaries. The load to the Mackay model was assumed to be equal to the total pond accumulation predicted by EXAMS, 11 kg.

Table 5-32 summarizes the concentration and distribution data predicted by the two models. The percentages of distribution predicted by the two models agree quite well; the estimated concentrations are also in agreement within an order of magnitude. The most important conclusion from these data is that there will be very little BaP in the dissolved state, and most of the aquatic load will be removed to the sediment.

TABLE 5-31. THE PERSISTENCE OF BENZO[a]PYRENE IN VARIOUS GENERALIZED AQUATIC SYSTEMS AFTER CESSATION OF LOADING^a

System	Time Period (days)	% Lost from Water	% Lost from Sediment	% Lost from Total System
Pond	1460	19.02	18.17	18.18
Eutrophic Lake	2190	71.93	56.26	56.55
Oligotrophic Lake	24	92.63	16.01	30.36
River	3.5	94.57	11.74	13.54
Turbid River	1	98.21	3.6	8.93
Coastal River	168	87.52	78.67	•78.27

All data simulated by the EXAMS (U.S. EPA 1980b) model. See text for further information.

TABLE 5-32. COMPARISON OF RESULTS FROM MACKAY'S EQUILIBRIUM MODEL AND EXAMS FOR BENZO[a]PYRENE IN A POND SYSTEM

EXAMS Results (Pond: 1.5x10 ll kg steady	0 ⁻³ kg/day loading, -state accumulation)	Mackay Results (11 kg in system)	
	Maximum Concentrations		Concentrations
Water (Total)	0.2 μg/l	Water (dissolved)	0.13 µg/1
Water Biota	43 mg/kg	Suspended Solids	1.6 mg/kg
Sediment Biota	43 mg/kg	Water Biota	28.4 mg/kg
Sediment	16 mg/kg	Sediment Biota	28.4 mg/kg
		Sediment	1.6 mg/kg
	Steady-State Accumulation	Percent of Chemica	l per Compartment
% in Water	0.24	% in water ^a	0.1
% in Sediment	99.76	% in Sediment	99.9

^aIncludes BaP dissolved in water, adsorbed on suspended solids, and in aquatic biota compartment.

5.2.4 Monitoring Data

PAHs, especially those in the BaP group, have been extensively monitored in environmental media. This section contains a discussion of the monitoring data contained in STORET, as well as a brief review of other data from the literature. In general, the most information is available for BaP, due to the emphasis on monitoring for this compound because of its carcinogenic nature.

5.2.4.1 STORET

Ambient and Effluent Water

STORET monitoring data for the benzo[a]pyrene group reflect sampling activities in twenty-nine states and the U.S. territory of Puerto Rico.

As Table 5-33 indicates, observations in STORET indicate low concentrations for the BaP group PAHs. Generally, levels are less than the most frequently used detection limit of 10 $\mu g/l$. For three of the compounds, benzo[g,h,i]perylene, dibenz[a,h]anthracene, and indeno-[1,2,3-c,d]pyrene, the detection limits most used during analysis were higher than 10 $\mu g/l$; however, levels in ambient waters do not exceed 100 $\mu g/l$. Table 5-34 presents the maximum detection limits used in monitoring and the ranges of unremarked observations 2 for each pollutant.

Table 5-33 shows the distribution of effluent concentrations, remarked and unremarked, for the benzo[a]pyrene group. Roughly 40% of the observations indicate detection limits equal to or less than 1 μ g/1, and 50% with detection limits between 1.1 μ g/1 and 10 μ g/1. The number of unremarked observations does not exceed eight for any one pollutant. Positive values range from less than 1 μ g/1 to over 1000 μ g/1.

Sediment and Soil

Data summaries are presented in Tables 5-34 and 5-35 for concentrations in sediment for the benzo[a]pyrene group contained within the STORET data base. For each pollutant, roughly 90% of the observations are remarked. The distribution indicates that, of the remarked observations, detection limits equal to or less than 1 $\mu g/kg$ (dry weight) are used about 30% of the time, detection limits between 100.1 and 1000 $\mu g/kg$ are used 10% of the time, and detection limits over 1000 $\mu g/kg$ are used 60% of the time. No more than a dozen unremarked observations are recorded for each pollutant, and these positive values are generally less than 1000 $\mu g/kg$.

 $^{^2}$ Remarked observations are generally undetected values, unremarked observations are positive values.

TABLE 5-33. DISTRIBUTION OF OBSERVED AMBIENT AND EFFLUENT CONCENTRATIONS OF THE BENZO(a)PYRENE GROUP PAHS

Number of Ambient Observations

				Mumber Of	vanorem	nt observations						
	Remarked Data					Unremarked Data						
	Total #			Conc. (pg	/1) .		Total #			Conc. (µ)		
Compound	Obs.	<u>≤1</u>	1.1 - 10	10.1-100	100.1-1000	>1000	Obs.	<u>≤1</u>	1.1-10	10.1-100	100.1-1000	>1000
							J					•
Benzo[a] pyrene	411	41	281	80	9		0	1				
Acenaphth ylene	423	46	326	38	13		6	6				
Benz[a]anthracene	347	41	268	35	3		5	1	2		2	
Benzo[b] fluoranthene	236	41	154	41			0	1				
Benzo[k] fluoranthene	354	39	194	113	4	4	5				3	2
Benzo[g,h,i]perylene	407	37	128	230	8	4	0	1				
Chrysene	246	47	184	15			i	1				
Dibenz[a,h]anthracene	416	50	108	237	13	8	Ô	l				
Indeno[1,2,3-c,d]pyren	e 404	45	121	222	12	4	lo	ļ				
		1						•				

					Number of 1	Effluen	Observ	atio	ons ^b			
	Remarked Data					Unremarked Data						
	Total i	1		Conc. (µ)	g/1)		Total #	1		Conc. (µg/1)		
	Obs.	<u>≤1</u>	1.1-10	10.1-100	100.1-1000	>1000	Obs.	<u>≤1</u>	1.1-10	10.1-100	100.1-1000	>1000
Benzo[a]pyrene	667	290	335	42			4	2	1			1
Acenaphthylene	575	196	362	17			8	4	4			
Benz[a]anthracene	559	190	3 52	17			4	[]	1			3
Benzo[b]fluoranthene	512	215	287	10			2	1	1			1
Benzo[k]fluoranthene	659	325	320	14			2	2	-			
Benzolg, h, ilperylene	677	296	305	76			1 3	2				1
Chrysene	508	188	315	5			7	2	1	1		3
Dibenz[a,h]anthracene	564	180	305	73	6		,	1~		i	1	J
ludeno [1,2,3-c,d]pyrer	ne 666	282	303	81			6	5		•	1	

aData as of December 1, 1980, except data for Dibenz[a,h]anthracene, which are as of November 20, 1980.

Source: U.S. EPA (1980c).

b_{Data} as of April 9, 1980.

TABLE 5-34. NUMBER, DETECTION LIMITS, AND RANGES OF OBSERVED CONCENTRATIONS IN AMBIENT WATER AND SEDIMENT FOR THE BENZO[a]PYRENE CROUP PAILS--STORET, 1980^a

Aubiona II a (12)	Benzo[a]- pyrene	Acenaph- thylene	Benz[a]- anthra- cene	Benzo[b fluoran thene]- Benzo[k - fluoran thene]- Benzo- - [g,h,i]- perylene		Dibenz [a,h] an- thracene	
Ambient Water (µg/1)									
Total No. Observations		429	352	236	359	407	247	416	101
Remarked	0 411	6	5	0	5	0	1	0	404
TO HELL KEY	411	423	347	236	354	407	246	416	0 404
Maximum Detection Limit	800	640	400	25	1000	1600	25	1600	1600
Range of Unremarked Observations		0.01-0.12	1-400		320-1500		0.02		
Sediment (µg/kgdry we	ight)								
Total No. Observations Unremarked Remarked	125 11 114	125 12 113	116 12 104	109 13 96	121 12 109	123 5 118	102 11 91	130 2 128	117 5
Maximum Detection Limit	10000	10000	10000	.10000	10000	10000	10000	10000	112 10000
Range of Unremarked Observations	0.02-1400	0.002-93	6.2-340	3.4-310	0.9-1300	4.3-40.9	0.06-120	15.7~2600]	1-35.8

aData as of November 20, 1980, except for Dibenz[a,h]anthracene, which are as of December 1, 1980. Source: U.S. EPA (1980c).

TABLE 5-35. DISTRIBUTION OF OBSERVED SEDIMENT AND TISSUE CONCENTRATIONS OF THE BENZO[a]PYRENE GROUP PAHS.

		1			Number of	Sedime	it Obser	vat	ions				
		 		Remarked Da	ta			Unremarked Data					
	Total #	1	Conc.	· ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' '			Total#			g/kg-dry v			
	Obs.	<u> </u>	1.1-10	10.1-100	100.1-1000	>1000	Obs.	<u>≤1</u>	1.1-10	10.1-100	100.1-1000	>1000	
							ĺ						
Benzo[a]pyrene	114	30			19	65	11	1 2	1	5	2	1	
Acenaphthalene	113	28		2	18	65	12	1 1	6	5	2-	1	
Benz[a]anthracene	104	27			16	61	12	3	9		0		
Benzo[b]fluoranthene	96	28		1	17	50	13	٦	2)	2		
Benzo[k]fluoranthene	109	28		-	21	60		١.	2	2	9		
Benzo[g,h,i]perylene	118	35		1			12	1	2	4	4	1	
Chrysene		28		1	12	70	5		3	2			
	91				13	50	11	1	2	6	2		
Dibenz[a,h]anthracene	128	36			18	74	2			1		2	
Indeno[1,2,3-c,d]pyrene	112	29		1	16	66	5	1	2	$\frac{1}{2}$		1	

Numbel	of Tis	s <u>ue Observ</u> a	ations			
Remarked Data		Unremarked Data				
		1 1 .	Conc. (mg/kg-wet wt.) 1.1-10 10.1-100 100.1-1000 >1000			
62	1000		$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			
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62		1 1				
62		0				
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55 9						
51 9		- 1				
	1	0				
	Į	ļ l				
6 6 6 6 5 5	Remarked Data Conc. (mg/kg-wet wt.) 1-10 10.1-100 100.1-1000 2 0 2 2 2 2 2 6 8 8 8 5 9	Remarked Data Conc. (mg/kg-wet wt.) 1-10 10.1-100 100.1-1000 >1000 2 0 2 2 2 2 2 6 8 8 8 7 9	Conc. (mg/kg-wet wt.) 1-10 10.1-100 100.1-1000 >1000 Obs. \$\frac{1}{2}\$ 0 1 2 2 2 2 6 8 8 8 7 9			

Source: U.S. EPA (1980c).

Fish Tissue

As shown in Table 5-35, most concentrations for the benzo[a]pyrene group in fish tissue are remarked. The exceptions are observations for acenaphthylene, with one unremarked observation no higher than 1 mg/kg (wet weight), and chrysene, with two unremarked observations no higher than 1 mg/kg. The distribution indicates that, in general, roughly 15% of the time fish tissue samples are analyzed with detection limits not exceeding 1 mg/kg, and 85% of the time the detection limits are between 1.1 mg/kg and 10 mg/kg-wet wt.

5.2.4.2 Data From Other Sources

Ambient and Effluent Water

White and Vanderslice (1980) have thoroughly reviewed literature reports of concentrations of PAHs in ambient waters. The results of their review are presented as frequency distributions of reported concentrations in Table 5-36. The data show that levels are almost always less than 1 μ g/l, and commonly less than 0.01 μ g/l, suggesting that actual levels are probably much less than the 10 μ g/l detection limit generally reported in the STORET data base.

A 1978 study by Basu and Saxena (1978) reports concentrations of BaP and of total PAHs in river waters that serve as municipal drinking water supplies. Their results are summarized in Table 5-37. The higher levels found in Monongahela probably result from coking operations in the area.

Andelman and Snodgrass (1974) have reported BaP concentrations of 2-300 $\mu\text{g}/1$ in aqueous effluents from industrial operations such as shale oil recovery, oil refining coke byproduct recovery, and acetylene manufacture. The fact that the upper end of this range exceeds the BaP water solubility suggests that total (dissolved plus suspended) PAHs were being analyzed.

Davies et al. (1976) reported that levels of BaP group PAHs in the spent scrubber water of a British municipal refuse incinerator were on the order of 0.03-0.64 μ g/l. These direct aqueous discharges represented only a small (<1%) percentage of the total estimated PAH emissions from this facility, compared with stack gas (approximately 10%) and ash (approximately 90%) emissions of PAHs.

A systematic study of sources of priority pollutants to POTWs in Cincinnati, St. Louis, Atlanta, and Hartford (Levins <u>et al</u>. 1980) reported no detectable levels of BaP or other PAHs in this group in raw wastewater from residential, commercial, or industrial urban areas, at $4-10~\mu g/1$ detection limits.

Drinking Water

Basu and Saxena (1978) investigated levels of BaP group PAHs in several United States municipal drinking water supplies. The results

TABLE 5-36. CONCENTRATIONS OF BENZO[a]PYRENE GROUP PAHS IN AMBIENT WATER

Sample/PAH	Total No. of <u>Observations</u>	No. Observations in Indicated Concentrations (pg/1) Range								
		0.0001-0.001	0.001-0.01	0.01-0.1	0.1-1	1 <u>-</u> 10	> 10			
River Water										
Benzo[a]pyrene	17	2	5	4	5		1			
Benz[a]anthracene	5			4	1		1			
Benzo[b]fluoranthene	5		1	3	1					
Benzo[k]fluoranthene	12	1	2	7	2					
Benzo[g,h,i]perylene	13		3	6	4					
Chrysene	1		J	1	4					
Indeno(1,2,3-c,d]pyrene	12		2	6	4					
Lake Water			-	Ū	4					
Benzo[a]pyrene	4	1	ı		ı	•				
Benzo[b]fluoranthene	1	1	-		ı	1.				
Benzo[k]fluoranthene	1		1							
Benzo[g,h,i]perylene	1		1							
Groundwater		3	1		2					
Benzo[a]pyrene	6	1	1		2					
Benz[a]anthracene	2	2	2							
Benzo[b]fluoranthene	4	2	2							
Benzo[g,h,i]perylene	4		_							
Precipitation										
Benzo[a]pyrene	2			2						
Benz[a]anthracene	2			1	,					
Benzo[b]fluoranthene	2			1) 1					
Benzo[k]fluoranthene	2			2	•					
Benzo[g,h,i]perylene	2			1	J					

Source: White and Vanderslice (1980).

TABLE 5-37. LEVELS OF PAHS IN THE MONONGAHELA, OHIO, AND DELAWARE RIVERS

	Concentration (ng					
River	<u>BaP</u>	Total PAH				
Monongahela, PA	42-77	600-660				
Ohio, WV	5.6	58				
Delaware, PA	4.1	350				

Source: Basu and Saxena (1978).

reported for tap water in selected locations in New York, Pennsylvania, West Virginia, and Louisiana are given in Table 5-38. The highest value reported in this survey was $0.004~\mu g/l$ for benzo[g,h,i]perylene. However, most values were less than $0.001~\mu g/l$.

Levins <u>et al</u>. (1980) reported no detectable levels of the BaP group PAHs in tap water samples from Cincinnati, St. Louis, Atlanta, and Hartford (detection limit of $4\text{--}10~\mu\text{g}/1$).

Sediment and Soil

Heit (1979) has reported BaP concentrations of 6-305 $\mu g/g$ (dry weight) in sediment samples collected at depths ranging from 0-55 cm from a lake near Los Angeles, CA. The largest concentration corresponds to a sample taken at a depth of 27-29 cm; this value was almost a factor of 10 above the concentrations at the other depths. Samples of other western lake sediments from rural areas showed BaP levels generally below the 2 $\mu g/g$ detection limit in this study.

Giger and Blumer (1974) and Hites et al. (1977) have documented the BaP levels in marine sediments in the vicinity of the 1969 oil spill near West Falmouth, MA. BaP concentrations of 370 $\mu g/kg$ and 340 $\mu g/kg$ dry weight of sediment were observed in 1969 and 1970, respectively. Sediment core samples showed apparent historical levels of 380 $\mu g/kg$ (in 1900) and 26 $\mu g/kg$ (in 1850). By contrast, a parallel examination of Charles River, MA, sediment by Hites et al. (1977) showed a BaP level of 8000 $\mu g/kg$ dry weight of sediment.

A research group at Research Triangle Institute (White and Vanderslice 1980) has reviewed reported data on PAH levels in soils. The results of that review for the BaP group PAHs are summarized in Table 5-39.

Fish Tissue

Shellfish tissue taken from estuarine waters indicated a higher level of BaP than those taken from marine environments (Pancirov and Brown 1977, Mix et al. 1976). Table 5-40 summarizes readily available data on the BaP group PAHs in edible marine organisms from a number of U.S. locations.

Air

Benzo[a]pyrene has been monitored in air in urban and industrial areas. Table 5-41 summarizes concentration ranges for BaP and related PAHs in U.S. urban air as cited in the U.S. EPA Criteria Document (U.S. EPA 1980a). Table 5-42 presents some earlier data, also from the Criteria Document, that illustrates the variability of PAH levels in different cities. Table 5-43 presents the frequency distribution of observations of BaP group PAHs in urban air, based upon the compilation of White and Vanderslice (1980).

TABLE 5-38. LEVELS OF BENZO[a]PYRENE GROUP PAHS IN DRINKING WATER

Location of Water Supply Syste		Sampling Date	Concentra Benzo[a] pyrene	tion data for Benzo[k] fluoranthene	Treated Drinki Benzo[g,h,i] perylene	ng Water ^a (ng/l) Indeno[1,2,3-c,d] pyrene
Syracuse, NY	Lake Skaneatelas (uncontaminated)	12/16/76	0.3	0.4	0.4	
Buffalo, NY	Lake Erie (contaminated with industrial discharge)	12/26/76	0.2		0.7	
Pittsburgh, PA	Monongahela River (contaminated with coke oven effluent)	1/19/77	0.4	0.2	0.7	1.2
Huntington, WV	Ohio River (downstream from coke oven plants)	1/20/77	0.5	0.2	2.5	1.2
Endicott, NY	Groundwater (uncontaminated)	2/22/77	0.2		2.9	0.7
lammondsport, NY	Keuka Lake (contaminated with agricultural waste)	2/28/77	0.3	0.1	1.9	0.9
'hiladelphia, PA	Delaware River (contaminated with municipal waste)	3/5/77	0.3		4.0	1.7
nidentified	Uncontaminated upland water	3/17/77	0.5	0.7	1.8	2.2
ake George, NY	Lake George (contamination from recreational sources)	3/26/77	0.3	0.1	2.6	0.9
ew Orleans, LA	Mississippi River(contaminated with industrial discharges)	5/1/77	1.6	0.6		·

^aData are for single analysis of one water sample taken from each of ten finished water supplies.

Source: Basu and Saxena (1978).

TABLE 5-39. FREQUENCY OF OBSERVATIONS OF BENZO[a]PYRENE GROUP PAHS IN SOIL AND SEDIMENT

Sample/PAH	Total No. of Observations	1	40. Observat	ions in	Indicated	Concentratio	on (µg/kg) Range	<u>.</u>
Rural Soil		0.01-0.1	0.1-1.0	1-10	10-100	100-1000	1000-10,000	10,000-100,000
Benzo[a]pyrene	52	3	3	19	23	4		
Benz[a]anthracene	3			3				
Benzo[b]fluoranthene	3				3			
Benzo[k]fluoranthene	3			2	1			
Benzo[g,h,i]perylene	6			3	3			
Indeno[1,2,3-c,d]pyrene	3			2	1			
Urban Soil								
Benzo[a]pyrene	23			1	7	7	7	1
Benz[a]anthracene	2					1		1
Benzo[k]fluoranthene	2]	1	
Benzo[g,h,i]perylene	2					1		1
Chrysene	2						1	1
Marine Sediments								
Benzo[a]pyrene	17				3	4	6	
Benz[a]anthracene	2			4	1 .	1		
Benzo[g,h,i]perylene	2				1	1		
Chrysene	2				1	Ł		
Soil Near Industrial Sources								
Benzo[a]pyrene	13				4	3	2	4

Source: White and Vanderslice (1980).

TABLE 5-40. REPORTED LEVELS OF BENZO[a]PYRENE GROUP PAHS
IN EDIBLE MARINE ORGANISMS

Concentration (ug/kg wet weight) Marine Benz[a]-Other Location of Sample Tissue anthracene BaP PAHsa Long Island Sound Oyster 8 2 <2-15 Chincoteague, VA Black Point Oyster 0.1 0.2 <0.5-0.7 Little Toms Cove Clam 0.3 0.3 <0.1-0.9 Darien, Conn. Scotts Cove Clam 1 <1 1-3 Fish Market, Linden, NJ Clam <1 <1 <0.5-<3 Chesapeake Bay Crab <1.5 0.5 <0.5-<1.2 Raritan Bay Crab 2 3 1-2 Menhaden 1.5 < 0.3 <0.3-1 Atlantic Ocean Long Branch, NJ Flounder 2 <1 <1-<2 S. of Long Island Flounder <1 1 1 Falmouth, MA Little Sippewisset Mussel <0.2 <0.5 <0.2-0.3 Wild Harbor Mussel <0.6 <0.5 <0.5-1.2 <0.3-<1 Palacios, Texas Shrimp <0.2 <1 <0.5-<5 Atlantic Ocean 25 mi. off Toms River, NJ Codfish <2 <1 <0.5-<1.5

Source: Pancirov and Brown 1977.

