6. POTENTIAL FOR HUMAN EXPOSURE

6.1 OVERVIEW

Hexachlorobenzene has been identified in at least 113 of the 1,699 hazardous waste sites that have been proposed for inclusion on the EPA National Priorities List (NPL) (HazDat 2007). However, the number of sites evaluated for hexachlorobenzene is not known. The frequency of these sites can be seen in Figure 6-1.

Hexachlorobenzene has not been manufactured in the United States (as an end-product) since its last registered use as a pesticide was voluntarily canceled in 1984. Small amounts of hexachlorobenzene are still released to the environment by industrial activities as fugitive and stack emissions, or in waste water. It is released as a by-product during the manufacture of several chlorinated solvents or as an impurity in several currently registered pesticides. Additional amounts of hexachlorobenzene are also formed during combustion processes such as incineration of municipal refuse (EPA 1986b) or through use of pyrotechnic mixtures (Karlsson et al. 1991). Bailey (2001) provided a detailed analysis of the global emissions of hexachlorobenzene during the 1990s from these various sources (see Section 6.2).

Hexachlorobenzene is among the most persistent environmental pollutants because of its chemical stability and resistance to degradation. If released to the atmosphere, hexachlorobenzene exists primarily in the vapor phase and degradation is extremely slow. Half-life estimations for hexachlorobenzene in the atmosphere are highly variable, ranging from 0.63 years in tropical/subtropical regions, to 1.94 years in temperate/boreal regions, to 6.28 years in polar regions (Wania and Mackay 1995). A global calculated half-life of 1.69 years was attained from a measured hydroxyl rate constant (2.7x10⁻¹⁴ cm³/molecule-second) (Brubaker and Hites 1998). Long-range global transport is possible from the temperate to the polar regions (Barber et al. 2005). Physical removal of hexachlorobenzene from the air may occur via washout by rainfall or snowfall, or via dry deposition. If released to water, hexachlorobenzene will partition from the water column into sediment and suspended particulate matter (EPA 1981).

In water, it is a persistent chemical not readily degraded by either abiotic or biotic processes. The halflife value of hexachlorobenzene is estimated to range from 2.7 to 5.7 years in surface water and from 5.3 to 11.4 years in groundwater (Howard et al. 1991). Volatilization of hexachlorobenzene from the water column will occur; however, the compound's strong adsorption to particulates and organic matter in water slow the rate of volatilization. If released to soil, hexachlorobenzene can volatilize from the soil surface, but will be strongly adsorbed to organic matter and is generally considered immobile with respect





6. POTENTIAL FOR HUMAN EXPOSURE

to leaching (HSDB 2012). Its half-life value in soils is estimated to range from 3 to 6 years (Beck and Hansen 1974). Hexachlorobenzene bioaccumulates significantly in both terrestrial and aquatic food chains. Bioconcentration factors (BCFs) as high as 17,000,000 and 21,900 have been reported for lichens and fish, respectively (Muir et al. 1993; Veith et al. 1979). Hexachlorobenzene has been used as a referent compound for BCFs in fish (Adolfsson-Erici et al. 2012).

Temporal trends of hexachlorobenzene levels in the environment vary, depending on the media and study location, and the time period measured, but the average half-life from all of these studies is about 9 years. Estimates made of the present-day burden in the environment range between 20 and 60,000 million pounds and are dominated by the loadings in treated and background soils, sediments, and oceans (Barber et al. 2005).

Monitoring data for hexachlorobenzene levels are extensive in the Great Lakes region where production of chlorobenzenes was historically high. Atmospheric monitoring detected the compound at mean and median concentrations of 36.68 (0.03668 ng/m³) and 30.94 (0.03094 ng/m³) pg/m³, respectively, from 56 air samples in Villeroy, Quebec in 1992 (Poissant et al. 1997). Hexachlorobenzene has also been detected in minute amounts (up to 0.174 ng/L [ppt]) in precipitation samples from the Great Lakes region (Chan et al. 1994) and in precipitation samples collected from Villeroy, Quebec in 1992 (0.04 ng/L) (Poissant et al. 1997). It was also detected in drinking water in three cities on Lake Ontario at a mean concentration of 0.1 ppt (Oliver and Nicol 1982a). Sediment samples (2 cm depth) collected from lakes (Allen-Gil et al. 1997) and landfills have also been contaminated with hexachlorobenzene (Yasuhara et al. 1999).

Concentrations of hexachlorobenzene have been reported for a variety of commercial fish species in the Great Lakes (Allen-Gil et al. 1997; Kuchlick and Baker 1998) with concentrations up to 17 ppb in raw fish fillets (Newsome and Andrews 1993; Zabik et al. 1995). In the National Pesticide Monitoring Program, concentrations as high as 700 ppb were reported in whole fish samples collected from contaminated areas (Schmitt et al. 1990). This chemical has also been detected in the fatty tissues and muscle of a wide variety of waterfowl (Foley 1992; Gebauer and Weseloh 1993; Swift et al. 1993), marine mammals (Becker et al. 1997; Langlois and Langis 1995), and mammals (Corsolini et al. 1999). In terrestrial ecosystems, hexachlorobenzene has been detected in lichens (Muir et al. 1993) and in caribou that graze primarily on lichens (Elkin and Bethke 1995). Concentrations of hexachlorobenzene in these fish and wild game species can be a source of hexachlorobenzene exposure to man.

6. POTENTIAL FOR HUMAN EXPOSURE

Hexachlorobenzene residues have been detected in 76% of samples analyzed as part of the National Human Adipose Tissue Survey (FY82) (EPA 1986c). These hexachlorobenzene residues are most likely the result of consumption of low levels of hexachlorobenzene in food, with a calculated yearly intake of 68, 22, and 5 µg for adults, toddlers, and infants, respectively (EPA 1986b). Compared to this, exposure to hexachlorobenzene via inhalation or through drinking water is relatively low. Human exposure may also occur via dermal contact with contaminated soil or sediment or via ingestion of contaminated soil by children. In occupational settings, exposure occurs primarily via inhalation or dermal contact.

Due to extensive research conducted on hexachlorobenzene, the data reported herein do not encompass complete and thorough research for this chemical.

6.2 RELEASES TO THE ENVIRONMENT

The Toxics Release Inventory (TRI) data should be used with caution because only certain types of facilities are required to report (EPA 2005). This is not an exhaustive list. Manufacturing and processing facilities are required to report information to the TRI only if they employ 10 or more full-time employees; if their facility is included in Standard Industrial Classification (SIC) Codes 10 (except 1011, 1081, and 1094), 12 (except 1241), 20–39, 4911 (limited to facilities that combust coal and/or oil for the purpose of generating electricity for distribution in commerce), 4931 (limited to facilities that combust coal and/or oil for the purpose of generating electricity for distribution in commerce), 4939 (limited to facilities that combust coal and/or oil for the purpose of generating electricity for distribution in commerce), 4939 (limited to facilities that combust coal and/or oil for the purpose of generating electricity for distribution in commerce), 4939 (limited to facilities regulated under RCRA Subtitle C, 42 U.S.C. section 6921 et seq.), 5169, 5171, and 7389 (limited S.C. section 6921 et seq.), 5169, 5171, and 7389 (limited S.C. section 6921 et seq.), 5169, 5171, and 7389 (limited section section section of a contract or fee basis); and if their facility produces, imports, or processes ≥25,000 pounds of any TRI chemical or otherwise uses >10,000 pounds of a TRI chemical in a calendar year (EPA 2005).

Additional releases of hexachlorobenzene to the environment occur from processes such as combustion of coal, cement, sewage sludge, or biomass; iron sintering; incineration of municipal, hazardous, or medical wastes; during the use of pyrotechnic mixtures; as a byproduct during the manufacture of several chlorinated solvents; or as an impurity in several currently registered pesticides (Bailey 2001; Barber et al. 2005). Quantitative information on releases of hexachlorobenzene to specific environmental media is discussed below.