Source: Pancirov and Brown (1977)

a Includes chrysene, benzo[b]fluoranthene, benzo[e]pyrene, perylene, and benzo[g,h,i]perylene.

TABLE 5-41. REPORTED LEVELS OF BENZO[a]PYRENE GROUP PAHS IN AIR OF U.S. CITIES

Compound	Concentration Range (ng/m ³) ^a	Reference
Benz[a]anthracene	0.18-4.6	Fox and Staley (1976), Gordon (1976)
Benzo[b]fluoranthene	0.1-1.6	Gordon and Bryan (1973)
Benzo[k]fluoranthene	0.18-5.2	Fox and Staley (1976)
Benzo[a]pyrene	0.13-3.2	Colucci and Begeman (1971)
<pre>Indeno[1,2,3-c,d]pyrene</pre>	0.03-1.34	Gordon (1976), Gordon and Bryan (1973
Chrysene	0.2-6.4	. Gordon and Bryan (1973)
Benzo[g,h,i]perylene	0.2-9.2	Gordon and Bryan (1973)

The concentration data for air can be converted to ppt (parts per trillion) by multiplying the value in ng/m³ by the following factors (F=24.47/M.W.): benz[a]anthracene and chrysene, F=0.107; benzo[a]pyrene and benzofluoranthenes, F=0.097; indeno[1,2,3-c,d]pyrene and benzo[g,h,i]-perylene, F=0.089.

TABLE 5-42. CONCENTRATIONS OF BENZO[a]PYRENE GROUP PAHS IN THE AIR OF SELECTED U.S. CITIES, AVERAGE OF SUMMER AND WINTER VALUES

City	Benzo[a]pyrene	Concentration (ng/mBenzo[k]fluoranthene	Benzo[g,h,i]perylene
Atlanta, GA	4.5	3.7	7.0
Birmingham, AL	15.7	8.8	13.2
Detroit, MI	18.5	12.5	21.3
Los Angeles, CA	2.9	3.1	10.2
Nashville, TN	13.2	8.0	10.2
New Orleans, LA	3.1	2.9	6.0
San Francisco, CA	1.3	1.0	5.1
Range	1.3-18.5	1.0-12.5	5.1-21.3
Mean	8.5	5.7	10.4

^aConcentration data for air samples can be converted to ppt (parts per trillion) by multiplying the value in ng/m^3 by the following factors (F=24.47/M.W.): benzo[a]pyrene and benzo[k]fluoranthene. F=0.097; benzo[g,h,i]perylene, F=0.089.

Source: Sawicki et al. (1962).

TABLE 5-43. FREQUENCY OF AMBIENT CONCENTRATIONS OF BENZO[a]PYRENE GROUP PAHS IN URBAN AIR

РАН	Total No. of Observations	No. Ob	servations i	n <u>I</u> ndicated	Concentrati	ion (ng/m ³) R	ange
		0.01-0.1	0.1-1	<u>1-10</u>	10-100	100-1000	> 1000
Benzo[a]pyrene	116	1	6	45	54	9	1
Benz[a]anthracene	24	4	5	8	5	2	
Benzo[b]fluoranthene	8		4	4			
Benzo[k]fluoranthene	29	3	7	13	6		
Benzo[g,h,i]perylene	54	1	6	21	24	2	
Chrysene	12	1	2	6	2	1	
Indeno[1,2,3-c,d]pyrene	2		1	1			

Source: White and Vanderslice (1980).

Air samples collected in areas surrounding combustion sources show the highest levels of BaP in air. Table 5-44 presents some data showing that oil refineries, for example, can be associated with air levels of BaP 30 times those near airports. Katz and Chan (1980) note that an earlier U.S. EPA study found BaP levels in urban air to be 1.42 to 3.34 times higher in cities with coke oven facilities than in cities without coke ovens. Table 5-42 shows relatively high ambient levels of BaP in cities such as Detroit and Birmingham, compared with San Francisco, for example. These differences are due probably in considerable part to the different degrees of industrialization, although meteorologic and climatologic factors are also important.

Rural areas have been shown to have detectable levels of PAHs in ambient air, though the levels are well below any levels measured in heavily populated or industrial areas (U.S. EPA 1980a). Table 5-45 presents a frequency distribution for reported concentrations in rural air compiled by White and Vanderslice (1980).

Moschandreas <u>et al</u>. (1980) examined BaP levels in a few locations, indoors and outdoors, on woodburning and non-woodburning days. In the Boston metropolitan area, they found mean levels for various residences of $0.4-1.1~\text{ng/m}^3$ indoors, and $0.4-0.9~\text{ng/m}^3$ outdoors. On woodburning days in one residence, the mean level indoors was $4.7~\text{ng/m}^3$, and the mean level outdoors was $1.3~\text{ng/m}^3$. These levels suggest that there is little difference between indoor and outdoor concentrations of BaP except on days on which wood is being burned.

In order to place the ambient air data in perspective, Table 5-46 presents some data on PAHs in mainstream cigarette smoke (Severson et al. 1976). Note that these data are in $\mu g/m^3 \text{ vs. ng/m}^3$ for ambient air concentrations and thus generally at least an order of magnitude higher.

5.2.5 Summary - Ultimate Fate and Distribution

Benzo[a]pyrene and the other PAHs in this group are released to the environment primarily as products of combustion. Virtually all of the BaP in the atmosphere is adsorbed onto airborne particulates. Even though photolysis of BaP in the atmosphere is expected to be rapid, wet and dry deposition of these PAHs are estimated to be the major input pathways to the aquatic environment. On the basis of transport model calculations, it was estimated that ~44 kkg/yr BaP may be deposited on the surface of the U.S.; only a fraction of that will be introduced directly or indirectly to aquatic systems.

The relative composition of the PAH mixtures released from high-temperature combustion sources indicates that production of the unsubstituted, parent compound is favored. The fact that, within a PAH series, unsubstituted PAHs are the most abundant homologs in sediments from industrial areas (despite their higher solubility) further supports the conclusion that combustion is the major source of PAHs in the environment.

TABLE 5-44. RELATIONSHIP BETWEEN CONCENTRATION OF BENZO[a]PYRENE IN AIR AND DISTANCE FROM EMISSION SOURCE

Concentration of BaP (μ g BaP/kg of particulate material)

Location	Distance from Source (m)						
	At Source	50	51-250	251-500	501-1500		
Airport ^a - USSR	400	64	46	17	1.3		
Oil refinery - USSR	12,000			1,200	120		
Highway ^b (in town) (rural)	176 120	100	6	21 15	5		

a Maximum values

Source: White and Vanderslice (1980).

bTotal POM µg/kg

5-36

TABLE 5-45. FREQUENCY OF AMBIENT CONCENTRATIONS OF BENZO[a]PYRENE GROUP PAHS IN RURAL AIR

РАН	Observations	Number of Observations in Indicated Concentration(ng/m Range						
		0.001-0.01	0.01-0.1	0.1-1	1-10	10-100		
Dibenz[a,h]anthracene	2		1	1				
Benzo[a]pyrene	16	2	2	2	6	6		
Benzo[g,h,i]perylene	10				1	1		

Source: White and Vanderslice (1980).

TABLE 5-46. CALCULATED CONCENTRATIONS OF BENZO[a]PYRENE AND RELATED PAHS IN MAINSTREAM CIGARETTE SMOKE

Concentration $(\mu g/m^3)$ Non-Filter Filter Cigarette <u>PAH</u>a Cigarette (10 mg) Cigarette Benzo[a]pyrene plus 85-86 28-32 16 Benzo[e]pyrene Benzo[a]anthracene plus 310-680 96-210 73 Chrysene plus Triphenylene

Source: Calculated from data of Severson et al. (1976).

Analysis method used did not give complete resolution of the PAHs; data are for partially resolved mixtures of PAHs.

The Mackay equilibrium partitioning model was run for BaP, and the calculations indicate that 99.9% of the environmental BaP will reside in the sediment compartment; very small amounts will be found in the air (vapor phase BaP), water, or biota. The U.S. EPA EXAMS model was used to predict the fate and distribution of BaP in six generalized aquatic systems. The results indicate that in all of the systems modelled, maximum BaP concentrations in sediment can be expected to be higher than total water concentrations by factors up to 10^4 . Aqueous BaP concentrations were estimated to range from 0.1 μ g/l to 1.9 μ g/l; sediment concentrations predicted by EXAMS range from 90 μ g/kg to 1.9 x 10^4 μ g/kg. The EXAMS data also indicate that more than 90% of the BaP will reside in the sediment for all aquatic systems examined, with the exception of the oligotrophic lake (81%).

The most significant fate processes for BaP in aquatic systems include adsorption (with subsequent transport to the sediment), and chemical degradation (photolysis). Volatilization and biodegradation, with half lives on the order of a number of days to months, are expected to be slow processes for BaP and the other PAHs in this group. The EXAMS calculations indicate that the relative importance of the various fate processes is determined by the actual conditions characteristic of the environmental systems. In the static pond and lake systems, chemical degradation will be the dominant fate process. However, in the more dynamic river systems, physical transport of BaP downstream is largely responsible for determining the ultimate fate. The persistence of BaP in aquatic systems is reflected in the self-purification times predicted by EXAMS. The times estimated for removal of 97% of the BaP accumulated in the turbid river system is 89 days, whereas the self purification time predicted for the static pond system is as high as 69 years.

Figure 5-9 gives a representation of the major inputs of BaP to the aquatic environment, as well as the dominant fate and transport pathways.

BaP has been extensively monitored in environmental media; there are fewer data for the other PAHs in this group. Most of the monitoring data in the STORET data base for the PAHs in the BaP group are remarked, i.e., reported to be below the detection limits. The PAH concentrations actually reported for sediment samples range from 0.002 $\mu g/kg$ to 2600 $\mu g/kg$. Ambient water concentrations recorded in STORET range from 0.01 $\mu g/l$ to 1500 $\mu g/l$. However, since sediment and aqueous samples were not taken from the same locations, very few conclusions can be based entirely on the limited data in STORET.

Data from other sources indicate that BaP concentrations in ambient waters are generally less than 1 $\mu g/1$; concentrations of BaP in industrial effluents were reported to be as high as 300 $\mu g/1$. Concentrations of BaP in drinking water were reported to range from 0.2 ng/1 to 1.6 ng/1; the highest BaP concentration was in New Orleans, LA. In this survey, the highest concentration for the PAHs in the BaP group was 4.0 ng/1, reported

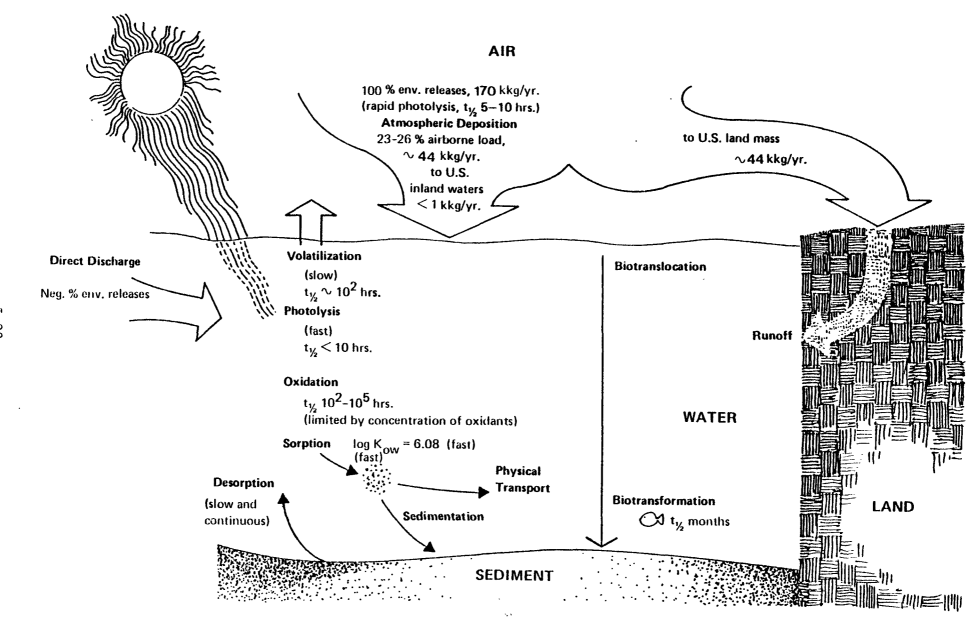


FIGURE 5-9 SOURCES AND FATE OF BENZO[a] PYRENE IN THE AQUATIC ENVIRONMENTS

for benzo[g,h,i]perylene in a drinking water sample taken from Philadelphia, PA. Benzo[a]pyrene has been reported at concentrations up to 300,000 µg/kg (dry weight) in lake sediment samples. In contrast, a Charles River, MA sediment sample showed a BaP level of 8000 µg/kg dry weight. Concentrations of these PAHs in urban soil were reported to be about two orders of magnitude higher than concentrations in rural soils.

Ambient air concentrations of these PAHs were also reported to be significantly higher in urban areas than in rural areas. In a study of seven urban areas, the highest PAH concentrations in air were reported for Detroit, MI, up to $21.3~\text{ng/m}^3$ benzo[g,h,i]perylene, probably due to the high degree of industrialization in that city. Data presented in a separate review of ambient levels of PAHs indicate that most rural air concentrations are less than $10~\text{ng/m}^3$; urban air concentrations up to $100~\text{ng/m}^3$ were reported for many of these PAHs.

5.3 HUMAN EFFECTS AND EXPOSURE

5.3.1 Human Toxicity

5.3.1.1 Introduction

Data on the effects of ingestion of individual polycyclic aromatic hydrocarbons included in the benzo[a]pyrene group (i.e., acenaphthylene, benz[a]anthracene, benzo[a]pyrene, benzo[b]fluoranthene, benzo[k]fluoranthene, benzo[g,h,i]perylene, chrysene, dibenz[a,h]anthracene, indeno-[1,2,3-c,d]pyrene) on man or laboratory animals are few. We have concentrated our efforts on benzo[a]pyrene (BaP), the prototype PAH carcinogen since regulatory action to reduce levels of BaP will also result in the simultaneous reduction of other PAHs as well as non-carcinogenic compounds, and BaP is a major source of exposure.

5.3.1.2 Pharmacokinetics

There are no pharmacokinetic data for this group of PAHs in humans. Considerable animal data are available, however, for BaP from which generalizations can be drawn for other PAHs in this group.

Absorption and Distribution

Animal studies indicate that BaP is readily transported across the intestines, primarily by passive diffusion (Rees et al. 1971). It is also easily absorbed through the lungs (Kotin et al. 1969, Vainio et al. 1976). Rodent studies indicate wide tissue distribution following absorption of BaP (USEPA 1980). In rats, BaP disappears from blood and liver very rapidly following intravenous administration; the half-time for BaP disappearance from liver is about 10 minutes. This rapid elimination phase is followed by a slower disappearance phase lasting 6 hours or more in which the concentration of BaP increases in body fat and fatty tissues (e.g., breast) (Schlede et al. 1970). Since BaP has been shown to induce microsomal enzyme activity, prior exposure can accelerate both the rate of disappearance from tissues and excretion of metabolites into bile.

Metabolism

Most tumorigenic PAHs in themselves are not direct carcinogens but require metabolic activation before biological activity such as PAH-induced carcinogenesis can be expressed. Current understanding of the mechanisms involved in this process (Yang et al. 1978; Sims 1976, Lehr et al. 1978, Selkirk et al. 1975a,b Jerina et al. 1977a,b, Selkirk 1977) is as follows: PAHs are metabolized by the cytochrome P-450 dependent microsomal mixed function oxidase (MFO) system (often designated as the aryl hydrocarbon hydroxylase system). This enzyme system is readily inducible by exposure to a variety of chemicals and is found in most

adult as well as fetal mammalian tissues, most especially in the liver. [Hepatic microsomal activity, although only a few percent of adult levels, has been detected in human fetuses as early as 6-7 weeks of gestation (Pelkonen 1976)]. The MFO catalyze the formation of reactive epoxide intermediates which are capable of forming covalent bonds with cellular constituents (DNA, RNA, proteins), and ultimately result in tumor formation. BP7,8-diol-9,10-epoxide is currently suspected to be the ultimate carcinogenic derivative of BaP. Two major groups of metabolites are formed: (1) water-soluble glutathione, glucuronide and sulfate conjugates, and (2) organo-soluble metabolites consisting of ring hydroxylated products such as phenols and dihydrodiols, quinones and labile epoxide intermediates. Figure 5-10 presents the possible metabolic pathways for BaP which are reasonably well understood.

A vital consideration in the extrapolation of risks associated with PAHs in general, and BaP specifically, is the metabolic differences, both qualitatively and quantitatively, between humans and other species. Numerous studies have demonstrated that these differences exist (Sims 1976, Wang et al. 1976, Selkirk et al. 1976). There is some evidence to indicate heritable variations in the inducibility of enzymes involved in the metabolic process occur in man but data are confounded by effects of age, seasonal variations and lack of a homogenous human population (Paigen et al. 1978). Freudenthal and coworkers (1978) reported considerable individual variations in BaP metabolism among lung microsome samples taken from the same species as well as qualitative and quantitative differences between man, monkey and rat. Similar findings have been reported by Cantrell $\underline{\text{et}}$ $\underline{\text{al}}$. (1979) and Marshall $\underline{\text{et}}$ al. (1979). The degree of individual genetic variation of some activating enzymes is suspected to be a key factor in an organism's susceptibility to PAH-induced carcinogenesis. This issue remains unresolved at this time.

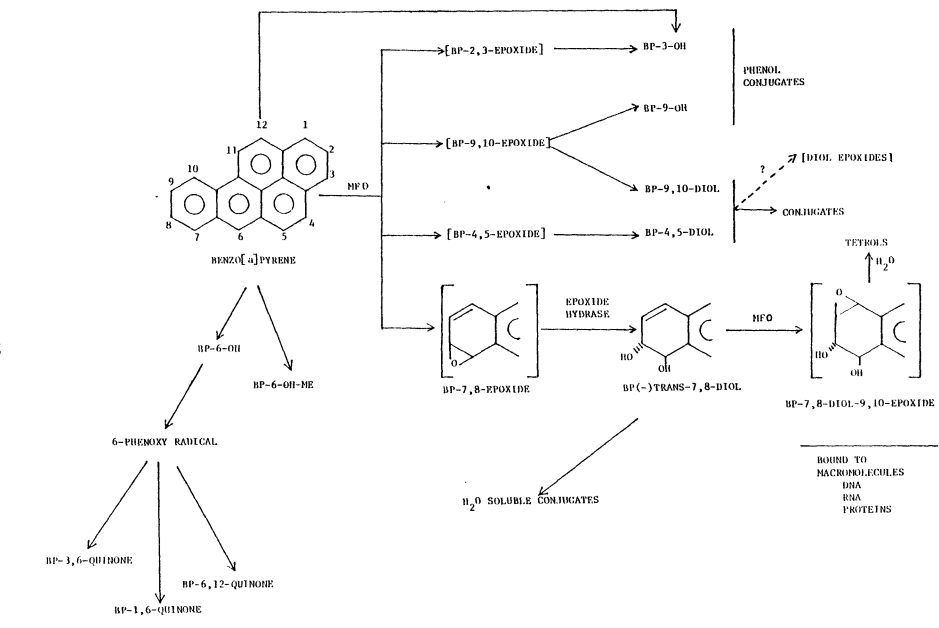
Excretion

The primary routes of BaP excretion in mice and rats are the hepatobiliary system and the feces, regardless of the route of administration. The dihydroxy-, 3-hydroxy-, and 6-hydroxy-derivatives have been detected in bile, liver and in the bowel (Berenblum and Schoenthal 1943, Falk et al. 1962, Sims 1967, 1970a,b). No extensive bioaccumulation as such is believed to occur.

5.3.1.3 Human and Animal Studies

Carcinogenicity

Several PAHs in the benzo[a]pyrene group are well established animal carcinogens, co-carcinogens and/or tumor initiators; others induce no tumorigenic responses. The capacity of individual PAHs to induce positive responses in humans is not so well established. This



Source: Lu et al. 1976; Selkirk 1977; U.S.EPA 1980.

FIGURE 5-10. POSSIBLE PATHWAYS OF BENZO[a]PYRENE METABOLISM

is primarily due to the fact that human exposures have not been to individual chemicals but rather to combinations as they occur in coke oven emissions, coal-tar, soot or from environmental exposures to tobacco smoke or exhaust fumes. Numerous studies have shown increased incidences of lung, skin and other types of cancer among workers exposed to coke oven emissions, coal gas, coal tar and pitch (IARC 1972). These studies, however, do not allow identification of the individual chemical(s) responsible, do not account for possible synergistic or co-carcinogenic effects resulting from other components, often are unable to clearly define exposure levels and generally are not amendable to quantifying human risk.

For the most part, available data for specific PAH compounds are from skin painting and subcutaneous or intramuscular injection experiments in mice. Few oral or inhalation experiments have been conducted, and those that are available are generally inadequate for risk assessment purposes.

Benzo[a]pyrene

Among the compounds in the benzo[a]pyrene group, BaP itself has been the most extensively studied. The published literature on BaP is vast. A few representative studies of the positive carcinogenic effects of BaP in a variety of animal models are presented in Table 5-47. BaP has been shown to be both a local and systemic carcinogen by oral, dermal and intratracheal routes. It is also a transplacental carcinogen, an initiator of skin carcinogenesis in mice and is carcinogenic in single dose experiments. Additional documentation of the carcinogenicity of BaP may be found in IARC (1972) or the Survey of Compounds . . . Tested for Carcinogenic Activity (1961, 1968, 1970, 1972, 1978).

Few studies have adequately examined the carcinogenic effects of orally administered BaP. Available studies are generally of very short duration with small test populations and frequently without appropriate control animals. Incidence data for the two best available oral studies for BaP (Neal and Rigdon 1967, Fedorenko and Yanysheva 1966) are presented in Table 5-48. Forestomach tumors were found in CFW mice fed BaP in their diet for an average of 110 days (Neal and Rigdon 1967) as well as in CC57 mice administered BaP by gavage once a week for 10 weeks and then maintained up to 19 months (Federenko and Yanysheva 1966). Although both these studies are by the oral route, several shortcomings raise questions as to their relevance to possible induction of cancer in humans from ingestion of BaP. Test populations in both studies were exposed for periods considerably less than lifetime and tumors were limited to the forestomach. The site of tumor development and the probable underlying mechanism are possibly unique to laboratory rodents. Unlike the fundus portion of the human stomach (which is homologous to the forestomach of rodents), the rodent forestomach is nonglandular and similar in histology to skin epithelium.

TABLE 5-47. CARCINOGENIC ACTIVITY OF BENZO[a]PYRENE BY VARIOUS EXPOSURES

Species (No.)	Major Effects	Dose	Exposure (# administrations)	Route	Incidence	Comments
A/HeJ female mice (12)	Forestomach tumors	1 mg/g d1et Ave. 4.8 mg BaP/mouse exposure	3 x/wk for 2 wks	Diet	12/12	Duration 29 wks
		Controls		• • • • • • • • • • • • • • • • • • •	0/12	
Ha/ICR female mice	Forestomach tumors	1 mg/mouse in corn	2 x/wk for 4 wks	Gavage	36/39	Duration 21 wks
		0.3 mg/mouse in corn oil	3 x/wk for 8 wks	Gavage	33/33	Duration 31 wks
		Control (no treat- ment)			0/19	Duration 31 wks
Mice	Tumors	0.2 mg	Single administration	Gavage	5/11	
		0.05 mg	11		0/9	
		0.012 mg in PEG	H (1		2/10	
Sprague Dawley female rats (9)	Mammary tumors	100 mg	Single administration	Oral	8/9	
C57B1./6J female mice (30)	Skin tumors	0.02 µmoles	Once every two wks for 60 wks	Skin painting	12/30	Mostly squamous
		0.1 µmoles	(Same as above)		26/27	

TABLE 5-47. CARCINOGENIC ACTIVITY OF BENZO[a]PYRENE BY VARIOUS EXPOSURES (continued)

Species (No.) CD-1 female mice (30)	Major Effects Papillomas	Dose 200 nunoles	Exposure (# administrations) Single application followed by twice weekly applications of 10 µg TPA for 23 weeks	Route Skin painting	Incidence 94%; 4.8 papillomas/ mouse	Comments Initiator of skin carcino- genesis
Sprague Dawley rats (13)	Sarcomas	0.2 µmoles	Alternate days for 30 doses	Subcutaneous in- jection	13/13	Average latency 101 days
Mice, male (14) Mice, female (16)	Sarcomas	2.4 µmoles	At monthly intervals for 3 treatments	Injection	13/14	Average latency: 129 days
(ID)		11 11	(Same as above)	11	8/16	160 days
Rats	Lung tumors	2.5 mg	Once monthly for 10 mo.	Intratracheal instillation	80%	Observed for 2
	·	0.25 0.05 0.01 0.002 0.0005 Mixed with a blood substitute BK-8 and India			43% 28% 14% 0% 0%	yr beyond treat- ment
Syrtan Golden hamsters	Respiratory tract tumors	1 mg 0.5 0.25 0.125 0.0625 1n saline	Once weekly for 52 weeks	Intratracheal Instillation	26/28 25/29 9/30 4/30 3/30	

TABLE 5-47. CARCINOGENIC ACTIVITY OF BENZO[a]PYRENE BY VARIOUS EXPOSURES (continued)

Species (No.)	Major Effects	Dose	Exposure (# administrations)	Route	Incidence	Comments
ST/A mice	Mammary tumors	10 mg	Single Injection	Intraperitoneal	2/10	At one year
Pregnant ICR/Ha mice	Lung adenomas; initiation of skin carcino- genesis	2-4 mg	Dams treated on days 11, 13 and 15 of gestation	Intraperitoneal or subcutaneous injections		Transplacental carcinogen

Source of data: USEPA 1980, IARC 1972.

TABLE 5-48. INCIDENCE OF ORALLY ADMINISTERED, BENZO[a]PYRENE-INDUCED FORESTOMACH TUMORS IN MICE

Strain	Treatment	Incidence of Forestomach Tumors	Comments	Reference
CC57 mice 2-3 mo.	1 mg BaP in 0.2 ml triethylene glycol, lx/wk for 10 wk by in-tubation (fasted)	23/27 (85%)	All mice were held for 19 months prior to sacrifice.	Fedorenko and Yanysheva (1966).
	As above except 0.1 mg	23/30 (77%)		
	As above except 0.01 mg	5/24 (21%)		
	As above except 1 μg	2/26 (8%)		
	As above except 0.1 μg	0/16 (0%)		
	Control	not given		
CFW mice	32.5 mg BaP mg/kg/day in the diet	66/73 (90%)	The various treatment groups were put on test	Neal and Rig- don (1967).
	As above except 13 mg	19/23 (83%)	at different times.	don (1707).
	As above except 6.5 mg	24/34 (71%)	The mice at different exposure levels varied in	
	As above except 5.85 mg	4/40 (10%)	age from 17 to 116 days	
	As above except 5.2 mg	1/40 (3%)	at the start of the experi- ment and were exposed for	
	As above except 3.9 mg	0/37 (0%)	an average of 110 days	
	As above except 2.6 mg	1/23 (4%)	(range: 70-197 days) over a 183 day average study	
	As above except 1.3 mg	0/24 (0%)	duration (range: 85-219	
	As above except 0.13 mg	0/25 (0%)	days).	
	Control	0/289 (0%)		

This suggests that the rodent forestomach may be unusually sensitive to repeated localized exposure to BaP akin to skin painting bioassays. This presumably local effect of ingested BaP on rodent forestomach epithelium, in the absence of tumors at other sites, appears to be of questionable relevance to humans with a wholly different stomach anatomy.

Other PAHs in the Benzo[a]pyrene Group

The carcinogenic activity of other PAHs included in the benzo[a]pyrene group is summarized in Table 5-49. No carcinogenicity data were found for acenaphthylene; the majority of other compounds in this group have been tested only in the mouse with exposures generally limited to skin painting experiments. Only benz[a]anthracene and dibenz[a,h]anthracene have been tested by the oral route. Both were found to be carcinogenic by the oral route (IARC 1972). Dibenz[a,h]anthracene induced forestomach tumors in mice following oral administration of this compound (IARC 1972, USEPA 1980). Repeated gavage administration of benz[a]anthracene (15 doses of 1.5 mg over 5 week period) produced a 95% incidence of lung adenomas, 64% incidence of hepatomas and a 3% incidence of forestomach papillomas in the test population of 59 B6AF1/J mice. Another group of 20 mice given only two treatments 3 days apart developed 16 hepatomas and 17 lung adenomas. Controls for these test groups had 2 hepatomas, 10 lung adenomas and no forestomach tumors among 59 control animals (IARC 1972).

Dibenz[a,h]anthracene, benz[a]anthracene, benzo[b]fluoranthene, chrysene and indeno[1,2,3-c,d]pyrene are 'complete carcinogens for mouse skin and all are also initiators of skin carcinogenesis in this species (USEPA 1980, IARC 1972, Habs et al. 1980). No carcinogenic response was noted in a skin painting study with benzo[k]fluoranthene in NMRI mice treated twice a week with up to 9.2 pg/mouse/application for their lifetime (Habs et al. 1980).

A pronounced co-carcinogenic effect was observed in a single experiment conducted with benzo[g,h,i]perylene (2000 μ g) plus 5 μ g BaP applied to skin of ICR/Ha Swiss mice 3 times per week for 52 weeks (USEPA 1980).

Local sarcomas have also been produced at the site of subcutaneous or intramuscular injections of benzo[b]fluoranthene, chrysene, dibenz-[a,h]anthracene and indeno[1,2,3-c,d]pyrene (USEPA 1980, IARC 1972).

Ultimate Carcinogenic Metabolites

Investigators have long sought a common molecular feature among carcinogenic PAHs such as BaP. Recent attention has focused on the increased reactivity of diol epoxide metabolites of PAHs in which the oxirane oxygen forms part of a "bay region." Carcinogenicity studies conducted with various potential oxidative metabolites of BaP, for example, suggest that the 7,8-diol-9,10-epoxide is the most active metabolite (Wislocki et al. 1977).

TABLE 5-49. SUMMARY OF THE CARCINOGENIC ACTIVITY FOR OTHER POLYCYCLIC AROMATIC HYDROCARBONS IN THE BENZO[a]PYRENE GROUP!

Chemical	Species (No.)	Major effects	Dose	Exposure (# administrations)	Route	Incidence	Comments
Benz[a]Anthracene	Mice (13)	No tumors in 16 months	0.5 mg in mineral oil	Single dose	Gavage		
Benz[a]Anthracene	Mice (27) Papilloma of forestomach		0.5 mg in mineral oil	8 or 16 X at 3-7 day intervals	Gavage	2/27	
	Mice (16)	No tunors	Mineral oil	Vehicle controls	Gavage	0/16	
Benz[a]Anthracene	B6AF1/J mice (59)	Lung adenomas; hepatomas; papil- lomas (stomach)	1.5 mg/mouse metho- celaerosol OF	15 treatments in 5 weeks	Gavage	56/59; 38/59; 2/59	2 hepatomas in 20 animals with median age of death 547-600 days
Benz[a]Anthracene	Hice (20)	Hepatomas; lung adenomas	1.5 mg/mouse	2 treatments, 3 days apart	Gavage	16/20; 17/20	Median death age 547-600 days
Controls	Mice (59)	Hepatomas; lung adenomas	Untreated			2/59; 10/59	
Benz[a]Anthracene	MIce (20)	Papilloma	0.4% in mineral oil	2 X/wk, 68 wks	Skin Painting	1/20	No control data
Benz[a]Anthracene	C3II/IIe Mice	Tumors (type not specified - mostly malignant)	.002% in toluene .02% " " .2% " " 1.0% " "	3 X/wk/50 wks	Skin Painting " "	0/32 1/18 3/32 8/29	
			.0002% in dodecane .002% " " .02% " " 0.2% " "	01 00 00 00 00 01 00 00 00 00 00 00 01 00 00 11 00 01 00 10	11 11 17 01 11 11 11 11	4/31 8/21 4/20 11/21 17/22	Dodecane is a cocarcinogen
Benz[a]Anthracene	Mice (75)	Tumors Papillomas	.05% in acetone +5% croton oil	Alternate once weekly applications	Skin Painting	17/17 at 9 month 18/9 at 12 month	
		Tumors	controls - 5% croton oil	,, ,,	** **	1/13	

TABLE 5-49. SUMMARY OF THE CARCINOGENIC ACTIVITY FOR OTHER POLYCYCLIC AROMATIC HYDROCARBONS IN THE BENZO[a]PYRENE GROUP (continued)

Chemical	Species (No.)	Major effects	Dose	Exposure (# administrations)	Route	Inc1dence	Comments
Benz[a]Anthracene	Mice (20)	Skin tumors	6 mg/mouse total followed by croton oil	10 X over 5 wks weekly applications		tumors in 18 mice	BA alone gave no tumors
Benz[a]Anthracene	Hamsters (10) (Syrlan gold.)	No tumors	0.5% in mineral oil	Bi-weekly for 10 wks	Skin Painting		6/10 alive at 50 wks, all dead at 85 wks
Benz[a]Anthracene	C57Bl, mice	Tumors	0.05 mg in tricaprylin 0.2 mg " " 1.0 mg " " 5.0 mg " " 10.0 mg " "	Single " " " "	Injection " " " " "	5/43 mice 11/43 15/31 49/145 5/16	Surviving at 9 months
Benz[a]Anthracene	C57BL mice	Sarcomas	l mg/mouse in arachis oil	10 weekly injections	Subcutaneous	14/20 in 146 179 days	-
Benz[a]Anthracene	Rats (20)	No tumors	1.9 mg in tri- caprylin	Single application	Subcutaneous	0/9 alive at 14.5 months	
Benzla Muthracene	Rats (29 °)	Mammary tumors	13 mg/kg	3 injections at 3 day intervals	Injection	0/29 at 98 d	ays
Benz La JAnthracene	MIce (52)	Carcinomas (bladder)	2 mg implant in paraffin wax	Single	Bladder implant.	17/52	In 40 weeks or more; 3.8% inci- dence in controls
Benz[a]Anthracene	BALB/C, new- born and 2-8 days old	Pulmonary adenomas, adenocarcinomas	50 μg in gelatin (aq)	Single injection	Injection	?	More tumors in newborns
Benzo[b]Fluoran- thene	Mice (20/ group)	Papillomas Carcinomas Papillomas Carcinomas Papillomas Carcinomas	0.5% in acetone " " " 0.1% in acetone " " " 0.01% in acetone	3 X/week " " " " " " " " "	Skin painting	g 100% 90% w/in 8 65% w/in 12 85% w/in 12 1/10 survi 14 mo	mos. No Control vors after

TABLE 5-49. SUMMARY OF THE CARCINOGENIC ACTIVITY FOR OTHER POLYCYCLIC AROMATIC HYDROCARBONS IN THE BENZO[a]PYRENE GROUP¹ (continued)

Chemical	Species (No.)	Major effects	Dose	Exposure (# administrations)	Route	Incidence	Comments
Benzo[b]Fluoran- thene	Mice (20)	No tumors	1.0 mg in acetone	Single application	Skin painting	***	63 weeks observ.
- Tener	Mice	Papillomas Carcinomas	1.0 mg in acctone and croton resin	Single application Repeated applica- tion	Skin painting	18/20 5/20	Initiator of skin carcinog.
Benzo[b] Fluoran- thene	16 d, 14 9 XVIInc/z mice	Sarcoma at injec- tion site	0.6 mg/injection	3 injections in 2 months	Subcutaneous	18/24	Avg. latent period of 4.5 months
Benzo[b]Fluoran- thene	NMRI (40)	Papillomas, sarcomas, and carcinomas	3.4 pg in acetone 5.6 pg " " 9.2 pg " "	2 X/wk for lifetime	Skin painting	5.3% 14.7% 54.1%	No tumors in controls
Benzo[k]Fluoran- thene	NMRI (40)	No tumors	3.4 pg in acetone 5.6 pg " " 9.2 pg " "	2 X/wk for lifetime	Skin painting """ """"		No tumors in controls
Chrysene	Mice (20)	Pap (11 omas Carcinoma	1.0% in acetone	3 X/wk for ?	Skin painting	9 After 8 8 mos.	Obvious shortened life-span
Chrysene	CF1 mice	Epithelloma	0.2% in acetone	Biweekly for 31wks	Skin painting	1/16	Service and the service of the servi
Chrysene	Mice	No tumors	1 mg in acetone	Single application	Skin painting		63 wks observ.
	Mice	Papillomas Carcinomas) mg in acetone and croton resin	Single application Repeated applica- tion	Skin painting	16/20 2/20	Initiator of skin carcinog.
Chrysene	Mice (50)	No tumors	2 mg (purified)	Single injection	Subcutaneous	And the same of	45 wks observ.
Chrysene	Mice (30)	Negative	10 mg/implantation	2 implantations	Subcutaneous		
Chrysene	Mice (40-50) (C57BL)	Sarcomas	5 mg in trycaprylene	Single injection	Subcutaneous	5/22?	Avg. induction time = 271 days 22 alive at 150 days

TABLE 5-49. SUMMARY OF THE CARCINOGENIC ACTIVITY FOR OTHER POLYCYCLIC AROMATIC HYDROCARBONS IN THE BENZO[a]PYRENE CROUP¹ (continued)

Chemical	Species (No.)	Major effects	Dose	Exposure (# administrations)	Route	Incidence	Comments
Chrysene	M1ce (57BL)	Tumors (?)	l mg in arachis oil	10 injections	Injection	2/20	No tumors in vehicle contro
Chrysene	Rats	Sarcomas	2-3 mg/injection	Repeated (?)	Injection	4/10	2/10 sarcomas in vehicle controls
Chrysene	Rats (10)	No tumors	0.05% (aq.)	Bl-weekly	Injection	0/4 at 18 mos.	
Chrysene	Mice (50)	No tumors	2 mg in lard	Single	Injection		50 wk observ.
Dibenz[a,h]An- thracene	M1 ce	Tumors (1 carcinoma of forestomach)	9-19 mg total dose	Over 5-7 months	Dietary	7/22 after 1 year	
Dibenz[a,h]An- thracene	Mice (20)	Squamous-cell carci- noma, Papilloma of forestomach	0.4 mg/day in oil emulsion	W/in 406 days	Drinking water	'2/20 11/20	****
	Mice	Lung, heart and intestinal tumor	0.4 mg/day in NaOH	?	Drinking water	?	
Ofbenz[a,h]An- thracene	Mice (42) DBA/2	Pulmonary adenoma- tosis Aveologenic carci- noma Hemangio-endothe- lioma Mammary carcinoma	0.76-0.85 mg/day in olive oil """	200 days	Drinking water """	27/27 24/27 16/27 12/138	
	Mice (35)	Mammary carcinoma Pulmonary adenoma- toses	c o	NTROLS		0	
Ofbenz[a,h]An- Thracene	Mice (BALB/c) Pseudopreg.	Mammary carcinoma	0.5% in almond ofl (15 mg total dose)	Twice weekly for	Stomach tube	1/20	

TABLE 5-49. SUMMARY OF THE CARCINGENIC ACTIVITY FOR OTHER POLYCYCLIC AROMATIC HYDROCARBONS IN THE BENZO[a]PYRENE GROUP (continued)

Chemical	Species (No.)	Major effects	Dose	Exposure (# administrations)	Route	Incidence	Comments
Dibenz[a,h]An- thracene	C3H mice	Mammary tumors	0.25% in benzene Controls	Twice weekly Untreated	Skin painting	10/11 50%	
	Swiss mice (20)	Skin tumors	30 ng/dose in ace- tone-benzene	Bi-weekly	Skin painting	16/20	
DIbenz[a,h]An- thracene	ICR/Ha Swiss mice	Tumors (some carcinomas)	0.001% 0.01% 0.1%	3 X weekly	Skin painting		carcinomas)
Dibenz[a,h]An- thracene	C3H mice	Local sarcomas	0.00019 mg/dose 0.0078 mg/dose 0.016 mg/dose 0.03 mg/dose 0.06 mg/dose 0.12 mg/dose 0.25 mg/dose	Single dose -	Injection	2/79 6/40 6/19 16/21 20/20 21/23 19/21	Lowest effective dose - 0.0019 mg Avg. latent period = 3.7 mos.
Dibenz[a,h]An- thracene	Mice (20)	Sarcomas	0.0125 mg in lard	5 injections	Injection	4/20	
Dibenz[a,h]An- thracene	Mice (new- born)	local sarcomas Lung Adenomas	> 0.08 μg > 0.2 μg	5 injections	Subcutaneous Subcutaneous		Dose-related appearance Increased Inci- dence
₿1benz[a,h]An-	Rats	Tumors	8 mg/injection	9 injections at monthly intervals	Injection	11/40	W/in l year
Dibenz[a,h]An- thracene	Rats	Sarcomas	0.1 mg in olive oil	Stugle dose	Injection "	3/9 6/10	
Dibenz[a,h]An- thracene	Guinea pigs	Sarcomas	8-48 mg (total dose) In oll	Multiple	Subcutaneous	2/25	After 19 mos.
DIbenz[a,h]An- thracene	Fowl	Sarcomas	0.4% in lard	Single dose	1.m.	15/31	W/in 45 mos.