6. POTENTIAL FOR HUMAN EXPOSURE

6.2.1 Air

Estimated releases of 1,211 pounds (~0.55 metric tons) of hexachlorobenzene to the atmosphere from 65 domestic manufacturing and processing facilities in 2013, accounted for about 5.5% of the estimated total environmental releases from facilities required to report to the TRI (TRI13 2014). These releases are summarized in Table 6-1.

Releases to the air from the production of chlorinated solvents, where hexachlorobenzene is a minor byproduct, have been estimated at 0.3 kg (0.7 pounds) annually and emissions from municipal refuse incineration have been estimated at 85–8,512 kg (187–1,870 pounds) annually (Bailey 2001). As incineration has emerged as a prevalent technology for reducing the bulk of hazardous and nonhazardous wastes, investigations have shown that even with careful controls it is virtually impossible to eliminate unwanted by-products (Products of Incomplete Combustion [PICs]) (Martens et al. 1998). Slight temperature differences on the surfaces of incinerator kiln and reactor components, or other reactions in flues, can lead to the formation of numerous chemical compounds. Where the original wastes contain organochlorines, one type of toxicant may be transformed into another (Dellinger et al. 1989). Hexachlorobenzene, for example, has been detected at concentrations ranging from 20 to 70 ppm in gases emitted by the thermal degradation of toxaphene (Lahaniatis et al. 1992). Since incineration of wastes is a growing global phenomenon, there are concerns that inadequate management attention is given to minimizing PICs. For hexachlorobenzene, and many other organochlorines that can be dispersed widely through atmospheric transport pathways, the virtual absence of data on PICs can lead to complications in estimating environmental releases and mass balances of hexachlorobenzene for regional areas or on a global scale (Lahaniatis et al. 1992).

Hexachlorobenzene emissions were reported as 0.94-3.8 g/year from siderurgies in Portugal and 0.013– 1.7 mg/year from metal non-siderurgy plants (Antunes et al. 2012). Hexachlorobenzene was detected in the emissions from cement plants (0.98–60.5 ng/m³) and from waste incinerators (3.21–2,500 ng/m³) (Wegiel et al. 2011).

Nonpoint source dispersal of hexachlorobenzene historically has resulted from its use as a seed fungicide (Beyer 1996) and results from the use of a number of registered pesticides in which it is a contaminant. Seven major pesticides (chlorothalonil, DCPA or Dacthal[®], pentachlorophenol or pentachlorophenol, picloram, PNCB or quintozene, atrazine, and simazine) in current use contain up to 0.3% hexachloro-

		Reported amounts released in pounds per year ^b								
							Total release			
Statec	RF^d	Air ^e	Water ^f	Οla	Land ^h	Other ⁱ	On-site ^j	Off-site ^k	On- and off-site	
AL	4	0	0	0	14	0	14	0	14	
AR	1	0	0	0	0	0	0	0	0	
CA	2	1	0	0	0	0	1	0	1	
CO	1	0	0	0	0	0	0	0	0	
DE	1	0	1	0	197	0	1	197	197	
FL	1	0	0	0	0	0	0	0	0	
ID	1	0	0	0	1	0	0	1	1	
IL	1	0	0	0	1	0	0	1	1	
IN	2	0	0	0	268	0	0	268	268	
KS	1	0	0	1	0	0	1	0	1	
KY	2	909	1	0	98	0	910	98	1,008	
LA	9	122	22	0	40	0	184	0	184	
MI	1	0	0	0	31	0	31	0	31	
MN	1	0	0	0	0	0	0	0	0	
MS	2	21	0	0	79	0	100	0	100	
MT	1	10	0	0	0	0	10	0	10	
NY	3	0	0	0	3	0	0	3	3	
ОН	5	3	0	6	25	0	3	31	34	
OR	4	0	0	0	546	0	542	4	546	
PA	2	1	0	0	9	0	1	9	10	
SC	1	0	0	0	0	0	0	0	0	
TN	1	0	0	0	159	0	159	0	159	
ТΧ	14	139	27	519	17,790	0	18,457	18	18,475	
UT	3	5	0	0	943	0	948	0	948	

Table 6-1. Releases to the Environment from Facilities that Produce, Process, orUse Hexachlorobenzenea

Table 6-1. Releases to the Environment from Facilities that Produce, Process, orUse Hexachlorobenzenea

				Reported amounts released in pounds per year ^b					
								Total rele	ease
Statec	RF^d	Air ^e	Water ^f	Οla	Land ^h	Other ⁱ	On-site ^j	Off-site ^k	On- and off-site
WA	1	0	0	0	0	0	0	0	0
Total	65	1,211	51	526	20,203	0	21,362	629	21,991

^aThe TRI data should be used with caution since only certain types of facilities are required to report. This is not an exhaustive list. Data are rounded to nearest whole number.

^bData in TRI are maximum amounts released by each facility.

^cPost office state abbreviations are used.

^dNumber of reporting facilities.

^eThe sum of fugitive and point source releases are included in releases to air by a given facility.

^fSurface water discharges, waste water treatment-(metals only), and publicly owned treatment works (POTWs) (metal and metal compounds).

⁹Class I wells, Class II-V wells, and underground injection.

^hResource Conservation and Recovery Act (RCRA) subtitle C landfills; other onsite landfills, land treatment, surface impoundments, other land disposal, other landfills.

Storage only, solidification/stabilization (metals only), other off-site management, transfers to waste broker for disposal, unknown

The sum of all releases of the chemical to air, land, water, and underground injection wells.

^kTotal amount of chemical transferred off-site, including to POTWs.

RF = reporting facilities; UI = underground injection

Source: TRI13 2014 (Data are from 2013)

6. POTENTIAL FOR HUMAN EXPOSURE

benzene as an impurity. Hexachlorobenzene was also an impurity in lindane; however, all of lindane's remaining uses were cancelled in 2006. When these pesticides are applied in sprays, they have the greatest potential for release into the air. Most of the pesticide, and the hexachlorobenzene impurities, end up on the top layer of the soil and can become airborne through volatilization of the vapor or adsorbed to soil particles. The hexachlorobenzene agriculturally applied through the use of these eight pesticides amounts to an estimated 1,270 kg/year (2,790 pounds/year); however, the total amount of hexachlorobenzene actually released into the air could not be estimated (Bailey 2001). At its peak, the amount of hexachlorobenzene emitted from treated soil to air may have been in the millions of pounds per year, which would have made it a significant source of hexachlorobenzene in the environment. While the amount of hexachlorobenzene being emitted from present day soil is much lower, only a small amount of re-emission of hexachlorobenzene from soil to air is required to maintain the current air concentrations given its persistence (Barber et al. 2005).

Other minor sources of hexachlorobenzene releases to the air come from the use of pyrotechnic mixtures that produce white obscurant screening smokes (Karlsson et al. 1991) and fireplace and woodstove combustion (Gullett et al. 2003). Screening smokes are used by the military to obscure vision and hide targets, and are used by civilian firefighters during fire training sessions. In a study of emissions from residential fireplace and woodstove combustion in the San Francisco Bay Region, mean emission factors of 13 ng/kg oak for woodstoves and 310, 380, and 990 ng/kg oak, pine, and artificial log, respectively, for fireplaces were measured (Gullett et al. 2003).

6.2.2 Water

Estimated releases of 51 pounds (~0.02 metric tons) of hexachlorobenzene to surface water from 65 domestic manufacturing and processing facilities in 2013, accounted for about 0.2% of the estimated total environmental releases from facilities required to report to the TRI (TRI13 2014). This estimate includes releases to waste water treatment and publicly owned treatment works (POTWs) (TRI13 2014). These releases are summarized in Table 6-1.