TABLE 5-49. SUMMARY OF THE CARCINOGENIC ACTIVITY FOR OTHER POLYCYCLIC AROMATIC HYDROCARBONS IN THE BENZO[a]PYRENE GROUP (continued)

Chemical	Species (No.) Major Effec		Major Effects	Dose	Exposure (# administrations)	Route I	ncfdence	Comments
Dibenz[a,h]An- thracene	Mice		Lung adenomas	> 0.1 mg colloidal dispersion (aq.)	Single dose	1.v.		Dose-response relationship
Dibenz[a,h]An- thracene	thracene nomas Othenz[a,h]An- Nice Lung 1		Renal adenocarci- nomas	0.3-0.5 mg in olive	Single dose	Intrarenal	26% 10/10 8/65 33/65	3% incidence controls
Dibenz[a,h]An- thracene			Lung Tumors	36 μmoles/kg	Single dose	1.v.		
Dibenz[a,h]An- Mice thracene			Papilloma Epithelioma	?	?	Skin painting		63% tumor incidence/239 day avg. latent period = 26 Iball Index
Dibenz[a,h]An- thracene	Mice (1 Mice (1	•	Aveologenic carcinoma and pulmonary adenomatosis	179 mg (total) 236 mg (total) (in olive oil)	237-279 days	Drinking water	77% 100%	
Indeno[1,2,3-c,d]1	'yrene	Mice (?) No tumors	0.05 or 0.1% in dioxane	3X/wk for 12 months	Skin painting		
Indeno[1,2,3-e,d]1	'yrene	MI ce	No tumors No tumors Papillomas Carcinomas Papillomas Carcinomas Skin carcinogen-	0.01% In acetone 0.05% " " 0.1% " " 0.1% " " 0.5% " " 0.5% " "	3 X/wk for 12 months """"""""""""""""""""""""""""""""""""	Skin painting """ """" """" """"	0 0 6/20 3/20 7/20 5/20	Dose-response Relationship """"""""""""""""""""""""""""""""""""
1ndeno[1,2,3-c,d]1	Pyrene	Mice of	Sarcomas Sarcomas	0.6 mg in olive	day interval 3 injections at 1 month intervals	Injection	10/14 1/14	W/in 265 days W/in 145 days
	Pyrene	NMR1 m1 (40)	ce No tumors	3.4 pg in acetone 5.6 pg " " 9.2 pg " "	2 X/wk for lifetime	Skin painting """ """		No tumors in controls.

¹ Source of data: USEPA 1980, JARC 1972, Habs et al. 1980.

Wislocki and coworkers (1977) found that BP7.8-oxide was the most active carcinogen among 4 BaP arene oxides tested (and only slightly less effective than BaP) following skin application of 0.3-0.4 μmol to C57BL/6J mice once every two weeks for 60 weeks. Since 4-,5-,6-,7-, 8-,9- and 10-hydroxyBP were all inactive as complete carcinogens at 0.4 µmol/dose (Kapitulnik et al. 1976), carcinogenic activity could not be attributed to phenolic isomerization products of BP7,8-oxide. Further tests with metabolic products of BP7,8-oxide implicated BP7,8-dihydrodiol as a more potent proximate carcinogen; the dihydrodiol was presumably formed when BP7,8-oxide was hydrated by epoxide hydrase (Jerina et al. 1977a,b). BP7,8-dihydrodiol induced 100% incidence of skin tumors (as did BaP itself) compared to <20% incidence induced by BP7,8-oxide at doses of 0.15 μ mol by the above schedule; at lower doses the dihydrodiol was more active than BaP (Levin et al. 1977a,b). The lack of carcinogenic activity of 7,8,9,10-tetrahydro-BP7,8-epoxide and 7,8,9,10-tetrahydro-BP7,8-diol at 0.3 µmol/dose indicated that the 9,10double bond was important in metabolic transformation to ultimate carcinogenic products (Levin et al. 1976a,b, Jerina et al. 1978). The most probable candidates for this ultimate metabolite, would again be the BP7,8-diol-9,10-epoxides formed by oxidation of the 7,8-dihydrodiol at the 9,10-position by the monooxygenase system (Conney et al. 1980, Jerina et al. 1977a,b). In tests with newborn Swiss-Webster mice, intraperitoneal injection of a 28 nmol dose of either BaP, BP7,8-dihydrodiol or BP7,8-diol-9,10-epoxide-2 resulted in a definite sequential increase in tumorigenic activity of BaP, BP7,8-dihydrodiol and BP7,8diol-9,10-epoxide-2, respectively, clearly implicating the dihydrodiol as the proximate carcinogenic metabolite and the epoxide as the ultimate carcinogenic metabolite. The BP7,8-diol-9,10-epoxide-1 isomer was too toxic for an adequate assessment of tumorigenic activity (Kapitulnik et al. 1977).

These and other studies have generated the "bay-region" (i.e., hindered area between the 10- and 11-positions) theory to correlate structure of polycyclic hydrocarbons with carcinogenic activity. Presumably benzylic carbonium ions derived from epoxides of a saturated angular benzo ring have a greater ease of formation when the carbonium ion is located in the bay region, thus enhancing the chemical reactivity and perhaps the mutagenicity and carcinogenicity of the epoxide (Jerina et al. 1977a, 1977b, 1978, Conney et al. 1930).

A number of recent experiments with bay region diol epoxides of benz[a]anthracene, dibenz[a,h]anthracene and chrysene indicate that they, also, are more active than the parent compounds or other oxidative metabolites.

Skin painting studies with various metabolites in CD-1 mice suggest that BA3,4-dihydrodiol is at least 10 times more tumorigenic than benz-[a]anthracene itself or other metabolites (Wood et al. 1977b,c, Levin et al. 1978, Slaga et al. 1978). In newborn mice treated with various

metabolites of benz[a]anthracene, BA3,4-dihydrodiol induced at least 30-fold more pulmonary adenomas than the parent compound or 4 other dihydrodiols tested (Wislocki et al. 1978).

Similarly, Buening and coworkers (1979b) found DBA 3,4-dihydrodiol to be equal to dibenz[a,h]anthracene and considerably greater than DBA 1,2-dihydrodiol, DBA 5,6-dihydrodiol and DBA H₄-3,4-diol in tumor-inducing activity on mouse skin following single applications. In the same study, the incidence of pulmonary adenomas induced in newborn Swiss-Webster mice following intraperitoneal injection was similar for dibenz[a,h]anthracene and DBA 3,4-dihydrodiol, while tumor incidences caused by DBA 1,2-dihydrodiol and 5,6-dihydrodiol were significantly lower. Again, the much lower tumorigenic activity on mouse skin of DBA 3,4-diol compared to DBA 3,4-dihydrodiol is consistent with the prediction of the bay-region theory that metabolism at the 1,2 double bond would yield the ultimate carcinogenic diol epoxide, DBA 3,-4-diol-1,2-epoxide (Buening et al. 1979b).

Initiation-promotion studies in CD-1 mice with chrysene and various chrysene metabolites applied once at doses of 0.4, 1.25 or 4.0 µmoles and followed by twice weekly applications with TPA, a phorbol ester, for 25 weeks resulted in significantly greater tumorigenic activity for chrysene 1,2-dihydrodiol above chrysene itself (Levin et al. 1978). Similar findings were reported by Buening et al. (1979a) who found enhanced lung tumor activity for chrysene 1,2-dihydrodiol in newborn mice given intraperitoneal injections on days 1,8 and 15 of life.

In summation, benzo[a]pyrene has been most extensively studied of all PAHs in the benzo[a]pyrene group. BaP has been shown to be both a local and systemic carcinogen by oral, dermal and intratracheal routes. It is also a transplacental carcinogen and an initiator of skin carcinogenesis in mice. Few studies have adequately examined the carcinogenic effects of orally administered BaP but it has been shown to induce forestomach tumors in mice exposed to BaP by the oral route. The absence of tumors at sites other than in the forestomach and the unique anatomy of rodent forestomach in contrast to humans raises questions as to the significance of the induction of forestomach tumors in mice to human risks associated with the ingestion of BaP.

With respect to other PAHs included in this group, benz[a]anthracene and dibenz[a,h]anthracene are both carcinogenic in mice by the oral route as well as complete carcinogens for mouse skin as are benzo-[b]fluoranthene and indeno[1,2,3-c,d]pyrene. Benzo[g,h,i]perylene is a co-carcinogen with BaP and benz[a]anthracene, benzo[b]fluoranthene, chrysene and indeno[1,2,3-c,d]pyrene are all initiators of skin carcinogenesis in mice.

Recent studies on the mechanisms of carcinogenic activity of some PAHs suggest increased reactivity of the diol epoxides of PAHs in which the oxirane oxygen is located in the "bay-region." Experiments

with possible metabolites of BaP indicate that one or more of the BP7,8-diol-9,10-epoxides are the ultimate carcinogenic form. The bay-region theory of polycyclic hydrocarbon carcinogenesis is further supported by findings of increased carcinogenic activity of the bay-region diol epoxides of benz[a]anthracene, dibenz[a,h]anthracene and chrysene above that of the parent compounds.

Adverse Reproductive Effects

BaP appears to exert little effect on the developing embryo (Bulay and Wattenberg 1970). Rigdon and Rennels (1964) found only one malformed fetus among 7 litters of rats from dams exposed to 1 mg BaP/g diet during gestation. Increased resorptions and dead fetuses were noted, however.

Juchau and coworkers (1978) reported that several human fetal tissues (i.e., liver, lung, kidney, adrenals and placenta) possess the requisite monooxygenase enzyme systems needed to bioactivate and biotransform BaP into metabolites that induce positive mutagenic effects in the Ames Salmonella assay.

No other data were available for compounds in this group.

Mutagenicity

A summary of the mutagenic activity exhibited by compounds comprising the benzo[a]pyrene group is presented in Table 5-50. No data were found for benzo[k]fluoranthene or indeno[1,2,3-c,d]pyrene. BaP is the most active mutagen of the group, exhibiting positive mutagenic responses in all of the test systems including induction of in vivo sister chromatid exchange in hamster cells (Bayer and Bauknecht 1977, Roszinsky-Köcher et al. 1979, Sirianni and Huang 1978) and chromosomal aberrations in both spermatogonia and bone marrow cells of hamsters in vivo (Basler and Rohrborn 1978; Roszinsky-Köcher et al. 1979). Studies conducted with various potential metabolites of BaP suggest that the BP7,8-diol-9,10-epoxide is the most potent mutagen in the presence of a metabolic activation system and may in fact be the ultimate carcinogen (Conney et al. 1977, Wislocki et al. 1976a,b, Wood et al. 1976, Huberman et al. 1976, Newbold and Brookes 1976).

Mixed responses have been found for several of the other compounds in this designated group. In vivo studies with benz[a]anthracene have been generally positive. Increased chromosomal aberrations were observed in mouse oocytes and hamster spermatogonia and bone marrow cells (Peter et al. 1979, Basler and Rohrborn 1978). Roszinsky-Köcher and coworkers (1979) noted a weak induction of sister chromatid exchange in hamster cells exposed in vivo to benz[a]anthracene. Both positive and negative results have been reported in host-mediated assays with Salmonella typhimurium TA1538 (Simmon 1979, Simmon et al. 1979, Rosenkranz and Poirier 1979). Benz[a]anthracene induced positive mutagenic

TABLE 5-50. SUMMARY OF MUTAGENIC ACTIVITY FOR COMPOUNDS COMPRISING THE BENZO[a]PYRENE GROUP

Compound Acenaphthylene		rations In Vitro	Sister Chromatid Exchange In Vivo	Host Mediated Assay	Mammalian Cells In Vitro			11a typ			Unscheduled DNA Synthesis	References Kaden <u>et al</u> . 1979.
Benz[a]anthra- cene	+		ŧ	<u>t</u>	-	+*	+*	±*	<u>*</u>	*	+	Basler and Röhrborn 1978; Huberman 1977; Huberman and Sachs 1976; Kaden et al. 1979; Martin et al. 1978; Peter et al. 1979; Rosenkranz and Poirler 1979; Roszinsky-Köcher et al. 1979; Simmon 1979; Simmon et al. 1979.
Benzo[b]fluor- anthene	_		<u> </u>				•					Roszínsky-Köcher <u>et al</u> . 1979.
Benzo[g,h,1] perylene						+**		_**	+**	+*		Gibson <u>et al</u> . 1978; Kaden <u>et al</u> . 1979.
Benzo [a] pyrene	4	+	+		+	+*	*		+*	*	+	Basler and Röhrborn 1978; Bayer and Bauknecht 1977; Hopkin and Perry 1980; Hsu et al. 1979; Huberman 1977; Huberman and Sachs 1976; Kaden et al. 1979; Lake et al. 1978; Martin et al. 1978; Matsucka et al. 1979; Mishra et al. 1978; Popescu et al. 1977; Rosenkranz and Poirier 1979; Roszinsky-Köcher et al. 1979; Rudiger et al. 1976; Salamone et al. 1979; Siranni and Huang 1978.

Legend: - = no significant induced mutations; + = significant induced mutations; t = variable or weak response.

^{*} Tested with liver microsomal activation.

^{**} Tested with $^{60}\mathrm{C}$ gamma irradiation activation.

TABLE 5-50. SUMMARY OF MUTAGENIC ACTIVITY FOR COMPOUNDS COMPRISING THE BENZO[a]PYRENE CROUP (continued)

Compound	Chromosomal Aberrations In In Vivo Vitro	Exchange	llost Mediated Assay	Mammallan Cells In Vitro	Cells In Vitro TA98 TA100 TA1535 TA1538 TM677		Unscheduled DNA Synthesis	References			
Chrysene	<u>.</u>	±	-	-	_*	* ±	_*	_*	+*		Basler et al. 1977; Basler and Röhrborn 1978; Huberman et al. 1972; Kaden et al. 1979; Kamei 1980; Poirier and de Serres 1979; Richter-Reichhelm et al. 1979; Rosenkranz and Poirier 1979; Roszinsky-Köcher et al. 1979; Salamone et al. 1979; Simmon et al. 1979; Simmon 1979.
Dibenz[a,h] anthracene	-	Ŧ		+	_**	*	**	+*	+	+	Gibson et al. 1978; Huberman 1977; Wood et al. 1978; Huberman and Sachs 1976; Kaden et al. 1979; Lake et al. 1978; Martin et al. 1978; Roszinsky- Köcher et al. 1979;

responses in in vitro bacterial tests with several strains of Salmonella typhimurium in the presence of liver microsomal activation (Kaden et al. 1979, Simmon 1979) and induced unscheduled DNA synthesis in Hela cells (Martin et al. 1978). No significant increase in the number of mutants was seen, however, in hamster cells exposed in culture (Huberman and Sachs 1976). Tests with possible metabolites of benz[a]anthracene indicate that the bay-region BA 3,4-diol-1,2-epoxides are the most potent mutagens and probable ultimate carcinogens for this compound (Wood et al. 1977a).

Weak induction of sister chromatid exchange but no induction of chromosomal aberrations of marrow cells was observed in hamsters exposed to dibenz[a,h]anthracene in vivo. Positive induction of unscheduled DNA synthesis was noted in both human epithelial and Hela cell cultures (Lake et al. 1978, Martin et al. 1978) and a significant number of mutations observed in V79 hamster cells exposed in culture in the presence of metabolic activation (Huberman and Sachs 1976). Mixed results have been reported in tests with various strains of Salmonella typhimurium (Kaden et al. 1979; Wood et al. 1978, Gibson et al. 1978).

In contrast to the generally positive findings for BaP, benz[a]—anthracene and dibenz[a,h]anthracene, results with chrysene have been either negative or, at most, only weakly positive. In vivo tests indicate no induction of chromosomal aberrations in hamster bone marrow cells (Basler et al. 1977, Basler and Rohrborn 1978), a weak sister chromatid exchange response in this species (Roszinsky-Köcher et al. 1979) and a slight increase in aberrations in mouse oocytes (Basler et al. 1977, Basler and Rohrborn 1978, Roszinsky-Köcher et al. 1979).

Generally negative findings were noted in tests with Salmonella typhimurium (Basler et al. 1977, Simmon et al. 1979, Rosenkranz and Poirier 1979) and negative results reported for a host-mediated assay (Poirier and de Serres 1979).

Limited data are available for acenaphthylene, benzo[b]fluoranthene and benzo[g, h, i]perylene. A single experiment with acenaphthylene produced positive findings in one strain of Salmonella typhimurium TM677 in the presence of rat liver activation but only at concentrations that were toxic to the bacteria (Kaden et al. 1979). An in vivo study with benzo[b]fluoranthene resulted in a weak induction of sister chromatid exchange but no significant induction of chromosomal aberrations in hamster marrow cells (Roszinsky-Köcher et al. 1979). Benzo[g,h,i]perylene has been tested in Salmonella typhimurium, producing negative results in strain TA1535 (base-pair mutant) and positive results in strains TA1538 and TA98 (frameshift mutants) with 60 Co gamma irradiation activation (Gibson et al. 1978). Positive results were also noted in S. typhimurium TM677 in the presence of liver microsomal activation (Kaden et al. 1979).

Other Toxic Effects

The major focus of studies conducted with PAHs in the BaP group has been their potential for inducing carcinogenic effects. Overt signs of toxicity are generally not evident except for the carcinogenic PAHs and then only at doses sufficient to produce a high tumor incidence (NAS 1977).

Normally proliferating tissues are the selected targets for carcinogenic PAHs such as BaP. This may be attributable to specific attack on the DNA of cells in the S-phase of the mitotic cycle (Philips et al. 1973). Other contributing factors are alterations of enzyme activity and immunologic competence. A single carcinogenic dose of BaP was reported to produce a prolonged depression of the immune response to sheep red blood cells (Stjernsward 1966, 1969). Damage to the hematopoietic and lymphoid systems has been frequently reported in experimental animals (USEPA 1980).

Bond and coworkers (1981) have noted the development of atherosclerotic plaques in groups of eleven 4-week-old Sc strain chickens injected weekly for 20 weeks with 0.1, 1 or 10 mg/kg BaP dissolved in dimethyl sulfoxide. The specific route of exposure was unspecified. Solvent and untreated control groups were also included in the study. At 20 weeks, detectable lesions were found in 44, 45 and 75% of animals in the 0.1, 1 and 10 mg/kg BaP groups, respectively, compared to 50% of solvent controls and 22% of untreated control animals.

5.3.1.4 Human Risk Considerations -- Benzo[a]pyrene Group

Carcinogenicity of Benzo[a]pyrene

The quantitative estimate of potential carcinogenic risk to humans resulting from ingestion of benzo[a]pyrene (BaP) is calculated below. Ideally, human data relating carcinogenic response at known daily dose levels in the dose range of interest would be utilized in an appropriate mathematical model to predict risk. Usually, such ideal data are not available and extrapolations of risk must be made from either:

- 1) epidemiological data involving relatively high level exposures such as may occur in occupational settings, or
- 2) controlled experiments on laboratory animals involving very high total dosages.

In the first case, the overriding uncertainty is often in the data themselves; the exposure levels, lengths of exposure, and even response rates are usually "best estimates," and, furthermore, unknown factors (e.g., co-existent carcinogenic exposure) may bias the data. In the second case, the data are usually more accurate, but the quantitative

extrapolation from animal models to humans is uncertain. At present, there is no universally accepted solution to this species to species extrapolation problem. In short, in the former case one has relevant data of questionable accuracy, whereas in the latter case, one has accurate data of questionable relevance. Further complicating the issue is the present lack of an indisputable basis for judging the relative merits of the various mathematical models relating dose and effect.

The available data concerning human and other mammalian effects have been presented above. Many experiments involving exposure of laboratory animals to BaP have been done, but surprisingly few experiments have been done in which significant populations of animals were given a wide range of doses of BaP. Two ingestion studies are considered whose data appear to lend themselves to dose/response extrapolation, one by Federenko and Yanysheva (1966) on CC57 mice, the other by Neal and Rigdon (1967) on CFW mice. Three commonly used mathematical models have been applied to each set of data to extrapolate the potential carcinogenic risk of BaP to humans. These assessments of potential human risk are subject to important qualifications:

- The experimental data chosen for extrapolation are for forestomach tumors induced either by weekly intubation (Federenko and Yanysheva 1966) or incorporation of BaP into the diet (Neal and Rigdon 1967). While apparently relevant with respect to route of administration, the site of tumor production and probable underlying mechanism are possibly unique to laboratory rodents because of the anatomy of the rodent stomach. The forestomach in both rat and mouse is non-glandular and has a stratified squamous epithelium with a cornified covering (Chiasson 1979). The fundus portion of the human stomach (homologous to the forestomach in rodents) is glandular and has a columnar epithelium. The similarities between the epithelium of rodents and that of skin suggest that the induction of forestomach tumors in mice may be analogous to the induction of skin tumors in skin-painting bioassays. This possibility calls into question the relevance of forestomach tumors in mice to the potential for human cancer from BaP ingestion. Although it is recognized that a carcinogen can cause different types of cancer in varying target organs in different species, even when the same route of exposure is used, it is considered possible that the forestomach tumors in mice induced by BaP may represent an unusually sensitive assay of the carcinogenicity of BaP by the oral route.
- The EPA guidelines call for the adjustment of dose between species to be based on body surface area not on body weight (USEPA 1979). In the case of forestomach tumor induction in rodents, neither adjustment would appear to reflect the presumably local effect of BaP on the rodent forestomach epithelium. If it were hypothesized that stomach tumors from BaP ingestion are unlikely in humans because of the differences in anatomy between rodent and human stomach, but other tumor sites were considered possible, one could reasonably adjust upwards

the human equivalent daily dose to account for the wider distribution in the body and consequent dilution of BaP. Conceivably, however, stomach tumors (or other intestinal tract tumors) are possible in humans due to ingested BaP. The analysis that is carried out below adjusts dose on a body surface basis, but it is regarded as the probable worst case — tending to overstate the risk due to BaP ingestion in humans.

 Both animal studies were of less than lifetime duration especially in the study by Neal and Rigdon (1967). Doses were administered for about half the duration of the experiment in both studies. In order to adjust animal dose schedule to an equivalent daily lifetime human dose, two assumptions were utilized as recommended by the USEPA (1979). First it was assumed that the response would be the same if the total dose was distributed evenly over the entire experimental lifetime of the animal. Second, it was assumed that a short-duration study attributable to either early mortality or early termination leads to an over-estimation of the dose required to produce the observed effect. The EPA guidelines (1979) suggest reducing the average daily dose by $(\text{Le/L})^3$ where Le is the actual experimental lifetime and L is the theoretical lifetime of the experimental animal. In treating the Neal and Rigdon data, this adjustment leads to a marked reduction $(\sim 1/40)$ in the effective daily dose level. These dose adjustments tend to ignore that detoxification processes and repair mechanisms generally reduce the effectiveness of low dose/long duration exposures.

The first set of data (Neal and Rigdon 1967) selected for extra-polation of possible risk is shown in Table 5-51 and indicates the incidence of forestomach tumors in CFW mice treated with BaP in their diet for an average of 110 days. The mice ranged in age from 17 to 116 days at the outset of the study.

To obtain a quantitative human risk, the human equivalent daily lifetime dose was computed as follows:

human dose (mg/kg/day)
$$=$$
 animal dose (mg/kg/day) $=$ $\frac{110 \text{ days}}{183 \text{ days}} \times \frac{183 \text{ days}}{90 \text{ wks } \times 7 \text{ d/wk}} \times \frac{2/3}{\text{animal weight}} \times \text{(animal kg weight)}$

That is,

1 mg/day human dose is approximately equivalent to $13.7 \, \text{mg/kg/day}$ animal dose.

The second set of data (Federenko and Yanysheva 1966) selected for extrapolation is shown in Table 5-52, and indicates the incidence of forestomach tumors in CC57 mice treated with BaP in triethylene glycol

TABLE 5-51. CARCINOGENIC RESPONSE IN CFW MICE TREATED WITH BENZO[a]PYRENE

Mouse Dose ^a (mg/kg/day)	Equivalent Human Dose (mg/day)	Incidence of Forestomach Tumors	Percent
0.0		0/289	0
0.13	0.0095	0/25	0
1.3	0.095	0/24	0
2.6	0.190	1/23	4
3.9	0.285	0/37	0
5.2	0.380	1/40	3
5.85	0.428	4/40	10
6.5	0.476	24/34	71
13.0	0.95	19/23	83
32.5	2.38	66/73	90

Source: Neal and Rigdon (1967).

^a Doses administered in diet for 110 days of a 183-day average study duration.

TABLE 5-52. CARCINOGENIC RESPONSE IN CC57 MICE TREATED WITH BENZO[a]PYRENE

Mouse Dose ^a (mg/kg/day)	Equivalent Human Dose (mg/day)	Incidence of Forestomach Tumors	Percent
1.0	2.68	23/27	85
0.1	0.268	23/30	77
0.01	0.0268	5/24	21
0.001	0.00268	2/26	8
0.0001	0.000268	0/16	0

Source: Fedorenko and Yanysheva (1966).

^a BaP in 0.2 ml triethylene glycol was administered by intubation one time per week for ten weeks of 83-week (19 months) study duration, beginning when mice were 2-3 months old.

solution by intubation once a week for 10 weeks. It is assumed that the control group response would be zero as in the previous study, since this was the response of the group treated with 0.1 μg , and the response of the group treated with 1 μg (8%) was also quite low. Thus, it is presumed that all forestomach tumors reported here are attributable to BaP ingestion.

To obtain the human equivalent lifetime daily dose, the following calculation was used:

human dose (mg/day) = animal dose (mg/mouse/wk) x $\frac{1 \text{ wk}}{7 \text{ days}}$ x $\frac{10 \text{ wks}}{83 \text{ wks}}$ x

$$\frac{83 \text{ wks}}{90 \text{ wks}}^{3} \times \frac{\text{human weight}}{\text{animal weight}}^{2/3}$$

That is, l mg/day (human dose) is approximately equivalent to .373 (mg/mouse/wk).

Estimation of Human Risk

The three dose/response models used to extrapolate human risk were the linear "one-hit" model, the multistage model and the log-probit model. The multistage model is actually a generalization of the one-hit model, in which the hazard rate function is taken to be a quadratic rather than a linear function of dose. The one-hit and multistage models assume that the excess probability (P) of a carcinogenic response to daily dose (D) is described by:

$$P = \frac{P(D) - P(O)}{1 - P(O)} = 1 - e^{-h(D)}$$

where h(D) is the "hazard rate" function, and P(D) and P(O) are the response rates at doses D and zero, respectively. For the one-hit model, the hazard rate function is simply BD, while in the multistage model, it is a quadratic function, $B_1D + B_2D^2$. The coefficients, B or B_1 and B_2 , of the hazard rate functions are solved by least squares regression of the data using linearized forms of the model equations.

For the "one-hit" model: $BD = log_e (1-P)$; for the multistage model, $B_1D + B_2D^2 = log_e (1-P)$.

The log-probit model assumes that human susceptibility varies with dose according to a log-normal distribution as follows:

 $P=\phi(z)$ where $z=A+\log_{10}D$ and ϕ is the cumulative normal distribution function. Values of z are obtained from cumulative normal probability tables relating response rate P such that $\phi(z)=P$. Once

z is determined, one can solve by linear regression for the "probit" intercept A in the linear equation:

$$z = A + \log_{10} D.$$

All of the data applicable to the models were utilized to calculate the parameters B, $B_1 + B_2$ or A. Although the EPA (USEPA 1979) has recommended that the data group which gives the largest value of B in the one-hit model should be used, this procedure was not followed here because of the relatively small number of animals per group. Table 5-53 presents the results of the mathematical calculations. Values of the parameters are given in the footnotes.

The observed and predicted responses at the equivalent human dose levels indicate the "goodness of fit". Within each data set, comparable fits are obtained by the three models. The extrapolated risk estimates based on the data of Neal and Rigdon (1967) and Fedorenko and Yanysheva (1966) are given in Tables 5-54 and 5-55, respectively. Due to different assumptions concerning the actual underlying mathematical relationship between dose and effect, the extrapolated risks are somewhat different at the relatively lower exposure levels typical of human environmental exposure. Regardless of the model, higher risk estimates (by about an order of magnitude) are predicted from the data of Fedorenko and Yanysheva (1966) due to the higher equation coefficients calculated for this study.

A possible cause for the difference in results for the two studies is thought to stem from the different modes of oral administration. In the study of Fedorenko and Yanysheva (1966), relatively higher doses were administered by intubation once a week, which probably resulted in higher exposure concentrations at the susceptible sites in the mouse forestomach. Alternatively, food may have diminished the bio-availability of BaP in the experiments of Neal and Rigdon (1967).

USEPA Risk Estimate

The U.S. EPA (1980) has established a zero ambient water concentration for the maximum protection of human health from potential carcinogenic effects of exposure to PAHs through ingestion of water and contaminated aquatic organisms. The water quality criterion is based on the assumption that each compound is as potent a carcinogen as BaP and that the carcinogenic effect of the compounds is proportional to the sum of their concentrations. Using a linear, non-threshold model, the EPA estimated, based on the findings of Neal and Rigdon (1967), that a concentration of 2.8 ng BaP/1 of surface water would result in an additional lifetime cancer risk of 10^{-6} .

TABLE 5-53. COMPARISON OF DOSE/EFFECT ANALYSES USING THREE MATHEMATICAL MODELS

	Response (%)					
Equivalent Human	Observed	Predicted				
Dose (mg/day)		One-hit	Multi-stage	log ₁₀ /probit		
Based on Neal and Rigdon (1967):						
.0095	0	0.9ª	1.6 ^b	1.4 ^c		
.095	0	8.8	14	11		
.19	4	17	26	18		
.285	0	24	36	23		
.38	3	31	44	27		
.428	10	34	48	29		
.476	71	37	52	31		
.95	83	60	74	41		
2.38	90	90	91	58		
Based on Federenko and Yanysheva (1966):						
2.68	85	100 ^d	86 ^e	92 ^f		
0.268	77	95	81	65		
0.0268	21	26	16	27		
0.00268	8	3	1.8	5.4		
0.000268	0	0.3	0.2	0.5		

a P = $1-e^{-.97D}$; B = 0.97b P = $1-e^{-(1.67D - .272D^2)}$; $B_1=1.67$; $B_2=-0.272$ c P = $\phi(-.182 + \log_{10} D)$; A = -.182d P = $1-e^{-11.5D}$; B = 11.5e P = $1-e^{-(6.78D-2.25D^2)}$; $B_1 = 6.78$; $B_2 = -2.25$ f P = $\phi(0.96 + \log_{10} D)$; A = 0.96

TABLE 5-54. PROBABLE UPPER BOUNDS ON EXPECTED EXCESS CANCERS PER MILLION POPULATION DUE TO BENZO[a]PYRENE INGESTION

(Based on Neal and Rigdon (1967) Study)

	Exposure Level (µg/day)					
	0.001	.01	0.1	1	10_	100
Linear Model	1	10	100	1,000	10,000	92,000
Log-Probit Model	10 ⁻³	10 ⁻¹	14	750	14,600	120,000
Multi-Stage Model	1.7	17	170 .	1,700	16,000	151,000

TABLE 5-55. PROBABLE UPPER BOUNDS ON EXPECTED EXCESS CANCERS PER MILLION POPULATION DUE TO BENZO[a]PYRENE INGESTION

(Based on Fedorenko and Yanysheva (1966) Study)

	Exposure Level (µg/day)					
	0.001	0.01	0.1	1	10	100
Linear Model	11	110	1,100	11,000	106,000	670,000
Log-Probit Model	0.3	26	1,200	21,000	150,000	480,000
Multi-Stage Model	7	70	680	6,800	65,000	480,000

Possible Shortcoming in BaP Risk Calculations Utilizing Less Than Lifetime Exposures

The range of possible excess cancers in humans ingesting BaP determined in the preceding section is based on an assumption that the total effective dose is equal to dose rate times dose duration. Additional information presented by Neal and Rigdon (1967) allowed an examination of the validity of this assumption in extrapolating short duration animal exposures to BaP to the human situation. As the following discussion indicates, it appears that in the case of the two available BaP studies, this assumption may not be valid.

Neal and Rigdon (1967) varied both the daily dose of BaP incorporated into the diet and the duration of feeding of the BaP-containing diet. Animals were returned to a basal, untreated diet after a variable period on the BaP-containing diet with most animals killed between 140 and 200 days from the start of the BaP exposure.

Their data, presented in Table 5-56, were analyzed by multiple regression analysis using linear transformations of the log/probit and one-hit models. In the log/probit model:

$$P(z) = \phi(B_1 + B_2 \log_e D^a t),$$

where: z =the value of $B_1 + B_2 \log_e D^a$ t at the response rate P(z), and

D = daily dose of BaP

t = number of days on BaP diet

 B_1 and B_2 = the probit intercept and slope, respectively.

The total effective dose is shown to be directly proportional to t but a power function (D^a) of daily dose. The exponent "a" should be close to 1 if total effective dose can be approximated by D times t.

The linear transformation of this equation is $z=B_1+(B_2 \text{ a log}_e \text{ D})+B_2 \log_e t$. The coefficients B_1 , B_2 a, and B_2 are obtained by least squares regression. From this analysis, "a" was determined to be 1.6 which is significantly greater than 1 (p <.01) suggesting that the total effective dose is not a simple product of D and t but is probably more accurately estimated by the power function, $D^{(1.6)}$ t. This result means that dose rate has a stronger influence on response than dose duration.

This relationship between dose rate and dose duration can also be demonstrated in the one-hit model:

$$P = 1 - e^{-(BD^a t)}$$

TABLE 5-56. COMPARISON OF DOSE/EFFECT ANALYSES WHERE EFFECTIVE DOSE IS A POWER FUNCTION

		Response			
Daily Dose (D)	Duration (t)	Observed	Predicted (1)	(2)	
0.10	110 (days)	0.017	0.10	0.07	
0.16	110	0.025	0.23	0.15	
0.18	110	0.10	0.27	0.19	
0.20	∿110	0.71	0.31	0.22	
0.4	~110	0.83	0.61	0.55	
1.0	~110	0.90	0.90	0.98	
1.0	2	0.11	0.07	0.07	
1.1	4	0.10	0.19	0.15	
1.0	5	0.44	0.20	0.16	
0.9	7	0.30	0.23	0.19	
0.40	30	0.67	0.27	0.20	
12.0	1	0.50	0.79	0.91	

Multiple Regression Results:

(2) one-hit model: $P = 1-e^{-(B D^a t)}$, B = .035 a = 1.71

Source of data for columns 1-3: Neal and Rigdon (1967).

⁽¹⁾ log/probit model: $P = \phi(B_1 + B_2 a \log_e D + B_2 \log_e^t)$, $B_1 = -1.957$; $B_2 a = 1.113$; $B_2 = 0.692$, a = 1.113/0.692 = 1.61

where D, t and "a" are defined as above and "a" should again be close to one if D times t is a good approximation of total effective dose. The linear transformation is:

$$\log_{e} [-\log_{e} (1-P)] - \log_{e} t = \log_{e} B + a \log_{e} D.$$

Solving for the coefficients by multiple regression "a" equals 1.71 and B equals 0.035. Thus, the effective dose is more appropriately given by: $D^{(1.7)}$ t. Again, the exponent of D is significantly greater than one $(p \sim .01)$.

Thus, both analyses indicate that in the case of BaP, dose rate is more important than dose duration in determining the incidence of BaP-induced forestomach tumors in mice. These results appear to contradict the validity of the assumptions inherent in the established guidelines (USEPA 1979) for computing risk to humans from ingestion of BaP, and suggest that in addition to the possible increased sensitivity of the rodent forestomach to tumor induction by BaP, extrapolation of high exposure/short duration animal studies to determine excess lifetime human risk may overestimate true risk.

Other Risk Considerations and Overview

Although the two animal studies discussed in the previous section demonstrate a carcinogenic response to BaP in mice by the ingestion route, the quantitative estimates of human risk must be treated with considerable caution since they are dependent on a number of stated assumptions that have high levels of uncertainty. Several considerations which seriously undermine the validity of quantitative human risk extrapolations for BaP have been discussed. For example, the mouse forestomach has a different anatomy than the human stomach and may be particularly sensitive to BaP. This difference in anatomy is indicative of the fundamental problem of species to species extrapolation. differences in dose-effect relationship were obtained between the Neal and Rigdon (1967) and the Fedorenko and Yanysheva (1966) studies which were primarily attributed to the difference in mode of administration but which may also have been caused by differences in dosing schedules. Analysis of additional data from the Neal and Rigdon (1967) study indicates that the total effective dose is probably not a simple product of daily dose times duration but, in fact, may be a more complex power function in which the dose rate has a greater influence on response than treatment duration.

Another vital consideration in the extrapolation of carcinogenic risks associated with PAH exposure is the metabolic differences, both qualitatively and quantitatively, in the inducibility of the P-450 dependent microsomal mixed function oxidase system which is involved in the activation of tumorigenic PAHs to their ultimate carcinogenic forms. Considerable variations are known to exist among various species and significant individual variations have been documented for humans.

Among the other PAH compounds in this group, benz[a]anthracene, dibenz[a,h] anthracene, benzo[b]fluoranthene, and indeno[1,2,3-c,d] pyrene are complete carcinogens for mouse skin; benzo[g,h,i]perylene is a co-carcinogen with BaP; and benz[a]anthracene, benzo[b] fluoranthene, chrysene and indeno[1,2,3-c,d]pyrene are all initiators of skin carcinogenesis in mice. Thus, for the most part, published reports on the other PAHs in this group deal with localized tumor production and are generally not considered relevant to the assessment of systemic carcinogenicity. We have, therefore, focused our attention on the risks associated with BaP because (1) it is the dominant substance in this group with respect to human exposure (see Section 5.3.2); and (2) reduction of BaP levels will presumably lead to reduction in exposure to the other PAHs in this group. In addition to ingestion studies, BaP has been shown to be animal carcinogen by dermal and intratracheal routes. It is also a transplacental carcinogen, an initiator of skin carcinogenesis in mice, and is carcinogenic in single dose experiments.

Benzo[a]pyrene is also the most active mutagen of the group, inducing in vivo chromosomal aberrations in both hamster spermatogonia and bone marrow cells and inducing positive mutagenic responses in sister chromatid exchange tests. Mixed mutagenic responses have been reported for the other PAHs in this group. Because of its mutagenicity, BaP exposure could also be expected to contribute to the genetic burden of a population, but since extrapolation procedures for genetic risks have not been well established, a quantitative risk assessment for these kinds of health hazards is not presently feasible.

BaP appears to exert little effect on the developing embryo; data were unavailable for the other compounds in this group regarding adverse reproductive effects.

The major focus of studies conducted with PAHs in this group has been their potential for inducing carcinogenic effects. Thus, little information is available on other possible toxic effects associated with exposure to these compounds. However, in assessing the risks to humans associated with exposure to these PAHs, one should not overlook possible augmentation of effects through synergistic or co-carcinogenic mechanisms. Current understanding of the co-carcinogenesis process, however, is not sufficiently adequate to allow estimation of human risk at this time.

5.3.2 Human Exposure

5.3.2.1 <u>Introduction</u>

As is apparent from the preceding sections, monitoring data are limited for the PAHs comprising the BaP group. This section examines human exposure to benzo[a]pyrene and the other chemicals in this group via ingestion (food and drinking water), inhalation, and dermal contact.

5.3.2.2 Ingestion

Drinking Water

Basu and Saxena (1978) sampled ten finished water supplies for several PAHs, including BaP, benzo[g,h,i]perylene, indeno[1,2,3-c,d]-pyrene, and benzo[k]fluoranthene. All of these chemicals were detected in at least seven of the 10 samples (see Table 5-38). Most values range from 0.2 ng/1-1 ng/1, and these as well as maximum reported values are shown in Table 5-57. Removal efficiencies ranged from 77-100% for plants using activated carbon; thus in plants of this type, concentrations of BaP and related compounds would be expected to be low.

No data on drinking water levels were found for the other chemicals in this group (chrysene, benz[a]anthracene, acenaphthylene, benzo[b]-fluoranthene, and dibenz[a,h]anthracene). In addition, these chemicals are rarely detected in ambient water (see Section 5.2.2). Table 5-57 shows estimated exposures for these other compounds based upon the limited data available.

Food

Levels of the PAHs in raw foods appear to be generally low. However, vegetables or fruits grown in the vicinity of air releases may contain higher levels (U.S. EPA 1980). The highest levels of the BaP group PAHs in food, however, appear to result from the cooking process, especially charcoal broiling and smoking. Table 5-58 estimates daily exposure by consumption of such foods, and lists the various assumptions made in deriving the estimates. The data on contamination levels were taken from the U.S. EPA Criteria Document (U.S. EPA 1980) and White and Vanderslice (1980). It is apparent that data are limited for many of these chemicals. As a result, the columns in Table 5-58 were not totaled, since the exposure estimates shown represent only an unknown portion of the actual total exposure. In addition, the maximum intakes represent a level of maximum contamination and consumption of the food item. the total is meant to represent a worst case and is not representative of widespread exposure. Charcoal broiling represents the major source of exposure to the BaP group PAHs via food. Consumption of large amounts of food cooked in this manner could result in a high exposure, perhaps up to about 6 µg/day (intake from charcoal-broiled beef and smoked pork).