The principal release of hexachlorobenzene into water in the past has been through direct discharges from chemical solvent manufacturing facilities. Total production of chlorobenzenes in the United States has declined from more than 300,000 metric tons (300 million kg or 661 million pounds) in 1970 to about 200,000 metric tons (200 million kg or 441 million pounds) in 1980. The total amount of hexachlorobenzene released as a by-product in production of all chlorinated solvents has been estimated to range

6. POTENTIAL FOR HUMAN EXPOSURE

from 70,343 to 241,311 kg/year (154,000–532,000 pounds/year) (EPA 1986b). Estimated hexachlorobenzene releases into water from these sources, however, were only 70 kg/year (154 pounds/year) (EPA 1986b).

6.2.3 Soil

Estimated releases of 20,203 pounds (~9.2 metric tons) of hexachlorobenzene to soils from 65 domestic manufacturing and processing facilities in 2013, accounted for about 97% of the estimated total environmental releases from facilities required to report to the TRI (TRI13 2014). An additional 526 pounds (~0.2 metric tons), constituting about 2.4% of the total environmental emissions, were released via underground injection (TRI13 2014). These releases are summarized in Table 6-1.

Historically, hexachlorobenzene was released to soils directly through its application as a fungicide on seed grains. Even after use of hexachlorobenzene as a pesticide ceased, an estimated 95% of hexachlorobenzene produced in the manufacture of chemical solvents was disposed of in land applications (EPA 1986b). Current releases to soils may occur through land disposal of hexachlorobenzene-containing wastes, from discharges from manufactured facilities, and from use of currently registered pesticides containing small amounts of hexachlorobenzene. Contamination of soil and sediment with hexachlorobenzene is frequently secondary to the discharge of contaminated water, from which the hexachlorobenzene is then adsorbed by the soil or sediment. Sediment samples (2 cm depth) were collected from four lakes (Feniak, Desperation, Schrader, and Elusive) from the foothills of the Brooks Range, Alaska. All lakes were oligotrophic, and the pH was neutral to slightly alkaline. The mean concentration among the six samples was 0.17 ng/g dry weight, derived from the concentrations of the sediments from Lakes Feniak (0.27 ng/g dry weight), Desperation (0.08 ng/g dry weight), Schrader (0.11 ng/g dry weight), and Elusive (0.21 ng/g dry weight) (Allen-Gil et al. 1997). Substantial levels of hexachlorobenzene, ranging from 0.5 to 460 ppb, have been detected in sediment cores sampled at 1 cm intervals to a sediment depth of 8 cm in Lake Ontario. The highest sediment contamination in Lake Ontario was found in sediments 1– 2 cm in depth which correspond to sediments laid down from 1971 to 1976 during a period of high U.S. production of chlorobenzenes (Oliver and Nicol 1982a). Although no studies concerning the release of hexachlorobenzene from landfills have been located for the United States, Yasuhara et al. (1999) sampled leachates from 11 landfills in Japan. Leachates were sampled at the outlet of the leachate collecting pipe from open and controlled landfills. Hexachlorobenzene concentrations in the 11 leachate samples from 11 landfills ranged from not detected to 0.054 ng/L. Of the 11 sites, 6 are currently under reclamation and at least 3 sites were sampled 12-17 years after completion of reclamation. Site 2, which was sampled

6. POTENTIAL FOR HUMAN EXPOSURE

after 14 years of reclamation, had no detectable levels of hexachlorobenzene. Leachates from sites 3 and 4, which were sampled after 17 and 12 years of reclamation, had hexachlorobenzene concentrations of 0.033 and 0.054 ng/L, respectively. Site 4 had the highest detection of hexachlorobenzene concentration. The median concentration among these sites was 0.03 ng/L.

The presence of hexachlorobenzene as an impurity in several registered pesticides appears to be a continuing source of exposure for the general population. The pesticides containing impurities of hexachlorobenzene include: picloram, PCNB or quintozene, chlorothalonil, DCPA or Dacthal[®], pentachlorophenol or pentachlorophenol, atrazine, simazine, and lindane (Bailey 2001; Kutz et al. 1991). Picloram is a herbicide used in agriculture and silviculture to control broad-leaf weeds and conifers in grasses (MacBean 2010). PCNB is a fungicide used to control diseases of turf, ornamentals, cole crops, potatoes, cotton, and other agricultural and horticultural crops (EPA 2009a). In February 2009, several uses of PCNB were cancelled. These uses included application of PCNB on golf course roughs; residential sites including lawns, yards, and ornamental plants and gardens around homes and apartments; grounds around day care facilities; school yards; parks (except industrial parks); playgrounds; and athletic fields (except professional and college fields) (EPA 2009a). Chlorothalonil (sold under the trade name Bravo®) is a fungicide used on horticultural crops, golf courses, and residential turf, and as a biocide in paints and wood preservatives. Dacthal® is a pre-emergent herbicide widely used in agriculture and on lawns and turf grass (EPA 1986b). Pentachlorophenol is an insecticide and fungicide used to protect timber from fungal rot and wood-boring insects (MacBean 2010). Atrazine and simazine are selective herbicides used to control broadleaf and grassy weeds in corn and other crops (MacBean 2010). Lindane is an insecticide and fumigant that was used on a wide range of soil-dwelling and plant-eating insects and as a seed treatment for barley, corn, oats, rye, sorghum, and wheat. All uses of lindane were cancelled in 2006 and its only current use in the United States is in shampoos and lotions to treat lice and mites.

Although hexachlorobenzene impurities in Dacthal[®] were as much as 10% in the early 1970s, current levels of hexachlorobenzene contamination in all five pesticides are much lower. A registration standard was issued for picloram in 1985 that specified a maximum hexachlorobenzene content of 0.02%. By the terms of an EPA PCNB rebuttal presumption against registration in 1982, PCNB registrants agreed to reduce hexachlorobenzene contamination levels in PCNB to 0.5% by 1983 and to 0.1% by April 1988. A registration standard was issued by EPA for chlorothalonil in 1984 requiring that hexachlorobenzene contamination not exceed 0.05%. Since 1973, the maximum allowable hexachlorobenzene content of technical grade DCPA (Dacthal[®]) has been 0.3% (EPA 1986b).

6. POTENTIAL FOR HUMAN EXPOSURE

The annual total emission of hexachlorobenzene in the United States during the 1990's was estimated to average 2,785 kg (Bailey 2001). These emissions are indirect or unintentional emissions from the manufacture and use of products that may contain varying amounts of hexachlorobenzene as a contaminant. This estimated total was calculated using emission subtotals of 1,270 kg for pesticide use, 0.3 kg for chlorinated solvent use, 399 kg from manufacturing processes, 156 kg from metal industries (using mean emission factors), and 960 kg from combustion sources (using mean emission factors).

Earthworms have been utilized as biosentinel for hexachlorobenzene soil presence (Vampre et al. 2010).