TABLE 5-57. ESTIMATED HUMAN EXPOSURE TO THE BENZO[a]PYRENE GROUP PAHS VIA DRINKING WATER

	Concentrati	on (ng/1) ^a	Estimate Exposure	ed Daily e (ng/day)
	Typical	Maximum	Typical	Maximum
Benzo[a]pyrene	0.3	1.6	0.6	3.2
Acenaphthylene	NAC	NA		
Benz[a]anthracene	NA	NA		
Benzo[b]fluoranthene	NA	NA		
Benzo[k]fluoranthene	0.3	0.7	0.6	1.4
Benzo[g,h,i]perylene	2.0	4.0	4.0	8.0
Chrysene	NA	NA		
Dibenz[a,h]anthracene	NA	NA		
<pre>Indeno[1,2,3-c,d]pyrene</pre>	1	1.7	2	3.4

Source: Basa and Saxena, 1978.

^aFor locations see Table 5-38. Typical concentrations are average values, the data on BaP from New Orleans was excluded from the average.

b Intake assumed to be 2 liters of water per day (ICRP 1975).

^cNA = Not Available.

TABLE 5-58. LEVELS OF BENZO[a]PYRENE GROUP PAIRS IN FOOD AND ESTIMATED EXPOSURE VIA INGESTION OF FOOD

Charcoal broile beel ^d	đ	Consum (g/da Typic <u>al</u>	iy)	Contami Lug/	nation kg)	(pg/	ake da <u>y)</u> Max.	Bo Contamir (0g/k Typica)	ation g)	<u>inthrace</u> n Int Int (ng/ Typical	ake day)	Contani	nation kg)	theryler Inta (ng/c Typical	ike Liy)	(րջ	rnar ron' /kg)		iake /day) Max.
tlamburge r		10	NA	NA	2.6	NA	0.03	NΛ	3	NΛ	.01	1	15	0.01	0.15	NA	1	NA	0.01
Steak		1	86	5	50	0. 02	4. 1	3	31	0.009	2.7	6	12	0.02	1.0	1	9	.003	0.8
Smoked pork ^b		1	27	2	55	0.002	1.5	3	33	0.003	0.9	2	25	0.002	0.7	NA	2.2	NΛ	0.06
Swoked saurage ^c		1.5	30	NA	4	0.006	0.12	0.2	2	0.0003	0.06	NΛ	3.0	0.005	0.09	NA	NA	NA	NΑ
Smoked fish ^d		0.1	14	1	37	0.0001	0.5	1	189	. 000 t	2.6	1	2.4	0.0001	0.03	1	173	0.0001	2.4
Ort		18	NΛ	ì	8	0.02	0.14	ì	30	0.02	0.5	1	4.0	0.02	0.07	NΛ	12	NA	0.2
Fruits		205	NA	0.02	6	0.004	1.2	NA	NA	NA	NΛ	NA	NA	NA	NA	NA	NA	HA	NA
Grains		256	NA	NA	0.3	NA	0.08	NA	NA	NA	N.A	HA	NA	NΛ	NA	NA	NA	NA	NA
Vegetables	lotal	248	NA	0.01	0.1	0.002	0.2	NA	NA	NA	NA	N.V	NA	NA	NA	NΛ	NΛ	NI	NA
	l.eat y	40	NA	NA	7.5	NA	0.3	NA	50	NA	NA	NA	B	NA	0.3	NA	80	nA	3.2

^{*}Consumption of beef = 86 g/day, 15% charcoal-broiled = 80% hamburger, 20% steak. Worst case maximum 86 g consumption of charcoal-broiled steak.

Note: Typical consumption refers to average consumption for males 23-34; typical concentrations are not actual arithmetic averages due to the widely varied nature of the studies, but are meant to approximate such an average.

Source: USDA (1978, 1980), U.S. EPA (1980), White and Vanderstice (1980).

 $^{^{\}rm b}$ Consumption of pork - 27 g/day, 5% smoked. Worst case maximum, $^{\rm o}$ / g/day smoked.

Consumption of sausage - 30 g/day, 5% smoked. Worst case maximum 30 g/day smoked.

densumption of fish - 14 g/day, 12 smoked. Worst case maximum, 14 g/day smoked.

5.3.2.3 Inhalation

The U.S. EPA (1980) has summarized the available data for PAH levels in air, mostly for urban areas. These data, although limited, were utilized in developing exposure estimates for an assumed respiratory flow of 20 $\rm m^3/day$ (ICRP 1975). These results are shown in Table 5-59. As was found for food and drinking water, BaP generally appears to represent the largest exposure of chemicals in this group.

As discussed in Section 5.2.4, Moschandreas et al. (1980) examined BaP concentrations indoors. These authors found concentrations indoors to be similar to those outdoors on non-woodburning days, but higher indoor concentrations on wood-burning days. The mean level indoors on woodburning days, however, was 4.7 ng/m^3 , which is within the range for urban and rural shown in Table 5-59.

In addition to ambient air, smoking can contribute to inhalation of PAHs. Schmeltz et al. (1975) reported that mainstream smoke contained 0.025 ug BaP and 0.044 µg benz[a]anthracene per cigarette. A sidestream:mainstream ratio of 3.4 for BaP was also reported, resulting in a sidestream smoke content of 0.085 µg BaP per cigarette. Thus, exposure of smokers to BaP in mainstream smoke could range from .025-2.5 µg/day, depending upon the type of cigarette smoked, the amount inhaled, and the number of cigarettes smoked, assuming a range of consumption of 1-100 cigarettes per day (U.S. DHEW 1979). An estimated 33.2% of adults over 17 years old smoke cigarettes or 54.1 million persons in the United States. Of smokers, 25-30% smoke more than 25 cigarettes per day (U.S. DHEW 1979); thus, a large segment of the population could be exposed to BaP at levels greater than 0.6 µg/day.

In addition, nonsmokers may be exposed to BaP through inhalation of sidestream smoke. Although only a few measurements of BaP have been taken in smoke-filled rooms, concentrations may be estimated from measurements of CO levels, which have been summarized by Burns (1975). The results are not consistent, and apparently depend upon a number of variables. They show levels of $44-92 \text{ mg/m}^3$ CO in rooms $(38-92 \text{ m}^3)$ where 30-80 cigarettes had been smoked with no ventilation. The Surgeon General's Report (U.S. DHEW 1979) reported levels up to 50 mg CO produced in sidestream smoke per cigarette, as compared with the 0.085 ug BaP produced per cigarette. Thus, by analogy, a room concentration of 0.08-0.2 $\mu g/m^3$ can be calculated. Alternatively, using 0.085 μg BaP/cigarette, and assuming a room size of 48 m^3 with no ventilation in which 40cigarettes were smoked, a concentration of about 0.07 $\mu g/m^3$ can be calculated. A nonsmoker exposed to such a situation 2 hours/day would receive about 0.25-0.7 µg/day, assuming a respiratory flow of 1.8 m³/hr. These values would tend to overestimate exposure since no ventilation was assumed, and a high respiratory flow was used in the calculation. Smokers in the same situation would receive the combined intake from sidestream and mainstream smoke.

TABLE 5-59. ESTIMATED EXPOSURE TO BENZO[a]PYRENE GROUP PAHS VIA INHALATION OF AMBIENT AIR

	Ambient Concent:	<u>Intake (r</u> Urban		
Benzo[a]pyrene	1-100	Rura1 0.01-10	20-2000	Rural 0.2-200
Benz[a]anthracene	0.1-20		2-400	
Benzo[b]fluoranthene	0.1 -10		2-200	
Benzo[k]fluoranthene	0.1-10		2-200	
Benzo[g,h,i]perylene	0.2 -50	7-50 ^c	4-1000	140-1000
Chrysene	0.2 -10		4-200	
Indeno[1,2,3-c,d]pyrene	0.3 -1.3		6-30	

^aU.S. EPA (1980), White and Vanderslice, 1980.

b_{Based} on respiratory flow of 20 m³/day (ICRP 1975).

^cThere are not enough monitoring data to explain this apparent lack of difference between urban and rural areas.

There are problems with using CO levels to estimate BaP levels in a smoke-filled room since they would be due to sidestream smoke and exhaled mainstream smoke. In addition, the concentration of CO and BaP would be influenced by type and amount of tobacco smoked, extent of inhalation, size of room, ventilation, and duration of exposure. The estimates provided above are probably on the high side since worst-case assumptions were frequently made.

The U.S. EPA (1978) estimated the sizes of populations exposed to various levels of BaP in the United States using SRI data for areas with coke ovens and ambient data or national average concentrations for other areas. Their results are shown in Table 5-60.

5.3.2.4 Dermal Contact

No direct information is available regarding the dermal exposure of humans to this group of PAHs. However, due to the low levels found in drinking water, dermal exposures are expected to be low.

5.3.2.5 Overview

Estimated typical daily exposures to the BaP group PAHs are shown in Table 5-61. Data are unavailable for estimating exposures to some of the compounds in the group. Smoking appears to be the most significant exposure route for benzo[a]pyrene and benz[a]anthracene. A smoker consuming 25 cigarettes per day could be inhaling 600 ng and 1100 ng of these compounds, respectively.

Food, primarily charcoal-broiled and smoked meats and fish, is also a significant exposure medium for BaP, benzo[g,h,i]perylene, and benz[a]anthracene, although inhalation exposures at the upper limit of the concentration ranges reported for urban areas are greater in every case. By contrast, although monitoring data are very limited, drinking water appears to contribute relatively small amounts to typical daily exposures.

Data concerning levels of these compounds in foods are variable, as are atmospheric concentrations reported for urban areas. Therefore, some subpopulations (i.e., those consuming large amounts of charcoal-broiled and smoked meat) may be exposed to considerably higher levels (possibly as high as 6 $\mu g/day$ from food consumption alone).

TABLE 5-60. ESTIMATED SIZE OF THE U.S. POPULATION EXPOSED TO RANGES OF BENZO[a] PYRENE CONCENTRATIONS IN AMBIENT AIR

Population (1000's) exposed to BaP Concentration (ng/m^3)

>5.0	1.0-5.0	0.5-1.0	< 0.5
1059	102,132	26,731	73,294

NOTE: For some locations for which monitoring data were unavailable (representing about 50% of the population) the upper limits of 95% confidence intervals of national average concentrations were used. These levels were as follows:

urban SMSA
$$1.3 \text{ ng/m}^3$$

urban non-SMSA 1.4 ng/m^3
rural 0.23 ng/m^3

Source: U.S. EPA (1978).

TABLE 5-61. ESTIMATED HUMAN EXPOSURE TO THE BENZOLA PYRENE CROUP PAILS

				Typical ^a	Exposure (ng/day	y)			A Politika da de
Exposure Route	BaP	Acenaphthylene	Benz[a]- anthracene	Benzo[b]= fluoranthene	Benzo[k]- (luoranthene	Benzolg,h,il- perylene	Chrysene	Dibenz[a,h]- _anthracene	Indeno[1,2,3-c,4]-
Ingestion									
Drinking Water	0.6	0.6	HA	NA	NA	0.4	NA)	NA	2
Food	50	NA	30c	АИ	NA	60¢	3°	NA	NA
Inhalation									
Urban	10-60	11A	4-90	2 - 30	4 100	4-180	4-130	NA	6 - 30
Rural	2								
Smoking	600		1100						

antypical" as used here is a qualitative estimate based on average consumption and a range of reported concentrations.

Source: For assumptions and references, see text.

bNA = not available.

every limited data are available concerning the dietary intake of these compounds. Thus, this only represents a partial estimate.

5.4 EFFECTS AND EXPOSURE--NON-HUMAN BIOTA

5.4.1 Effects on Non-Human Biota

5.4.1.1 Introduction

No data from acute toxicity bioassays were available for benzo[a]-pyrene or any other PAHs in this group, but several studies have been done that indicate the general concentration ranges that cause toxic effects in marine and freshwater organisms.

5.4.1.2 Plants and Microorganisms

In studies on the effects of PAHs on Escherichia coli, BaP was found to be highly toxic when in colloidal suspension in the presence of light and oxygen. The authors suggest that the toxicity of BaP was probably due to its photo-oxidation products. The concentrations of BaP at which toxicity occurred, however, were not reported (Harrison and Raabe 1967).

Haas and Applegate (1975) studied the effect of PAHs at concentrations of 10^{-5} , 10^{-6} , and 10^{-7} M on E. coli; (approximate calculated concentration range, $25-2500~\mu g/1~BaP$). Benz[a]anthracene, dibenz[a,h]anthracene, and benzo[a]pyrene all promoted growth in bacteria at these concentrations.

Similar growth-stimulating effects of PAHs have been shown with freshwater algae. Graff and Nowak (1966) reported that concentrations of 10-20 µg/l of BaP, benz[a]anthracene, benzo[b]fluoranthene, indeno[1,2,3-c,d]pyrene, and benzo[g,h,i]perylene promoted growth in the algae Chlorella vulgaris, Scenedismus obliquus, and Ankistrodesmus oraundii.

5.4.1.3 Animals

No standard toxicity tests were found for the compounds in this group. In the only study examining mortality, benz[a]anthracene was found to cause 87% mortality to the freshwater fish, bluegill (Lepomis macrochirus) at 1.0 μ g/l in 6 months (Brown et al. 1975).

Various other effects have been reported in the literature. For example, Haranghy (1956) reported that concentrations of 0.2-1.0 $\mu g/1$ BaP injected into the visceral sac of freshwater mussels resulted in a significant decrease in filtration rate. The author suggested that BaP inhibited ciliary activity or possibly the entire metabolism of the mussels. Exposure of marine calcarious sponges, Leucosolenia complicata and L. variabilis, to 5 g/l BaP caused tissue damage and abnormal growth (Korotkova and Tokin 1968).

Several studies on aquatic invertebrates and lower vertebrates, including sponges, newts. and toads, have shown that PAHs in this group can produce cancer-like growths and cause teratogenetic and

mutagenic effects. The number of species examined is small; however, though no exact concentration levels for these effects were given, it is believed that they were caused by concentrations higher than those generally found in the environment (Neff 1979).

Flat fish (English sole, Parophrys vetulus) exposed to hydrocarbon-contaminated sediments near the industrialized City of Seattle were found to have a significant number (32% of the adult population) of liver tumors (Malins 1979). It appeared that these fish had absorbed PAHs and PCBs from the sediment. Fish of the same species taken from relatively "clean" areas of the Washington coast were almost free of this disease. PAHs specifically were not detected in liver samples of these fish, but the authors theorized that these compounds had been absorbed to some degree and metabolized. It is known that metabolites (e.g., diol-epoxides) of certain higher-molecular-weight aromatic hydrocarbons are carcinogenic, but these compounds were not identified in liver samples in this study (Malins 1979).

In related work, Malins (1979) showed that liver enzymes of English sole extensively metabolize benzo[a]pyrene to intermediates that interact with DNA. Such interactions are thought to be a starting point for tumor formation. Studies with freshwater trout (Salmo trutta lacustris) have shown that trout liver microsomes actively biotransformed benzo[a]pyrene into a variety of electrophilic metabolites and catalyzed binding of activated BaP to DNA. Similar studies showed that liver enzymes from coho salmon (Onchorhynchus kisutch) and starry flounder (Platichthys stellatus) extensively metabolized BaP into reactive intermediates that bind to DNA (Varnasi et al. 1979). Although no data on BaP concentrations found in the coastal Washington sediments were available, it is strongly felt (Malins 1979) that the liver disease seen in the benthic English sole was associated with contaminated sediments.

5.4.1.4 Conclusions

Since the toxicity data for the BaP group PAHs are very limited, it is not possible to make any conclusions as to the toxicity of BaP or any other PAHs. The toxicity data that are available are summarized below:

Concentration (µg/1)	Compound	Effect
1.0	(benz[a]anthracene)	Caused 87% mortality in Bluegill in 6 months.
10-20	(BaP, benz[a]anthracene, benzo[b]-fluoranthene, indeno[1,2,3-c,d]-pyrene, benzo[g,h,i]perylene)	Promoted growth in freshwater algae.
25-2500	(benz[a]anthracene, dibenz[a,h]-anthracene, and BaP)	Promoted growth in bacteria.

5.4.2 Exposure of Non-human Biota

5.4.2.1 Introduction

PAHs are found in various aquatic and terrestrial environments. From monitoring data, it appears that higher concentrations are generally found near industrialized areas, major ports, and areas of petroleum or other fossil-fuel related activities. In some remote locations, such as the coast of Greenland and areas of peat deposits, concentrations of PAHs, particularly BaP, may be higher than general background levels in "pristine" areas. The source of these high concentrations is believed to be biosynthesis of PAHs from naturally-occurring quinones in anaerobic sediments, and leaching of BaP from peat deposits, which are known to contain high concentrations of PAHs (Neff 1979).

This section will examine the exposure of biota to PAHs, considering both the levels and the types of locations where it may occur.

5.4.2.2 Monitoring Data

The STORET monitoring data provide a basis for examining aquatic exposure. Summary descriptions and concentration distributions for each PAH in the BaP group are given in Section 5.2.2. Observations of values greater than the detection limit (unremarked data) are infrequent, with only 17 such observations of a total of 3,261 observations in water for this group. A greater percentage (14%) of the sediment values were unremarked; 83 of 600 total observations. Unremarked observations for both sediment and surface water are presented in Table 5-62.

Overall there were three observations of the PAHs in the BaP group greater than 1000 $\mu g/kg$ in sediment.

Many of the higher sediment values occurred consistently in several locations, including Puget Sound, coast of Washington State and Oregon, San Francisco Bay and north coastal California, a hazardous waste site in North Carolina, and the Houston ship channel.

5.4.2.3 Aquatic Fate

Environmental conditions have a significant influence on the disposition of BaP and related PAHs in aquatic systems and, hence, on the exposure conditions. These compounds are quite insoluble in water and chemical transformation processes; e.g., photolysis, are not as significant as with the lower-molecular-weight PAHs. EXAMS data for BaP indicate that in five of the six environments examined, nearly all ($\sim 98\%$) of the compound is accumulated in the sediments. These observations are confirmed by monitoring data that indicate a propensity of these PAHs to accumulate in the sediments. In the oligotrophic lake, a greater percentage (18.75%) of BaP remained in the water column, but

TABLE 5-62. SUMMARY OF UNREMARKED OBSERVATIONS OF CONCENTRATIONS OF BENZO[a]PYRENE GROUP PAHS IN SURFACE WATER AND SEDIMENT--STORET, 1980

Compound	Number of Observations	Range of Observations
	Surface Water (µg/1)	
Acenaphthylene	6	0.01-0.12
Benz[a]anthracene	5	1-400
Benzo[k]fluoranthene	5	320-1500
Chrysene	1	0.02
	Sediment (µg/kg dry weight)	
Benzo[a]pyrene	11	0.02-1400
Acenaphthylene	12	0.002-93
Benz[a]anthracene	12	6.2-340
Benzo[k]fluoranthene	12	0.9-1300
Benzo[g,h,i]perylene	5	4.3-40.9
Chrysene	11	0.06-120
Dibenz[a,h]anthracene	2	15.7-2600

Source: See Table 5-34.

the remainder still went to the sediments. BaP in sediment is less likely to be physically transported than BaP in the water column. It is likely to persist chemically unchanged because the conditions conducive to chemical degradation (light, oxygenation) are not present. In addition, microbial biodegradation is not a significant fate process for any of the BaP group compounds. These characteristics suggest that although monitoring data for sediment are limited, this medium may present an important exposure route. Therefore, uptake from sediment was examined.

5.4.2.4 Factors Affecting Bioavailability of Sediment Concentrations

It is known that aquatic biota are exposed to PAHs directly from water, but the extent to which they may be exposed to sediment concentrations of these compounds is less well understood.

Accumulation of hydrocarbons, including PAHs from oil-contaminated sediments by the English sole Parophrys vetulus has been investigated (McCain et al. 1978). Results from several separate studies (Roesjadi et al. 1978, Fucik et al. 1977, Anderson et al. 1974) indicate that overall uptake of PAHs is not significant, but is greater by suspension feeding benthic invertebrates than deposit feeders, and that accumulation by fish is greater from water than from sediment. The conclusion has been made that any PAH that is taken up comes, to a greater extent, from dissolved PAH in interstitial waters and from PAH in the water column (dissolved and/or adsorbed onto suspended solids) rather than desorbed from the sediment itself (Neff 1979).

These studies also indicated that sediment adsorbed PAHs are not readily metabolized by benthic invertebrates. However, based on the few species studied, it is impossible to quantify the degree to which accumulation of PAHs in sediment represents a major source of exposure to aquatic biota.

5.4.2.5 Conclusions

Scattered observations for the PAHs in this group reveal that they are detected infrequently in water and when found are in the low $\mu g/l$ range. Unremarked observations indicate that these PAHs may be found to a greater extent and in higher concentrations in sediment. The possible contribution of naturally-occurring PAHs to these levels is unknown. Sediment concentrations do appear to be consistently higher near industrialized areas, however, and levels as high as 2600 $\mu g/kg$ have been detected.