6.3 ENVIRONMENTAL FATE

6.3.1 Transport and Partitioning

Hexachlorobenzene has a moderate vapor pressure $(1.09 \times 10^{-5} \text{ mmHg})$ (O'Neil et al. 2006) and has a very low solubility (0.005815 mg/L) (Yalkowsky 1992) in water (see Table 4-2). If released to the atmosphere, hexachlorobenzene can exist in both the vapor phase and in association with particulates (Eisenreich et al. 1981); however, monitoring studies have demonstrated that the vapor phase generally predominates (Ballschmiter and Wittlinger 1991; Bidleman et al. 1989; Lane et al. 1992). Hexachlorobenzene concentrations in the vapor phase represented 92–100% of the total hexachlorobenzene concentration in air samples collected in a monitoring study conducted in Ontario, Canada (Lane et al. 1992). Although physical removal of hexachlorobenzene from the atmosphere is possible via both wet and dry deposition (Howard 1990), the compound is hydrophobic, and somewhat resistant to wet deposition scouring unless it becomes sorbed to airborne dust or cloud condensation nuclei. Its resistance to wet deposition tends to slow down its transfer across the equatorial areas between the northern and southern hemispheres. At high latitudes, the typically cold air conditions encourage dry deposition of aerosols (Ballschmiter and Wittlinger 1991; Lane et al. 1992; Wania and Mackay 1993). These factors lead to atmospheric pathways as a major transport mechanism for hexachlorobenzene. The net residence time of hexachlorobenzene in the atmosphere is significantly less than 1 year, and is based on physical translocation and not on chemical transformation (Ballschmiter and Wittlinger 1991). The atmospheric mechanisms can operate over large distances, perhaps on a hemispheric scale (Kelly et al. 1991). One study modeled the atmospheric residence time of hexachlorobenzene and calculated the transport distance (the distance over which 50% of the chemical is removed) to be 10^5 km (Barber et al. 2005). At higher latitudes, transfers and partitioning back to aqueous systems may be accentuated by the cold, dry air. Since these areas are not zones of hexachlorobenzene production or use, the presence of such contaminants has attracted considerable attention in research studies (Ballschmiter and Wittlinger 1991).

6. POTENTIAL FOR HUMAN EXPOSURE

The Henry's law constant value for this compound (5.8x10⁻⁴ atm-m³/mol) (ten Hulscher et al. 1992) suggests that releases of hexachlorobenzene to surface water will volatilize at a moderate rate, and that volatilization can be a significant transfer mechanism (Thomas 1990). If released to water, adsorption of hexachlorobenzene to sediment or suspended particulate matter is also expected to be significant on the basis of the high organic carbon partition coefficient (K_{oc}) value (EPA 1981) (see Table 4-2). This tendency to partition to sediment and adsorb to suspended solids in the water column attenuates the rate of volatilization. Since hexachlorobenzene will adsorb strongly to soil particles and sediments, it may build up in the bottom sediments of large aquatic systems such as the Great Lakes. The concentration of hexachlorobenzene in Lake Ontario sediment is about 1 million times higher than in Lake Ontario water (Oliver and Nicol 1982a). In Lakes Superior, Michigan, and Huron, the very large sizes of the water bodies, considerable depths, and appreciable retention times have allowed much of the historical organochlorine burden to become immobilized in bottom sediments, with a concomitant reduction in the levels of hexachlorobenzene found in surface waters. In the upper Great Lakes, the vast majority of the ongoing mass balance inputs seem related to atmospheric deposition (Eisenreich et al. 1981). For other parts of the Great Lakes system, and especially the connecting waters and Lakes Erie and Ontario, mass balance studies can give different impressions. Detailed studies on Lake Erie suggest that well over half of the hexachlorobenzene inputs come from wet or dry atmospheric deposition processes (270 kg/year or 600 pounds/year) (Kelly et al. 1991). However, a significant portion (110 kg/year or 240 pounds/year) also comes from river pathways such as the highly polluted Detroit River via surface runoff and contaminated sediments resuspended during their passage through the connecting waters. Because of its strong adsorption to sediment, most of the hexachlorobenzene is transported with silt and sediment particles during floods, and very little is dissolved in the water. Quemerais et al. (1994) reported that 23% of the hexachlorobenzene in whole water samples collected from the St. Lawrence River was associated with the dissolved phase, while 77% was associated with the particulate phase. Although Rostad et al. (1988) did not quantify the percentage of hexachlorobenzene found in the dissolved versus the particulate phase, they did report that hexachlorobenzene was one of the organic compounds associated with suspended sediment particles in several river systems within the Mississippi River drainage area. In a 1999 study, Rostad et al. (1999) measured hexachlorobenzene concentrations in suspended sediment particles within the Mississippi River in the spring and summer of 1989 and 1990. Concentrations of hexachlorobenzene fluctuated between Winfield, Missouri and Belle Chasse, Louisiana during the spring and the summer; however, in both seasons, the concentration was higher at Chasse, Louisiana (1.9 ng/g in the spring; 2.1 ng/g in the summer) than at Winfield, Missouri (1.2 ng/g in the spring; 0.13 ng/g in the summer). Furthermore, Rostad et al. (1999) estimated annual transport of hexa-

6. POTENTIAL FOR HUMAN EXPOSURE

chlorobenzene via suspended sediments to the Gulf of Mexico by averaging the St. Francisville, Louisiana and Belle Chasse, Louisiana daily loads, averaging the spring and summer values for hexachlorobenzene, and projecting annual transport. The Gulf of Mexico receives an estimated 145 kg/year hexachlorobenzene via suspended sediment particles of the Mississippi River (Rostad et al. 1999).

Because of its high sorption characteristics, hexachlorobenzene is expected to be immobile in soil and unlikely to leach into groundwater (Swann et al. 1983). At waste disposal sites, where bioremediation techniques are proposed to reduce the mass of carbon-containing contaminants, there is the potential for augmenting the leaching properties of hexachlorobenzene and other organochlorines. The lipid materials in bacterial cell membranes may lead to a repartitioning of hexachlorobenzene sorbed to soil colloids. This can lead to a phenomenon called facilitated transport where the mobility of hydrophobic pollutants such as hexachlorobenzene adsorbed to soils may be enhanced by biosorption on bacteria and move into aquifers along with the bioremedial bacterial cultures (Lindqvist and Enfield 1992). Except at NPL sites, however, this potential source of groundwater pollution would seem to be remote.

The Henry's law constant value suggests that hexachlorobenzene released to moist soil will volatilize at a moderate rate. Several studies have indicated that volatilization may be a significant mechanism for loss of hexachlorobenzene released to soils. Beall (1976) studied the persistence of aerially applied hexachlorobenzene (equivalent to 10 ppm in the top 5 cm of soil) in a simulated pasture maintained for 19 months in a greenhouse. Twenty hours postapplication, the top 2 cm of soil contained 5.6 ppm (airdry basis). Hexachlorobenzene concentrations in the top 2 cm of soil found after 0.5, 1, 6.5, 12, and 19 months were 45.2, 24.4, 7.9, 4.7, and 3.4% of day 1 values, respectively. However, no significant change in the deeper 2–4 cm layer of soil which averaged hexachlorobenzene residues of 0.11 ppm was observed over the 19-month study. Concentrations of hexachlorobenzene in pasture grass on day 1 were 1,060 ppm, but 2 weeks postapplication only 15.6 ppm (1.5% of day 1 residues) was detected. Although hexachlorobenzene volatilized fairly rapidly from plant and soil surfaces, it could be persistent within the soil if treated surface soil were mixed into deeper soil layers by plowing. Nash and Gish (1989) studied the volatilization and dissipation of several halogenated pesticides from moist sandy loam soil under controlled conditions in micro-agroecosystem chambers maintained in a greenhouse for 154 days. As soil temperature increased from 5 to 35 $^{\circ}$ C, the percentage of originally applied hexachlorobenzene that was detected in the soil compartment decreased, while the percentage detected in the air increased suggesting that hexachlorobenzene volatilizes more rapidly with increased soil temperature. Reported soil/air partition coefficient values for hexachlorobenzene range from 5.0 to 7.3, signifying that, at equilibrium, soil will contain a much greater mass of hexachlorobenzene than air (Barber et al. 2005).