Environmental fate and monitoring data indicate that BaP group PAHs may accumulate to potentially high levels in the sediments. The extent to which these compounds are available to biota from the sediments is not well understood or quantifiable at this time, but experimental evidence indicates that although some PAHs are taken up by benthic organisms directly from sediment, the PAHs present in interstitial waters and adsorbed to suspended particulates are more available for uptake by biota.

5.5 RISK CONSIDERATIONS

5.5.1 Introduction

An objective of the risk assessment process is the quantification of risks to various subpopulation groups of humans and other classes of biota. Quantifying such risks requires:

- Careful identification of the subpopulations at risk and the populations exposed;
- Evaluation of the ranges of exposure for each subpopulation;
- Determination of the effects levels or dose-response data in the species of concern and/or proxies for these species (for example, laboratory animals as a proxy for humans);
- Extrapolation of dose/response data to the subpopulations at risk.

The largest data base available for estimating the risks associated with human exposure to PAH compounds in the benzo[a]pyrene group is that available on BaP. Dose-response data for BaP-induced forestomach tumors in mice and information on the range and types of human exposure to BaP are sufficient to allow estimation of the potential risks associated with continuous lifetime exposure to some level of BaP. Information on exposure and effects for the other PAHs in this group is lacking and no quantitative estimates of risks associated with exposure to these compounds is possible at this time.

Extensive uncertainties are inherent in the extrapolation of carcinogenic findings in laboratory animals to humans. Additionally, because of an inadequate understanding of the mechanisms of carcinogenesis, a scientific basis has not been developed for selecting among several alternative dose-response models, which yield widely differing results. Therefore, we have applied three dose-response models to the data, and the results provide a range of potential human risk associated with exposure to BaP.

In the case of risks to other forms of biota, insufficient data are available on most toxic effects and on exposure levels to assess risk quantitatively.

5.5.2 Human Exposure

A series of possible exposure routes for humans with an indication of the size of the population at risk and typical exposure levels are presented in Table 5-61 and also in Table 5-63 with the estimated population sizes. Exposure levels of BaP appear to be the largest of the chemicals in the group via drinking water, food, and air (see Section 5.3.2), but this may be because monitoring data are limited for the other compounds. Smoking appears to be the most significant exposure route for benzo[a]-pyrene and benz[a]anthracene. A smoker consuming 25 cigarettes per day could inhale 600 ng and 1100 ng of these compounds, respectively.

Food, primarily charcoal-broiled and smoked meats and fish, is also a significant exposure medium for BaP, benzo[g,h,i]perylene, and benz[a]-anthracene, although inhalation exposures at the upper limit of the concentration ranges reported in urban settings are greater in every case. By contrast, although monitoring data are very limited, drinking water appears to contribute relatively small amounts to typical daily exposures.

Data concerning levels of these compounds in foods are variable, as are atmospheric concentrations reported for urban areas. Therefore some subpopulations (i.e., those consuming large amounts of charcoal-broiled and smoked meat) may be exposed to considerably higher levels (possibly as high as 6 $\mu g/day$ from food consumption alone).

5.5.3 Human Risk

5.5.3.1 Carcinogenicity

Several PAHs in the benzo[a]pyrene group are well established animal carcinogens, co-carcinogens and/or tumor initiators; others have not been demonstrated to induce tumorigenic responses (see Section 5.3.1.3). The capacity of individual PAHs to induce positive responses in humans is not so well established. This is primarily due to the fact that human exposures have not been to individual chemicals but rather to combinations as they occur in coke oven emissions, coal-tar, soot or from environmental exposures to tobacco smoke or exhaust fumes. Numerous studies have shown increased incidences of lung, skin and other types of cancer among workers exposed to coke oven emissions, coal gas, coal tar and pitch (IARC 1972). These studies, however, do not allow identification of the individual chemical(s) responsible, do not account for possible synergistic or co-carcinogenic effects resulting from other components, often are unable to clearly define exposure levels and generally are not amenable to quantifying human risk.

For the most part, available data for specific PAH compounds are from skin painting and subcutaneous or intramuscular injection experiments in mice. Few oral or inhalation experiments have been conducted, and those that are available are generally inadequate for risk assessment purposes.

TABLE 5-63. ESTIMATED HUMAN EXPOSURE TO THE BENZO(a) PYRENE GROUP PARS

			Typical Exposure (pg/day)								
Exposure Route	Population ^d Size	Benzo[a] pyrene	Acenapht by Lene	Benz[a]- anthracene	Benzo[b]+ fluoranthene	Benzo[k]- ! luoranthene	Benzo[g,h,i]- perylene	Chrysene	Dibenz[a,h] - anthracene	Indeno 11,2,3-c,d1- pyrene	
Ingest ion											
Drinking Water	220.6 x 10 ⁶	0.0006	0.0006	NA '	NΛ	NA	0,0004	NA	NΛ	0.002	
Food	220.6×10^6	0.05	NA	0.03^{d}	NA	NA	0.064	0.003 ^d	NA	ħΑ	
Inhalat ion											
Urban	165.6 x 10 ⁶	0.02 - 2	NA	0.002 - 0.4	0.002 - 0.2	0.002 - 0.2	0.004 - 1	0.004 - 0.2	NΛ	0.006 - 0.03	
Rural	55.1 x 10 ⁶	0.0002 - 0.2					0.14 - 1				
Smoking	54.1 x 10 ⁶	0.6		1.1							

[&]quot;U.S. Department of Commerce (1980).

SOURCE: For assumptions and references, see Section 5.3.2.

bully bical" as used here is a qualitative estimate based on average consumption and a range of reported concentrations.

^{&#}x27;NΛ = not available.

 $^{^{}m d}$ Very limited data are available concerning the dietary intake of these compounds. Thus, this only represents a partial estimate.

Benzo[a]pyrene has been most extensively studied of all PAHs in the benzo[a]pyrene group. BaP has been shown to be both a local and systemic carcinogen by oral, dermal and intratracheal routes in animals. It is also a transplacental carcinogen and an initiator of skin carcinogenesis in mice. Few studies have adequately examined the carcinogenic effects of orally administered BaP but two ingestion studies whose data appear to lend themselves to dose/response extrapolation, one by Neal and Rigdon (1967) on CFW mice, the other by Fedorenko and Yanysheva (1966) on CC57 mice, resulted in the production of forestomach tumors in this species. Dose-response values based on these two studies were estimated using three extrapolation models (see Section 5.3.1.4). A summary of the risk estimates obtained for these two data sets from the various models is presented in Table 5-64. Risk estimates are shown for continuous lifetime daily exposures ranging from one nanogram to 100 micrograms. The gap between the estimates is large in the low-dose region. However, present scientific methods do not permit a more accurate or definitive assessment of human risk.

The interpretation of these calculated risk estimates is subject to a number of important qualifications and assumptions which were discussed in Sections 5.3.1.3 and 5.3.1.4:

- The site of tumor production and probable underlying mechanisms are possibly unique to laboratory rodents. The rodent forestomach has a different anatomy than the human stomach and may represent an unusually sensitive assay of the carcinogenicity of BaP by the oral route. The absence of tumors at sites other than in the forestomach raises questions as to the significance of these findings to the risks to humans from ingestion of BaP.
- Assuming that the positive findings indeed provide a basis for extrapolation to humans, although dosing route may be significant, we have assumed that absorbed dose by inhalation or ingestion has the same effect.
- A vital consideration in the extrapolation of risks associated with PAHs in general, and BaP specifically, is the existence of metabolic differences, both qualitatively and quantitatively, between humans and other species.

The degree of individual genetic variation of some activating enzymes is suspected to be a key factor in an organism's susceptibility to PAH-induced carcinogenesis. The highly heterogeneous nature of human populations exposed to BaP introduces a confounding factor in reliably predicting excess cancer incidences due to BaP exposure.

 Both animal studies, and especially the study by Neal and Rigdon (1967), were of less than lifetime duration. Doses were administered for about half the duration of the experiment in both studies. In order to adjust animal dose

TABLE 5-64. ESTIMATED LIFETIME EXCESS PROBABILITY OF CANCER TO HUMANS DUE TO BENZO[a]PYRENE INCESTION AT VARIOUS EXPOSURE LEVELS BASED ON THREE EXTRAPOLATION MODELS^a

Exposure Level (µg/day	Estimated	Lifetime Excess	Probability of	Cancer at Indic	ated Exposure L	evel
napostire never (hg/day	0.001	0.01	0.1	1	10	100
Extrapolation Model Linear Model						
Neal and Rigdon data	1×10^{-6}	1 x 10 ⁻⁵	1×10^{-4}	1×10^{-3}	1×10^{-2}	9.2×10^{-2}
Fedorenko and Yanysheva data	1.1 x 10 ⁻⁵	1.1×10^{-4}	1.1×10^{-3}	1.1×10^{-2}	1.1 x 10 ⁻¹	6.7×10^{-1}
Log-probit Model Neal and Rigdon data	1 x 10 ⁻⁹	1 × 10 ⁻⁷	1.4×10^{-5}	7.5×10^{-4}	1.5 x 10 ⁻²	1.2 x 10 ⁻¹
Fedorenko and Yanysheva data	3×10^{-7}	2.6×10^{-5}	1.2×10^{-3}	2.1×10^{-2}	1.5×10^{-1}	4.8×10^{-1}
Multi-stage Model Neal and Rigdon data	1.7×10^{-6}	1.7×10^{-5}	1.7 x 10 ⁻⁴	1.7×10^{-3}	1.6 x 10 ⁻²	1.5 x 10 ⁻¹
Fedorenko and Yanysheva data	7×10^{-6}	7×10^{-5}	6.8×10^{-4}	6.8×10^{-3}	6.5×10^{-2}	4.8×10^{-1}

The lifetime excess probability of cancer represents the <u>increase</u> in probability of cancer over the normal background incidence, assuming that an individual is continuously exposed to BaP at the indicated daily intake over a 70-year lifetime. There is considerable variation in the estimated risk due to uncertainty introduced by the use of laboratory rodent data, by the conversion to equivalent human dosage, and by the application of hypothetical dose-response curves. In view of several conservative assumptions that were utilized, it is likely that these predictions overestimate the actual risk to humans.

schedule to an equivalent daily lifetime human dose, two assumptions were utilized as recommended by the USEPA (1979). First it was assumed that the response would be the same if the total dose was distributed evenly over the entire experimental lifetime of the animal. Second, it was assumed that a short-duration study attributable to either early mortality or early termination leads to an overestimation of the dose required to produce the observed effect. The average daily dose for the Neal and Rigdon data was adjusted to reflect the actual experimental lifetime versus the theoretical lifetime of the experimental animal. In treating the Neal and Rigdon data, this adjustment lead to a marked reduction (∿1/40) in the effective daily dose level.

- Slight differences in dose-effect relationship between the Neal and Rigdon (1967) and the Fedorenko and Yanysheva (1966) studies which were primarily attributed to the difference in mode of administration (diet vs. gavage, respectively). These may also have been caused by differences in dosing schedules.
- It appears that the total effective dose for the Neal and Rigdon study is probably not a simple product of daily dose times duration but, in fact, may be a more complex power function in which the dose rate has a greater influence on response than treatment duration.
- Development of PAH-induced tumors can be altered by components in the diet, exposure to inducers or inhibitors of microsomal enzymes, as well as cocarcinogenic effects such as from substances present in cigarette smoke.

Keeping these qualifiers in mind, the risks shown in Table 5-64 were combined with the known BaP exposure levels (Table 5-63) to estimate per capita lifetime risks and incidence (excess cancers/year) for BaP exposure routes as shown in Table 5-65.

The highest estimated carcinogenic risks appear to be associated with the subpopulation that smokes. In the general population, the highest estimated risks are associated with exposure to BaP in the diet (e.g., charcoal-broiled meats and fish). Inhalation exposure at the upper limit of the reported concentration ranges in urban areas are higher but are based on limited monitoring data. In contrast, continuous lifetime consumption of BaP-contaminated drinking water at typical concentration levels presents the lowest potential carcinogenic risks.

TABLE 5-65. ESTIMATED RANGES OF CARCINOGENIC RISK TO HUMANS DUE TO BENZO[a]PYRENE EXPOSURE FOR VARIOUS ROUTES

Route	Average Lifetime BaP Exposure (pg/day)	Size of Exposed Population	Estimated Lifetime Excess Probability of Cancer	Estimated Incidence (excess cancers/year)
Typical Diet	0.05	221 × 10 ⁶	4×10^{-6} to 6×10^{-4}	13 - 1,900
Drinking Water	0.0006	221 x 10 ⁶	1×10^{-10} to 7×10^{-6}	<1 - 22
Ambient Air - Urban	0.02 - 2	166 x 10 ⁶	6×10^{-7} to 4×10^{-2}	£ - 95,000
- Rural	0.0002 0.2	55 x 10 ⁶	3×10^{-11} to 3×10^{-3}	<pre>< 1 = 2,400</pre>
Smoking	0.6 b	54 × 10 ⁶ b	3×10^{-4} to 1×10^{-2}	230 - 7,700

A range of probability is given, based on several different dose-response extrapolation models. The lifetime excess probability of cancer represents the <u>increase</u> in probability of cancer over the normal background incidence, assuming that an individual is continuously exposed to BaP at the indicated daily intake over a 70-year lifetime. There is considerable variation in the estimated risk due to uncertainty introduced by the use of laboratory rodent data, by the conversion to equivalent human dosage, and by the application of hypothetical dose-response curves. In view of several conservative assumptions that were utilized, it is likely that these predictions overestimate the actual risk to humans.

but was assumed that the total population of smokers (54 million) smoked on average 25 digarettes per day. From 25% to 30% of smokers consume more than 25 digarettes per day and consequently may receive a higher daily exposure.

With respect to other PAHs included in this group, data were inadequate for quantitative risk assessment purposes. Benz[a]anthracene and dibenz[a,h]anthracene are both carcinogenic in mice by the oral route. They are also complete carcinogens for mouse skin as are benzo[b]-fluoranthene and indeno[1,2,3-c,d]pyrene. Benzo[g,h,i]perylene is a co-carcinogen with BaP and benz[a]anthracene, benzo[b]fluoranthene, chrysene and indeno[1,2,3-c,d]pyrene are all initiators of skin carcinogenesis in mice. No carcinogenicity data were found for acenaphthylene.

5.5.3.2 Non-Carcinogenic Risks

The major focus of studies with compounds in the benzo[a]pyrene group has been their potential for inducing carcinogenic effects. Little information is available on other potential risks to humans from exposure to these compounds (see Section 5.3.1). BaP has been shown to be the most active mutagen of the group, inducing in vivo chromosomal aberrations and sister chromatid exchange in hamsters. Mixed mutagenic responses have been noted with other compounds in this group. Because of its mutagenicity, BaP exposure could also be expected to contribute to the genetic burden of a population, but since extrapolation procedures for genetic risks have not been well established, a quantitative risk assessment for these kinds of health hazards is not presently feasible.

BaP appears to exert little effect on the developing embryo; data were unavailable for the other compounds in this group.

Little information is available on other possible toxic effects associated with exposure to these compounds. However, in assessing the risks to humans associated with exposure to these PAHs, one should not overlook possible augmentation of effects through synergistic or co-carcinogenic mechanisms. Current understanding of the co-carcinogenesis process is not sufficiently adequate to allow estimation of human risk at this time.

5.5.4 Risk to Biota

Data on acute and chronic effects of PAHs are limited and water quality criteria have not been set (USEPA 1980). The only laboratory toxicity test for any of these compounds indicates that 1.0 $\mu g/l$ of benz[a]anthracene caused 87% mortality in bluegill sunfish in six months. Concentrations in this general range (0.2 - 20 $\mu g/l$) have also been found to stimulate growth in some organisms including baceria and freshwater algae. Several field studies have attributed certain tumors found in benthic marine fish to BaP from hydrocarbon-contaminated sediments, but the concentrations of BaP causing these effects were not known.

Monitoring data at detectable levels were very scarce, but those which were available indicate that benz[a]anthracene and benzo[k]fluoranthene have been found in surface water at concentrations from 1 - 1500µg/1. Sediment concentration data were somewhat more extensive and indicated that all of the eight PAHs in this group have been detected in concentrations exceeding 1.0 $\mu g/kg$. Maximum sediment concentrations for this group ranged from 35.8 μ g/kg (indeno[1,2,3-c,d]pyrene) to 2600 µg/kg (dibenz[a,h]anthracene). Uptake of PAHs from sediments by benthic organisms is believed to occur, but the extent of bioavailability of sediment-bound PAHs is not known at this time. Although no data were found attributing specific sediment PAH concentrations to toxic effects, it is believed that PAHs from hydrocarbon-contaminated sediments have caused liver tumors in fish. Many of the higher sediment monitoring observations were found in Puget Sound, the area in which liver tumors in fish have been observed. PAH sediment levels, therefore, which are often found near industrialized areas, may pose some risk to aquatic biota.

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APPENDIX A

NOTE 1:

Fireplaces

Emissions were calculated from emission factors (see Table A-1) that represented the average of three tests (EPA 1980b). The total mass of wood burned in fireplaces in 1976, 2.7 x 10^6 kkg (EPA, 1980a) was extrapolated to 1977, based upon the number of housing units in each region of the country, the percentage of those housing units with fireplaces an average consumption of 98.3 kg/housing unit (EPA 1980a). These assumptions led to the estimate that 2.9 x 10^6 kkg of wood were burned in 1977.

NOTE 2:

Carbon Black

An estimated 1.0 x 10^6 kkg of carbon black were produced in 1977 (SRI, 1979). The emission factors in Table A-1 (Serth and Hughes, 1980) were used to estimate PAH emissions, although the resulting estimates are limited by the fact that testing was performed upstream of an emissions control device (burner).

An estimate of PAH production associated with carbon black manufacture is presented in Table A-3.

Tire Wear

A crude estimate of PAH emission from tire wear associated with carbon black (see Table A-4) was developed from the following data:

- 365 da/yr.
- 7.4 \times 10⁶ bb1/da gasoline consumption (Oil and Gas Journal 1979).
- 14.7 miles/gal average (EPA 1975).
- 42 gal/bbl.
- 0.19 g/vehicle-mile is airborne particulates and 0.15 g/vehicle-mile is deposited on road surface (EPA 1979c).
- Rubber composed of 33% carbon black (SRI 1979).
- Average PAH composition of carbon black in Table A-3 (Locati et al. 1979).

NOTE 3:

Motor Vehicles

Motor vehicle PAH emissions were calculated by using the emission factors in Table A-1 (Hangebrauck, 1967), and estimated 2.7 x 10^{15} meters/yr travelled (see note on tire wear for estimation of vehicle miles travelled), and an assumed emission reduction of two thirds. this emission reduction was based upon the following information:

Fraction of Automobile

<u>Population</u>	<u>Control</u>	Present Reduction
0.32 0.58	catalytic Conventer Engine Modification	99 65
0.10	None	0

The automobile population is from EPA (1978b), and the percent reductions are based on ranges mentioned in that document.

The concentrations of PAHs in used crank case oil are shown in Table A-5. Releases to sewers and landfills were assumed to account totally for 2 x 10^9 ℓ of oil disposed of by the public (Tanacredi 1977); however, this figure does not take into account used oil that is recycled.

NOTE 4:

a. Oil:

Water figure based on 3.6 \times 10⁷ $_{2}$ of various oils -crude (36%), diesel (18%), fuel (42%), waste (2%), lube (0.3%) other (1.7%) - spilled in navigable waters in 1978 (U.S. Coast Guard, 1980).

Land figure based on 5.1 x 10^5 £ of crude oil spilled in 1978 by common carrier (23%), private carrier (22%), rail (6%), and "other" (49%) (U.S. Dept. of Transportation 1980). Average oil Density = 0.85.

b. Gasoline:

Water - 1.1. x 10^7 $_{\rm L}$ spilled: aviation/automobile gasoline (98%) and natural (Casinghead) Gasoline (2%) (U.S. Coast Guard 1980).

Land - 3.7 x 10^6 & spilled: common carrier (53%), private carrier (47%), rail (0.8%), "other" (<0.01%) U.S. Dept. of Transportation 1980). Gasoline density: 0.73.

Table A-1. Emission Factors

	Residential Coal Combustion (g/kg)	Fireplaces (g/kg)	Primary an Auxiliary Wood Heating (9/kg)	d Cigarettes (µg/cig)	Coal Refuse Piles (kg/kg) POM	Forest Fires (ng/y) (dry fuel)	Carbon Black P9/kg	Gasoline (ng/meter)
Acenaphthene	0.039	0.0012	0.0076				•	
Acenaphthylene		0.010	0.057				800	
Anthracene ^a	0.008	0.010	0.076	0.17	0.1	2,500	35	4.3
Benzo[a]anthracene	0.002	0.0008	0.0071	0.02		3,100	4.5	
Benzo[b]fluoranthene ^C	0.002	0.0008	0.0058		0.01	1,300	15	
Benzo[k]fluoranthene	0.002	0.0008	0.0058		0.01	1,300	15	
Benzo[ghi]pery]ene		0.0009	0.0053			2,500	12	47
Benzo[a]pyrene ^d	0.0015	0.0008	0.0040	0.01	0.005	740		11.5
Chrysene ^b	0.002	0.0008	0.0071	0.02		3,100	4.5	
Dibenzo[a,h]anthracene ^e	0.003	0.0001	0.0007		<0.001	•		
fluroanthene	0.005	0.0028	0.019	0.01	0.05	5,500	60	75
Fluorene	0.026	0.0047	0.016			•		, -
Indeno[1,2,3-cd]pyrene	0.002	ND	ND	0.006	<0.001	1,700	<2	
Naphthalene	0.15	0.0403	0.25	3		•		
Phenanthrene	0.008	0.010	0.076	0.36	0.1	2,500	35	30
Pyrene	0.005	0.0028	0.016	0.16	0.05	4,600	500	110

Sources listed in Section 5.1 text.

a) Reported as anthracene/phenanthrene, assumed equal division between them.
b) Reported as chrysene/benzo[a]anthracene, assumed equal division between them.
c) Reported as benzo fluoranthenes, assumed divided solely between benzo[b]fluoranthene and benzo[k]fluoranthene.

d) Reported as benzopyrene(s) and perylene, assumed to be 50% benzo[a]pyrene.
e) Reported as dibenzanthracene, assumed to be solely dibenzo[a,h]anthracene.

Table A-2. Fireplace Population

Region	Number of Housing Units (1977)	Percentage w/Fireplace	Fireplaces
Northeast	17,707,000	47	8,300,000
North Central	21,181,000	33	7,000,000
South	26,422,000	29	7,700,000
West	15,406,000	46	7,100,000
Total			30,100,000

Sources: Census 1979 and EPA 1980a

Table A-5. Concentrations of PAHs in Used Crankcase Oils (mg/l)

 Anthracene	0.3
Benzo[a]anthracene	0.9
Benzo[k]fluoranthene	1.4
Benzo[ghi]perylene	1.7
Benzo[a]pyrene	0.4
Chrysene	1.2
Fluoranthene	4.4
Fluorene	1.5
Phenanthrene	7.8
Pyrene	6.7

Source: Peake and Parker 1980.

Table A-6. Municipal Incinerators Release Factors ($\mu g/kg$ refuse)

	Air ^a	Land	Water ^b
Benzo[a]anthracene ^C	1.5	18	0.08
Benzo[b]fluoranthene ^d	0.5	21	0.01
Benzo[k]fluoranthene ^d	0.5	21	0.01
Benzo[ghi]perylene	1.8	10	0.007
Benzo[a]pyrene ^e	0.04	16	0.016
Chrysene ^C	1.5	18	0.08
Fluoranthene	2.5	12	0.14
Indeno[1,2,3-cd]pyrene	0.77	<2.1	<0.002

Source: Davies 1976.

a) after scrubber
 b) taken as one half reported benzo[a]anthracene + chrysene emissions
 c) taken as one third of benzo[b+k+j]fluoranthene emissions
 d) taken as one half of benzo[a+e]pyrene emissions

e) scrubber water

Table A-3. PAH Associated with Carbon Black $(\mu g/g)a$

Carbon PAH Black Type	Vulcan J	Regal 300	330 HAF	660 GPF	339	Avg	Contained in Carbon Black (kkg)
Anthracene ^b	0.5	ı ND	0.05	ND	1	0.3	0.5
Benzofluoranthenes ^C	10	ND	<0.9	4	7	4	6
Benzo[ghi]perylene	166	16	25	41	164	82	100
Benzopyrenes	20	1	3	8	32	17	30
Fluoranthene	68	9	10	13	52	30	50
Indenopyrene	24	1	0.3	7	35	13	20
Phenanthrene	0.5	ND	0.05	ND	1	0.3	0.5
Pyrene	314	58	47	52	207	140	200

Source: Locati et al. 1979

a) Based 1.6 x 10^6 kkg carbon black production (SRI 1979). b) Reported as anthracene/phenanthrene, assumed equal division among them. c) Excluding benzo[ghi]fluoranthene, reported separately.

Table A-4. PAH Releases from Tire Wear (kkg)a

		rborne	Sedimentary or directly trans-		
	Initial	Reentrained	ferred to Roadway	Total	
Anthracene					
Benzofluoranthenes					
Benzo[ghi]perylene	1	7	7	20	
Benzopyrenes	0.3	1	1	2	
Fluoranthene	0.5	3	2	6	
Indenopyrene	0.2	1	1	2	
Phenanthrene					
Pyrene	2	10	10	20	

See Appendix text for calculations and sources

a) Blanks indicate <1 kkg/yr. For all entries, totals may not add due to rounding.

Table A-9. Concentrations of Various PAHs in Coal and Coal Tar Derivatives (mg/kg)^a

	Coal	Coal Tar	Coal Tar Pitch	Creosote Oil
Acenaphthene Anthracene Phenanthene Benzo[a]anthracene Benzo[a]pyrene Chrysene Fluoranthene Fluorene Naphthalene Pyrene	0.7	10,000 ^b 9,000 ^b 30,000 ^d <0.007 ^d 30 ^d 4,000 ^d 6,000 ^b 10,000 ^b 90,000 ^f 3,000 ^g	<10 ^d 10 ^d <10 ^d	40,000 ^c 20,000 ^c 100,000 ^c <3 ^d neg ^d ,e <1 ^d 40,000 ^c 30,000 ^c 200,000 ^c 30,000 ^c

a) All numbers rounded to one significant figure; blank spaces = data not available.

b) Sources: Rhodes 1954; Lowry 1945 (averaging of data).

c) Source: Stasse 1954; as cited in Weiler 1963.

d) Source: IARC 1973.

e) Less than 0.01 ppm.

f) Source: Rhodes 1954; and Lowry 1945.

g) Source: Rhodes, 1954.

Table A-10. PAH Wastewater Discharge: By-Product Cokemaking^a

	D	ischarge Factors (kg/kkg	coke)	
	Aumonia Liquor	Cooler Blowdown	Benzol Plant	(kkg) Total Discharge
Acenaphthylene Benzo[a]anthracene Benzo[a]pyrene Chrysene Lluoranthene Lluorene Maphthalene Pyrene	0.00073 0.000006 0.000032 0.000045 0.000196 0.000145 0.00395 0.000393	0.000097 0.000032 0.000024 0.000018 0.000323 0.000048 0.0115 0.000026	0.000129 0.000125 MD 0.000155 0.000189 0.000049 0.00341 0.000109	40 7 3 10 30 10 800 20

a) Based on the total of three factors and a 1978 coke production of 44.5×10^6 kkg (DOE 1979). Distribution: 33%-direct, 25%-POIWs, 2%-deep well, 40% quenching (20% land, 20% air) (EPA 1979d).

Source: EPA 1979d.

Table A-7. Emissions of PAHs from Coal-fired Plants and Intermediate/small Oil-fired Units, $\mu g/10^9$ J Fuel

Type of Unit	Benzo[a]- pyrene	Pyrene	Benzo[ghi]- perylene	Phenan- threne	Fluoran- thene
Pulverized coal (vertically-fired dry-bottom furnace)	, 18 - 123	70 - 218	79		80 - 389
Pulverized coal (front-wall-fired dry-bottom furnace)	, 16 - 20	152 - 190	13	190	12 - 152
Pulverized coal (tangentially- fired, dry-bottom furnace)	123	133	142	30	370
Pulverized coal (opposed-, down- ward inclined burners; wet bottom furnace)	20 - 133	37 - 114	142 - 1,042		52 - 199
Crushed coal (cyclone-fired, wet-bottom furnace)	72 - 351	237 - 1,706	34 - 341		42 - 104
Spreader stoker (traveling grate)	<14 - 23	20 - 56			20 - 56
<u>Oil-fired</u> :					
Steam atomized Low pressure air atomized Pressure atomized Vaporized	<19 - 45 853 <38 - <57 <95	46 - 284 5,780 14 - 1,700 1,140	285	1,700 3,320 8,440	53 - 256 1,800 72 - 4,470 14,200

NOTE: Blanks indicate data not available.

Source: Hangebrauck et al. 1967.

Table A-8. Coke-Oven Tar Produced in the United States, Used by Producers, and Sold in 1978 by State (Thousand Liters)

		roduced Used by Producers			Sold for	Sold for Refining into Tar Products			
State	Total	L/kkg of Coal Coked	For Refinery or Topping	As Fuel	Other	Quantity	Valu Thousand Dollars	Average Per Liter	On-hand Dec. 31
Alabama	130,000	27	a			130,000	\$11,970	\$ 0.09	9,700
Calif., Colo., Utah	- 130,000	35	-	_	_	130,000	11,905	0.09	
Illinois	- 59,000	25	-	_	_	60,000	5,593	0.09	13,000
Indiana	- 350,000	33	à	a	a	120,000	12,840		5,700
Ken., Mo., Tenn., Tex	- 34,000	25	-	a	-	31,000	3,021	0.11	21,000
Maryland, New York		32	-	a	_	120,000	•	0.10	2,700
Michigan		a	_	_	_		12,911	0.11	25,000
Minnesota, Wisconsin	- 20,000	23	-	- a	_	a 19,000	1 020	a 10	à
Ohio	- 330,000	31	_	170,000	-		1,938	0.10	6 000 FB
Pennsylvania	- 580,000	35	a	270,000	a	180,000	17,529	0.10	27,000
Virginia, West Virginia	- a	a	a	a	a	280,000	29,425	0.11	55,000
Undistributed	- 240.000	30	500,000	190,000	22 000	150 000	a	à	5,700
	210,000		300,000	190,000	32,000	150,000	12,904	0.09	14,000
Total (1978) ^b	2,100,000	44	500,000	360,000	32,000	1,200,000	120,036	0.10	180,000
At Merchant Plants	89,000	34	С			00 000			
	2,000,000	44		360 000	22 000	89,000	8,986	0.10	4,900
· · · · · · · · · · · · · · · · · · ·	2,000,000	77	500,000	360,000	32,000	1,100,000	111,050	0.10	170,000
Total (1977) ^b	2,200,000	32	570,000	550,000	38,000	1,100,000	106,728	0.10	160,000

Source: DOE 1979.

<sup>a) Included with "Undistributed" to avoid disclosing individual company data.
b) Data may not add to totals shown due to independent rounding.
c) Included with "Furnace Plants" to avoid disclosing individual company data.</sup>

Table A-11. Concentration of Select PAHs in Petroleum Products, mg/kg

	Crude Oil	Gasoline	Kerosene	Petroleum Asphalt	Diesel Fuel	Number 2 Heating Oil
Acenaphthene	ND ^a					
Acenaphthylene	400					
Anthracene	trace	3	0.4		3	4
Benzo(a)anthracene	trace	3 3	<0.1	0.04	0.1	0.04
Benzo(b)fluoranthene	<5		-		• • •	
Benzo(k)fluoranthene	<5					
Benzo(ghi)perylene	•02	2	<0.1		0.03	0.03
Benzo(a)pyrene	1	2 2 2	0.01	0.01	0.07	0.03
Chrysene	<100	2	ND	0.02	0.5	0.6
Dibenzo(a,b)anthracene						
Fluoranthene	100	7	0.09		0.5	2
Fluorene	200					-
<pre>Indeno[1,2,3-cd]pyrene</pre>					•	
Naphthalene	1,000					
Phenanthrene	100		ND		ND	ND
Pyrene	100	5	0.2		0.4	1

a) ND means not detected.

Source: Guerin 1978; Guerin et al. 1978; EPA 1979f

Table A-12. Emissions of PAHs from Petroleum Refining

Process	Emissions, ug/100 m3a Benzo(a)- Benzo(ghi)- pyrene perylene			
Straight run distillation	0.015	0.11		
Pyrolysis	0.30	0.170		
Asphalt production	0.74	1.58		
Petroleum coke production	25.5	0.4		
Petroleum products purification	0.024	_b		

a) Mean values of multiple samples.

Source: Samedov and Kurbanov 1971, as cited in EPA 1979h.

b) Not available.

Emissions of PAHs from Catalyst Regeneration in Petroleum Cracking, $\mu g/m^3$ Oil Charges Table A-13.

Type of Unit	Benzo(a) pyrene	Pyrene	Benzo(ghi)- perylene	Anthra- cene	Phenan- threne	Fluoran- thene
FCC: ^a Regenerator outlet	0.7 - 73	6.4 - 4.450	24 - 67		63,560	7.0 - 3,180
Carbon monoxide boiler outlet	1.7 - 3.4	3.9 - 26	8.8	330		3.2 - 13
нсс:b						
Regenerator outlet	32,600 - 36,700	20,700 - 20,800	47,700 - 60,400	146 - 318	3,340 - 4,600	1,320 - 1,810
TCC:C						
Air lift, regenerator outlet	8,900 - 19,100	21,000 - 41,300	7,000 - 11,450	1,640 - 1,685	52,500 - 56,000	1,685 - 4,610
TCC:			•			
Brucket lift, regen- erator outlet	5	46 - 57				9.17

NOTE: Blanks indicate data not available.

Source: Hangebrauck, et al. 1967.

a) Fluid catalytic cracking.b) Houdriflow catalytic cracking.c) Thermofor catalytic cracking.

Wet deposition is controlled by the precipitation scavenging ratio, r, which expresses the ratio of pollutant concentration in precipitation (ng/1) to pollutant concentration in air ($\mu g/m^3$). The scavenging ratio is calculated by contributions from the vapor and sorbed fractions, i.e.:

$$r = r_{S} (\phi) + r_{V} (1-\phi)$$

where r, r_s and r_v are scavenging ratios for the total airborne mass, the sorbed contaminant, and the vapor phase, respectively. Once the scavenging ratio is known, the wet deposition flux is given by wet flux = rRC_{air} , where R is the rainfall rate. These parameter values have been estimated, as shown below:

		Precipitat	ion Sca	venging	Ratios	
			 -			
РАН	r * _s	$\frac{r_{v}}{}$	Rural	<u>Urban</u>	<u>Combustion</u> <u>Sources</u>	
Naphthalene	6x10 ⁴	53	53	71	2.5x10 ³	
Anthracene	6x10 ⁴	14	130	3.6×10^{3}	1.2×10^{5}	
Benzo[a]pyrene	6×10 ⁴	$5.4x10^4$	6x10 ⁴	6x10 ⁴	1.2×10^5	

For the purpose of further analysis a generic urban environment has been modeled on the basis of characteristics of Philadelphia and Cleveland. The average wind speed is 10.25 kts (5.3 m/s) and the annual rainfall is 1.0 m/yr. The urban area is 100 mi 2 (2.6 x 10 8 m 2).

The generic rural area is defined as a volume of air within which an associated urban source would contribute to rural concentrations. The size of the area is constrained by the half-life in air, such that the concentration is typical of an area significantly affected by the urban source. At a half-life of 5 hours, and typical wind speed of 5.3 m/sec., naphthalene contamination from an urban area could be significant over an area of 10^{10} m² (roughly 40,000 square miles).

Median observed ambient concentrations for three PAHs in urban and rural areas are presented below (White and Vanderslice 1980):

^{*}Tabulated r_s values apply for rural and urban conditions. Near a combustion source the adsorbed phase is expected to be associated with larger particles resulting in a scavenging ratio, $r_s = 1.2 \times 10^5$.

	Median Observed			
70.4.17	Concentration	on (ug/m^3)		
PAH	Rural	Urban		
Naphthalene	7×10^{-5}	5x10 ⁻⁴		
Anthracene	1x10 ⁻³	8×10^{-3}		
Benzo[a]pyrene	1×10^{-3}	1×10^{-2}		

Using the pathway evaluation method we have estimated the emission that would result in a specific ambient concentration, given the degradation rate, deposition velocity, rainfall rate, and precipitation scavenging ratio. Applying the equation for wet flux and dry flux we also estimated the amount deposited in the generic rural and urban study areas. Then, by comparison of the deposition rates with emission rates, we estimated the fraction of total atmospheric emissions of each of the three PAHs which would be deposited within the urban and rural study areas. These results are shown below:

РАН	% of Emissions Dry Deposited Rural Urban*		Deposition Rate % of Emissions Wet Deposited Rural Urban		% Deposited Rural Urban	
Naphthalene	2	2-3	<1	<1	2	2-3
Anthracene	1	4-19	<1	1-7	1	5-26
Benzo[a]pyrene	22	19	4	4-7	26	23-26

From the results above, it is apparent that a very small fraction of atmospheric emissions of naphthalene are deposited on land or water surfaces. The fraction of air emitted anthracene that is eventually deposited is uncertain, but could be fairly large percentage. Approximately one-fourth of all benzo[a]pyrene emitted into the atmosphere will eventually be deposited on land and water surfaces, where it would contribute to the contamination of surface runoff and surface water bodies.

^{*}The range for urban areas reflects alternative assumptions that the chemical has equilibrated with ambient aerosol or remains associated with particulates in the plume from the combustion zone.

APPENDIX B. APPLICATION OF THE AIR-TO-SURFACE PATHWAY EVALUATION METHOD FOR POLYNUCLEAR AROMATIC HYDROCARBONS

A method has been developed for estimating airborne toxicant deposition rates (air-to-surface pathway evaluation method, Arthur D. Little, Inc., 1981). The method accounts for both wet and dry deposition to land and water surfaces. When deposition to a watershed or other land area is estimated, this can be interpreted as an upper bound on the chemical loading to an associated surface water body resulting from air deposition, since only a fraction of the mass deposited on land surfaces will be delivered to the water body. The air-to-surface pathway evaluation method accounts for the partitioning of an airborne contaminant between adsorbed and vapor phases, with differing deposition rates inferred for the separate phases. It relies on fundamental physicochemical properties and is designed to use available data, while filling in data gaps with estimated values of various parameters. The evaluation method has been applied to naphthalene, anthracene, and benzo[a]pyrene. Each of the three PAHs modeled has an atmospheric chemical degradation of roughly 0.1 hr.-1 leading to halflives of 5-10 hours (Radding et al. 1976). Under typical meteorologic conditions this corresponds roughly to the travel time across major urban areas. Since urban areas also would be expected to have much greater emission densities than rural areas, significant urban/rural differences in air concentrations of PAHs are expected, and indeed observed. These factors suggest that deposition rates under urban and rural conditions should be considered separately.

One of the most important chemical properties influencing air-to-surface transfer is the vapor pressure. The vapor pressure affects the partitioning of airborne contaminants between vapor and adsorbed phases. The deposition rate is typically much greater for the adsorbed fraction of the airborne contaminant. The effect of vapor pressure is expressed by equation (5) of the Arthur D. Little report (1981):

$$\phi = \frac{.165\theta}{p_o + .165}$$

where

\$ is the adsorbed fraction of the total airborne mass,

 θ is the available aerosol surface area $\frac{\text{cm}^2}{\text{cm}^3}$ and

 $\mathbf{p}_{\mathbf{0}}$ is the saturation vapor pressure of the contaminant at ambient temperature (torr)

The available aerosol surface area is typically greater in urban areas where concentrations of total suspended particulates are higher

than in rural areas. In the plume from a combustion source, the aerosol surface area is even higher than typically found in urban air. Application of the above equation results in aerosol partitioning for the three PAHs as shown below:

	Adsorbed	Fraction of	Total Airborne Mass
PAH	Rural	Urban	Near Combustion Sources
Naphthalene	$7x10^{-6}$	$3x10^{-4}$	0.02
Anthracene	0.002	0.06	0.97
Benzo[a]pyrene	.99	1.00	1.00

Benzo[a] pyrene is strongly partitioned with the aerosol phase, regardless of ambient conditions, while at the other extreme naphthalene exists primarily as a vapor in the atmosphere. Anthracene exhibits intermediate properties, and the adsorbed fraction is sensitive to ambient conditions.

The dry deposition flux is proportional to the dry deposition velocity, $\boldsymbol{V}_{\underline{d}},$ i.e.,

where

 $\mathbf{C}_{\mbox{air}}$ is the ground-level air concentration.

The dry deposition velocity with respect to the total airborne contaminant is calculated as the (mass) weighted average of the deposition velocity for the vapor and sorbed fractions, i.e.,:

$$V_d = V_{d,s}^{\phi} + V_{d,v}^{\phi}$$
 (1- ϕ) (Eq. 6 of Arthur D. Little 1981)

According to the air-to-surface pathway method (Arthur D. Little 1981), the respective dry deposition velocities are given below:

	Dry Deposition Velocity					
РАН	Vd,v (<u>cm/sec</u>)	Vd,s (<u>cm/sec</u>)	Rural	Urban	Combustion Source	
Naphthalene	0.04	1	0.04	0.04	0.06	
Anthracene	0.02	1	0.02	0.08	1.00	
Benzo[a]pyrene	0.02	1	1.00	1.00	1.00	

APPENDIX REFERENCES

Arthur D. Little, Inc. Air-to-surface pathway evaluation methodology. Draft final report. Contract N. 68-01-5949. Washington, DC: Monitoring and Data Support Division, Office of Water Regulations and Standards, U.S. Environmental Protection Agency: 1981.

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