6. POTENTIAL FOR HUMAN EXPOSURE

The high octanol/water partition coefficient (K_{ow}) value (Hansch et al. 1995) for hexachlorobenzene (see Table 4-2) suggests that bioconcentration and biomagnification of hexachlorobenzene are likely to occur to a significant degree. Veith et al. (1979) measured bioconcentration factor (BCF) values of 16,200 for fathead minnows, 21,900 for green sunfish, and 5,500 for rainbow trout exposed to hexachlorobenzene at 15 °C for 32 days. Oliver and Niimi (1983) studied bioconcentration in rainbow trout exposed to water containing 2 concentrations of hexachlorobenzene (0.32 and 8 ng/g [nph]) for 119 and 105 days

containing 2 concentrations of hexachlorobenzene (0.32 and 8 ng/g [ppb]) for 119 and 105 days, respectively. The BCF values were 12,000 and 20,000 at the 0.32 and 8 ng/g (ppb) exposure levels, respectively. Chaisuksant et al. (1997) conducted a bioconcentration experiment using mosquito fish (*Gambusia affinis*) as well. The fish were exposed to three concentrations of eight chemicals, and the highest concentration used consisted of a mixture with each chemical present in a concentration equal to 1/20 of the LC₅₀. After 96 hours of exposure, the BCF of hexachlorobenzene in mosquito fish was 3,730. After an 8-week exposure period of carp (*Cyprinus carpio*) to concentrations of 0.5 and 0.05 µg/L of hexachlorobenzene, the BCFs were 11,000–27,000 and 6,000–30,000, respectively (NITE 2010).

In a model aquatic ecosystem to which hexachlorobenzene was introduced, the BCF averaged 740 for algae (Oedogonium cardiacum), 1,500 for the snail (Helisoma sp.), 910 for the daphnid (Daphnia magna), 1,610 for the mosquitofish (G. affinis), and 10,610 for the catfish (Ictalurus punctatus) (Isensee et al. 1976). The authors concluded that biomagnification was also occurring within the food chain because the catfish (highest trophic level species) accumulated over 10 times more hexachlorobenzene than the next highest trophic level (snails and mosquitofish), and these species accumulated 1.5–2 times more than the lowest food chain species, the daphnids (primary consumers) and the algae (primary producers). In studies of natural populations of white bass in Lake Erie, Russell et al. (1995) concluded that biomagnification of hexachlorobenzene did not occur. These authors did report biomagnification in Lake Erie fish populations was occurring for several other organic chemicals with log Kow values greater than 6.1. Hexachlorobenzene bioaccumulation factors (BAFs) in aquatic fish species has been measured by Burkhard et al. (1997) in the Bayou d'Inde of the Calcasieu River system near Lake Charles, Louisiana. This field study resulted in log BAF values of 5.80 for blue crab (*Callinectes Sapidus*), 6.03 for mummichog fish (Fundulus Heteroclitus), 6.30 for Atlantic croaker (Micropoganias undulatus), and 6.68 for gulf menhaden (Brevoortia Patronus). The author further compared the measured values obtained to previously reported and predicted BAF values. A comparison of these data with that of Pereria et al. (1988) reveals a difference that was not considered significant by the author. Pereria et al. (1988) determined log BAF values of 4.03 for blue crab, 4.56 for Atlantic croaker, 4.12 for spotted sea trout (Cynoscion nebulosis), and 4.61 for blue catfish (Ictalurus furcatus). Hexachlorobenzene was

6. POTENTIAL FOR HUMAN EXPOSURE

shown to biomagnify in lake trout food webs in lakes in the northeastern United States using lake trout and other food web organisms collected between 1998 and 2001, as measured by trophic magnification factors (Houde et al. 2008).

Connell et al. (1988), using data derived from terrestrial laboratory microcosm studies with two oligochaete worms (*Limnodrilus hoffmeisteri* and *Tubifex*), suggest that interstitial water may be the source from which lipophilic compounds such as hexachlorobenzene in sediment are bioconcentrated by oligochaetes. The concentration factor was 0.54 for hexachlorobenzene during a 110-day exposure test. In a similar study of the earthworm (*Eisenia andrei*) raised in field-contaminated soil, Belfroid (1995) reported a biota-to-soil accumulation factor of 0.507 for hexachlorobenzene. These authors also noted an initial elimination half-life of 1.9 days followed by a period of slower elimination with a half-life of 47 days. In a terrestrial food web study conducted on the Niagara Peninsula of Ontario, Canada from 1987 to 1989, Hebert et al. (1994) reported concentrations of hexachlorobenzene increased from the lower trophic level species to higher trophic level predator species. Concentrations of hexachlorobenzene were not detected in soil or plant material; however, concentration ranges were $0.2-0.3 \mu g/kg$ (ppb) (wet weight) in earthworms, not detected to $1.0 \mu g/kg$ (ppb) in mammals, $2.0-2.4 \mu g/kg$ (ppb) in starlings, $1.8-2.5 \mu g/kg$ (ppb) in robins, and $2.1-5.1 \mu g/kg$ (ppb) in kestrels at the top of the food web.

Several agricultural species of plants have been shown to bioaccumulate hexachlorobenzene in their roots and in portions of the plant growing closest to the soil, such as low stems (Scheunert et al. 1983; Smelt and Leistra 1974). There were marked differences in the BCFs among the various plant species with higher residues associated with those species with the higher lipid content (Schroll et al. 1994; Smelt and Leistra 1974). The roots of the plants generally accumulate higher concentrations of soil-applied hexachlorobenzene than do the aerial parts of the plants. This has been demonstrated for hexachlorobenzene in sugar beets, carrots, turnips, wheat, and pasture grass (Scheunert et al. 1983; Smelt and Leistra 1974). The edible root portion of carrots accumulated the highest concentration of hexachlorobenzene (1,250 ppb with a plant/soil BCF of 19) for a human food source. The measured BCF for hexachlorobenzene was 210 and 470 in soy bean plants via root uptake from water containing 0.2 and 0.4 μ g/L hexachlorobenzene, respectively (Kraaij and Connell 1997). Concentrations of hexachlorobenzene were also high in grass roots (810 ppb) and the lower (0–5 cm) part of the blade (220 ppb) (Smelt and Leistra 1974). It is assumed that hexachlorobenzene in soil is mobile mainly in the gas phase. Gaseous hexachlorobenzene can diffuse directly into the plant root or evaporated hexachlorobenzene can be taken up by plant foliage (Ecker and Horak 1994). Some studies have reported no marked translocation of the hexachlorobenzene from roots to shoots or vice versa (Schroll et al. 1994). Residues in the roots were associated only with

6. POTENTIAL FOR HUMAN EXPOSURE

root uptake from the soil; those residues in the shoots were only from foliar uptake from the air. Studies by Ecker and Horak (1994), however, suggest that root uptake of hexachlorobenzene by oil pumpkins occurred and that the hexachlorobenzene was translocated into the shoots. These authors report that uptake of dissolved hexachlorobenzene from soil solution into the roots may not have been considered earlier as a source for the translocated compound. Pollutants entering the plant from contaminated soil via roots would be translocated in the plant by the xylem while gas- and particle-phase deposition onto leaves or uptake by the stomata would be translocated by the phloem (Simonich and Hites 1995). Concentrations of hexachlorobenzene in agricultural crops can be directly transferred to humans via direct consumption, while concentrations in grass and other forage crops can be indirectly transferred to humans via consumption of dairy products or meat from cattle grazing on contaminated pastures.

Lichen from Northwestern Ontario and South Central Ontario exhibited BCF values of 1.7×10^7 and 8.8×10^6 , respectively, for hexachlorobenzene. These BCF values were calculated as the concentration of hexachlorobenzene in the lichen (ng/m³ wet weight) compared to the concentration in the air (ng/m³) (Muir et al. 1993). Furthermore, bioconcentration of hexachlorobenzene by lichen, a major forage food for caribou can transfer hexachlorobenzene to recreational hunters and natives peoples that consume caribou in their diets (Elkin and Bethke 1995).

6.3.2 Transformation and Degradation

6.3.2.1 Air

Photodegradation of hexachlorobenzene in its vapor phase, or as an adsorbable on silica gel, has been reported as not occurring when hexachlorobenzene was irradiated with ultraviolet radiation of wavelength 290 nm for 6 days (Parlar 1978); however, production of HCl and CO_2 was observed when hexachlorobenzene was irradiated at 230 nm (Parlar 1978). In the troposphere, hexachlorobenzene is probably photochemically stable, but degradation in the stratosphere by photo-dissociation by shorter-wavelength, higher energy-ultraviolet light may be a mechanism for atmospheric degradation in the stratosphere. Hexachlorobenzene, 20 µg, was degraded completely within 20 minutes when dipped onto a titanium dioxide film surface and irradiated with an ultraviolet radiation high-pressure mercury lamp with a wavelength of 365 nm, demonstrating how nano-titanium dioxide can increase the degradation of hexachlorobenzene under ultraviolet light radiation (Yu et al. 2007).

The photo-oxidation half-life (first-order kinetics) of hexachlorobenzene based on the vapor phase reaction with hydroxyl radicals in air was estimated to range from 156.4 days to 4.2 years by Howard et

6. POTENTIAL FOR HUMAN EXPOSURE

al. (1991) and from 158 days to 4.3 years by Kwok and Atkinson (1995). Wania and Mackay (1995) estimated the degradation half-life (first-order kinetics) of hexachlorobenzene to be 0.63 years (230 days), 1.94 years (708 days), and 6.28 years (2,292 days) in air in tropical/subtropical regions, temperate/boreal regions, and polar regions, respectively. Brubaker and Hites (1998) measured a hydroxyl rate constant of 2.7×10^{-14} cm³/molecule-second at 25 °C, corresponding to a calculated half-life of 1.69 years. Thus, atmospheric degradation is extremely slow and is not an efficient method of hexachlorobenzene removal.

6.3.2.2 Water

Hexachlorobenzene is a persistent compound and is not significantly degraded by either abiotic or biodegradation processes in water. It is resistant to the types of hydrolysis reactions that can help degrade other organochlorines or organophosphates, and it is not markedly subject to photolytic decay (Mill and Haag 1986). Biodegradation of organic priority pollutants in a waste water inoculum system amended with yeast extracts was studied by Tabak et al. (1981). Among the 57 environmental pollutants tested, hexachlorobenzene at concentrations of 5 and 10 ppm was among the more slowly biodegraded compounds tested. Biodegradation of hexachlorobenzene in waste water treatment systems is expected to be slow.

An aquatic ecosystem study conducted by Schauerte et al. (1982) shows that hexachlorobenzene will mainly absorb onto particulate matter in the water and then be transported to the bottom sediment. After 145 weeks, the study found a significant amount (10–20%) of hexachlorobenzene remaining in the upper sediment layers (0–10 cm). The half-life (first-order kinetics) of hexachlorobenzene was estimated to range from 2.7 to 5.7 years in surface water and from 5.3 to 11.4 years in groundwater based on unacclimated aqueous aerobic biodegradation (Howard et al. 1991).

Hirsch and Hutzinger (1989) conducted surface water photolysis test with hexachlorobenzene in a laboratory setting and found that this process may occur. A first order rate constant (1.3x10⁻⁶/sec) corresponding to a half-life of 6.17 days for the photolysis of hexachlorobenzene in distilled water in a photochemical reactor equipped with mercury arc lamps was reported. Hexachlorobenzene in an acetonitrile:water mixture exposed to wavelengths of 290 nm for 8 hours resulted in a 33.5% loss of hexachlorobenzene. 1,2,3,4,5-Pentachlorobenzene (76.8%), 1,2,3,5-tetrachlorobenzene (1.2%), 1,2,4,5-tetrachlorobenzene (1.7%), and 1,2,4-trichlorobenzene (0.2%) were found as hexachlorobenzene transformation products (Choudhry et al. 1986). In another experiment, hexachlorobenzene in a water:acetonitrile solution was exposed to sunlight and resulted in a half-life of 70 days (Mill and Haag

1986). The studies above found photolysis of hexachlorobenzene a feasible loss process with half-lives ranging from 6.17 to 70 days.

Hydrolysis is not expected to be an important fate process. EPA (1987) observed zero hydrolysis after 13 days for pH values of 3, 7, and 11 at 85 °C.

Hexachlorobenzene can also be eliminated by ozone reactions. Roche and Prados (1995) conducted a study and compared the efficiencies of ozone and ozone-hydrogen peroxide systems in removing hexa-chlorobenzene from water treatment processes. The concentration of ozone during the experiments was 70 mg O_3/L . When ozone was applied, 11–14% of an initial concentration of 1.0 µg/L hexachlorobenzene was removed. This removal increased to 15–48% when hydrogen peroxide and ozone were applied together.

6.3.2.3 Sediment and Soil

Hexachlorobenzene is a persistent compound and is not significantly degraded in soils by either abiotic or biodegradation processes. In a year-long laboratory study, soil treated with 0.1, 1.0, and 10 ppm of hexachlorobenzene was stored under aerobic (sterile and nonsterile) conditions and under anaerobic nonsterile conditions in covered containers to retard hexachlorobenzene volatilization (Isensee et al. 1976). No loss in the soil-incorporated hexachlorobenzene was observed at any treatment concentration or under any storage condition. Beck and Hansen (1974) measured a half-life (first-order kinetics) of 3-6 years for hexachlorobenzene in soils. Anaerobic biological dechlorination of hexachlorobenzene has also been demonstrated in anaerobic sewage sludge (Fathepure et al. 1988). These authors reported that hexachlorobenzene was dechlorinated to tri- and dichlorobenzenes under anaerobic conditions when sewage sludge was maintained in serum bottles and incubated in the laboratory. Complete biotransformation of a 50 ppm inoculum occurred within 3 weeks. Two routes of dechlorination were observed. The major route was hexachlorobenzene \rightarrow pentachlorobenzene \rightarrow 1,2,3,5-tetrachlorobenzene \rightarrow 1,3,5-trichlorobenzene; the minor route was hexachlorobenzene \rightarrow pentachlorobenzene $\rightarrow 1,2,4,5$ -tetrachlorobenzene \rightarrow 1,2,4-trichlorobenzene \rightarrow dichlorobenzenes (i.e., 1,2-, 1,3-, and 1,4-dichlorobenzene). Yuan et al. (1999) also conducted an anaerobic biological dechlorination study using sewage sludge obtained from the Di-Hua Municipal Sewage Treatment Plant in Taipei, Japan. All experiments were performed using 25 mL serum bottles containing 9 mL of sewage sludge and various concentrations of hexachlorobenzene. After a 20-day incubation period, 98% of the 2 mg/L hexachlorobenzene remained, while addition of 1,2,3-trichlorobenzene adapted consortium accelerated dechlorination which occurred at a

6. POTENTIAL FOR HUMAN EXPOSURE

calculated rate of 0.29 mg/L/day. At hexachlorobenzene concentrations of 2, 5, and 10 mg/L, complete dechlorination occurred within 6 days and at the 50 mg/L concentration, dechlorination occurred in 8 days. Optimal dechlorination occurred at a rate of 0.29 mg/L/day, pH of 7.0, and 30 °C. According to Yuan et al. (1999), dechlorination occurred via the following path: hexachlorobenzene \rightarrow pentachlorobenzene \rightarrow 1,2,3,4-tetrachlorobenzene + 1,2,3,5-tetrachlorobenzene \rightarrow 1,2,4-trichlorobenzene + 1,2,3-trichlorobenzene + 1,2-dichlorobenzene + 1,4-dichlorobenzene. From this and other studies, it is clear that in a time frame of days to years, anaerobic biodegradation may remove hexachlorobenzene from soils.

Under aerobic conditions, Uhlik et al. (2014) reported detection of bacterial populations capable of deriving carbon from hexachlorobenzene. The bacteria populations were primarily Proteobacteria, including *Methylobacterium* and *Pseudomonas* collected from contaminated soils at a chemical factory in Neratovice, Czech Republic; however, the exact biodegradation pathways and degradation rates were not determined. Additionally, hexachlorobenzene, present at 100 mg/L in an aerobic sludge study to test biodegradation, reached 0% of its theoretical biological oxygen demand (BOD) in 2 weeks using an activated sludge inoculum at 30 mg/L and the Japanese Ministry of International Trade and Industry (MITI) test (NITE 2010); thus, aerobic degradation is not an important fate process. Contaminated soil from Klatovy-Luby, Hajek, and Neratovice, Czech Republic contained bacterial strain, *Stenotrophomonas sp.*, capable of degrading hexachlorobenzene (Lovecka et al. 2014). A 34.9% decrease of the original concentration of hexachlorobenzene was detected in the sample of isolate H1D7.

Likewise, in areas of the Great Lakes region with a long history of hexachlorobenzene contaminated waste water discharges affecting aquatic sediments, the concentrations of hexachlorobenzene in the sediments can be significant (see Section 6.4.2). Susarla et al. (1997) examined the transformation of hexachlorobenzene in fresh water lake (Lake Kasumigaura, Japan) sediments under anaerobic conditions. Dechlorination occurred after a 4-day lag phase and was complete in 32 days. The calculated rate of dechlorination was 0.110/day. Hexachlorobenzene transformation pathway under sulfidogenic conditions resulted in hexachlorobenzene \rightarrow pentachlorobenzene \rightarrow 1,2,3,5-tetrachlorobenzene \rightarrow 1,3,5-tri-chlorobenzene \rightarrow 1,3-dichlorobenzene. Under methanogenic conditions the pathway was as follows: hexachlorobenzene \rightarrow pentachlorobenzene \rightarrow 1,2,3,4-tetrachlorobenzene \rightarrow 1,2,4-trichlorobenzene \rightarrow 1,4-dichlorobenzene (Susarla et al. 1997). In another experiment, dechlorination of hexachlorobenzene in an estuary sediment collected from the mouth of Tsurumi river occurred at a rate of 0.0256/day with a

half-life of 27.1 days (Masunaga et al. 1996). Thus, aquatic sediment degradation of hexachlorobenzene occurs in a month to a year.

6.3.2.4 Other Media

The explosion and collapse of the World Trade Center (WTC) on September 11, 2001 produced aerosols that dispersed into the environment. Two bulk samples of the dust that settled at indoor locations surrounding the WTC site contained mean hexachlorobenzene concentrations of 2.14 and 2.66 ng/g. Six bulk samples of dust from various outdoor locations around the WTC site contained an average hexachlorobenzene concentration of 1.3 ng/g (Offenberg et al. 2004).

6.4 LEVELS MONITORED OR ESTIMATED IN THE ENVIRONMENT

Reliable evaluation of the potential for human exposure to hexachlorobenzene depends in part on the reliability of supporting analytical data from environmental samples and biological specimens. Concentrations of hexachlorobenzene in unpolluted atmospheres and in pristine surface waters are often so low as to be near the limits of current analytical methods. In reviewing data on hexachlorobenzene levels monitored or estimated in the environment, it should also be noted that the amount of chemical identified analytically is not necessarily equivalent to the amount that is bioavailable. The analytical methods available for monitoring hexachlorobenzene in a variety of environmental media are detailed in Chapter 7.

6.4.1 Air

Due to its long persistence, hexachlorobenzene is subject to long-range transport in the atmosphere and can be detected in nonsource air samples including rural and remote locations. Hexachlorobenzene measured in air in Villeroy, Quebec in 1992 had mean and median concentrations of 36.68 (0.03668 ng/m³) and 30.94 pg/m³ (0.03094 ng/m³), respectively, from 56 air samples (Poissant et al. 1997). A meteorological station located in a semirural area outside Lancaster, England was the site of air samples. Four air samples per day (taken at 6-hour intervals) were taken for 7 days. The minimum, maximum, and mean concentrations of hexachlorobenzene in these samples were <28.8, 76.1, and 39.3 pg/m³, respectively (Lee et al. 2000b). Air samples taken along a North-South Atlantic transect during five legs in transit from the island of Texel in the Netherlands to Walvis Bay, Namibia and Cape Town, South Africa contained hexachlorobenzene concentrations of 145, 96, not detected, <0.1, and 0.1 pg/m³ for each leg of the trip, respectively (Booij et al. 2007). Huang et al. (2014) evaluated air and water concentrations

6. POTENTIAL FOR HUMAN EXPOSURE

of hexachlorobenzene in samples collected from the Indian Ocean. Hexachlorobenzene concentrations in the air ranged from 2.2 to 17 pg/m^3 and the average concentration was 6.2 pg/m^3 .

Eisenreich et al. (1981) reported atmospheric concentrations of hexachlorobenzene in the Great Lakes region ranging from 0.09 to 0.28 ng/m³. Results of airborne samples collected between 1990 and 1993 from the Great Lakes region by the Integrated Atmospheric Deposition Network are provided by Hoff et al. (1996). The annual mean gas-phase and particulate-phase concentrations of hexachlorobenzene were 98 and 0.2 pg/m³, respectively, in samples from Lake Superior near Eagle Harbor, Michigan, 120 and 0.1 pg/m³, respectively, in samples from Lake Michigan near Sleeping Bear Dunes, Michigan, 80 and 0.2 pg/m³, respectively, in samples from Lake Erie near Sturgeon Point, New York, and 130 and <0.1 pg/m³, respectively, in samples from Lake Ontario near Point Petre, Ontario. From July 1988 to September 1989, 143 air samples were collected at Egbert, Ontario, Canada and were analyzed for PCB and organochlorine concentrations. Hexachlorobenzene was detected at concentrations ranging from a minimum of 0.04 pg/m³ (0.00004 ng/m³) to a maximum of 640 pg/m³ (0.64 ng/m³) (annual mean >54 pg/m³ (0.054 ng/m³) (Hoff et al. 1992).

The variability in hexachlorobenzene air concentrations has decreased as emissions from point sources have declined due to the banning of hexachlorobenzene from agricultural use in the 1970s. Regionally, levels of hexachlorobenzene in the atmosphere are becoming uniform (Barber et al. 2005). Studies of hexachlorobenzene concentrations at a number of sites within a region tend to show little variation. For example, a passive sampling survey in 2002 showed that hexachlorobenzene concentrations varied by less than a factor of 6 over the continent of Europe (Barber et al. 2005). Hexachlorobenzene air concentrations measured along a latitudinal transect of remote/rural sites from the south of the United Kingdom to the north of Norway during 1998 to 2000 indicated that the differences between highest and lowest concentrations were only a factor of about 3 (Meijer et al. 2003a). Hexachlorobenzene concentrations showed a relatively uniform global distribution during a study under the Global Atmospheric Passive Sampling Network, which employed passive samplers at 20 sites globally for a 3-month spring sampling period during 2009 (Koblizkova et al. 2012). Hexachlorobenzene concentrations ranged from 37 to 240 pg/m³.

Hexachlorobenzene air concentrations have been measured in urban and rural areas in France. Atmospheric fallout from the urban area, Paris, and the rural area, La Ferté-sous-Jouarre, was collected in raw form as bulk precipitation. Hexachlorobenzene concentration in rural fallout measured in February– July 1992 and January –September 1993 ranged from 2.5 to 4.5 ng/L and from 0.3 to 4 ng/L, respectively.

6. POTENTIAL FOR HUMAN EXPOSURE

For the same time periods, urban fallout measured 1.8–17 and 0.3–5.6 ng/L, respectively (Chevreuil et al. 1996). The mean concentrations of hexachlorobenzene in precipitation samples collected in the Great Lakes region from 1986 to 1991 ranged from 0.145 ng/L (ppt) at Sibley Park on Lake Superior, to 0.108 ng/L (ppt) at Pelee Island in Lake Erie, to 0.174 ng/L (ppt) at Wolfe Island in Lake Ontario (Chan et al. 1994). The mean and median concentrations of hexachlorobenzene from eight precipitation samples collected from Villeroy, Quebec in 1992 were 0.04 and 0.05 ng/L, respectively (Poissant et al. 1997). Precipitation samples collected between 1990 and 1993 from the Great Lakes region by the Integrated Atmospheric Deposition Network were analyzed. The annual mean concentration of hexachlorobenzene in these precipitation samples were 0.1 ng/L in samples from Lake Superior near Eagle Harbor, Michigan, 0.06 ng/L in samples from Lake Michigan near Sleeping Bear Dunes, Michigan, 0.04 ng/L in samples from Lake Erie near Sturgeon Point, New York, and 0.3 ng/L in samples from Lake Ontario near Point Petre, Ontario (Hoff et al. 1996). The concentrations of hexachlorobenzene in air, particulate matter, and rain from Galveston Bay, Texas were 87.3±103.3, 0.4±0.5, and 42.5 pg/m³ (not detected–48.1 pg/m³), respectively, between 1995 and 1996 (Park et al. 2001). The median levels of hexachlorobenzene in ambient air samples collected in Zagreb, Croatia in 1997 were 29 pg/m^3 (range, 0.5–49 pg/m^3) in the northern residential region of Ksaverska and 31 pg/m³ (range, 15–61 pg/m³) in the southern region near a landfill (Romanic and Krauthacker 2000). The average concentration of hexachlorobenzene in air at Lake Malawi, in southeast Africa, from February 1997 to May 1998, was 11 ± 7.5 pg/m³ (Karlsson et al. 2000). Seasonal snowpack samples collected in spring 2003 from six national parks in the western United States contained mean hexachlorobenzene concentrations of 0.0065, 0.015, 0.017, and 0.035 ng/L for the four contiguous national parks, and 0.0055 and 0.0077 ng/L for the national parks in Alaska (Hageman et al. 2006). Average concentrations of hexachlorobenzene in the air samples collected from cities in China have been reported. Seasonal variation was evaluated and the concentrations of hexachlorobenzene detected were 91.2, 49.7, 85.4, and 420 pg/m³ in Suzhou, 241, 22.6, 218, and 468 pg/m³ in Wuxi, and 123, 46.5, 102, and 257 pg/m³ Nantong for spring, summer, autumn, and winter, respectively (Zhang et al. 2013). In Beijing, China, average concentrations from samples of urban atmosphere were 200, 68, 180, and 400 pg/m³ for spring, summer, autumn, and winter, respectively (Zhang et al. 2011).

Nonoccupational exposure to hexachlorobenzene for residents of two U.S. cities (Jacksonville, Florida and Springfield, Massachusetts) was studied over three seasons: summer 1986, spring 1987, and winter 1988 (Whitmore et al. 1994). The study focused primarily on inhalation exposures with primary environmental monitoring consisting of 24-hour indoor and outdoor air. For the Jacksonville, Florida, population, the estimated mean hexachlorobenzene concentration ranged from 0.3 to 1.3 ng/m³ for indoor air and from not detected to 0.2 ng/m³ for outdoor air. For the Springfield, Massachusetts population,

6. POTENTIAL FOR HUMAN EXPOSURE

mean exposures were much less. The estimated hexachlorobenzene concentrations ranged from not detected to 0.1 ng/m^3 for indoor air and no detectable concentrations of hexachlorobenzene for outdoor air.

Extremely high concentrations of hexachlorobenzene in air have been detected in areas close to production and disposal sites in both outdoor and indoor air. Mann et al. (1974) measured hexachlorobenzene concentrations ranging from 70 to 23,296 ng/m³ near chlorinated solvent and pesticide manufacturing facilities; air levels near a chemical waste landfill were as high as 16,000 ng/m³ (EPA 1975b). Hexachlorobenzene has been detected at 11,000 ng/m³ in flue gas effluents from a municipal refuse-fired steam boiler in Virginia, and at 9.5 ng/m³ in flue gas effluents from a refuse-derived fuel fired power plant in Ohio (Tiernan et al. 1985). Air concentrations of hexachlorobenzene inside industrial plants can be as high as 150,000 ng/m³ (Currier et al. 1980); air concentrations inside a pesticide production facility were measured at 22,000 ng/m³ (Davis and Morgan 1986).

Due to the moderate volatility and extremely slow atmospheric degradation of hexachlorobenzene, evaporation for environmental reservoirs can sustain atmospheric levels of hexachlorobenzene within an order of magnitude of those resulting from primary emissions, which may explain why atmospheric concentrations have remained steady decades after its main primary emissions have been largely eliminated (Choi and Wania 2011).

6.4.2 Water

Drinking water in three cities in the Lake Ontario vicinity contained hexachlorobenzene ranging from 0.06 to 0.2 ppt (mean of 0.1 ppt), about the same as water from the lake (Oliver and Nicol 1982a).

Hexachlorobenzene was detected in ambient water samples from two of the Great Lakes and their tributary rivers. Mean concentrations of hexachlorobenzene in Lake Ontario, Lake Huron, and the Grand River were 0.06 ppt (range, 0.02–0.1 ppt), 0.04 ppt (range, 0.02–0.1 ppt), and 0.06 ppt (range, 0.02–0.1 ppt), respectively. In the Niagara River, concentrations of 0.02–17 ppt were detected with the highest value measured downstream of a waste disposal site (Oliver and Nicol 1982a). Widely varying measurements in this river may be attributed to the fact that measurements were near the analytical detection limit of 0.01 ppt. Hexachlorobenzene was detected in 42% of whole water samples (dissolved plus particulate phases) collected during 1991 in the St. Lawrence River and several of its tributaries. Hexachlorobenzene concentrations detected in the St. Lawrence River ranged from not detected to

6. POTENTIAL FOR HUMAN EXPOSURE

0.09 ng/L (mean 0.01 ng/L [ppt]) (Quemerais et al. 1994). Hoff et al. (1996) obtained and presented water concentration data from a 1992 sampling study. Hexachlorobenzene concentrations in Lakes Superior, Michigan, Huron, Erie, and Ontario were 0.01, 0.014, 0.007, 0.014, and 0.045 ng/L, respectively. In remote European mountain lake waters, the concentrations of hexachlorobenzene were $8.4\pm11 \text{ pg/L}$ at Redó, Spain, $4.0\pm1.8 \text{ pg/L}$ at Gossenkölle, Austria, and $6.2\pm1.0 \text{ pg/L}$ at Øvre Neådalsvatn, Norway (Vilanova et al. 2001).

The concentrations of hexachlorobenzene in microlayer and subsurface Mediterranean seawater off the coast of Alexandria, Egypt were on average, 27.3±17 and 12±6.9 ng/L, respectively (Abd-Allah 1999). Seawater samples taken along a North-South Atlantic transect during five legs in transit from the island of Texel in the Netherlands to Walvis Bay, Namibia and Cape Town, South Africa contained hexachlorobenzene concentrations of 9.0, 6.3, 1.9, 3.3, and 2.7 pg/L for each leg of the trip, respectively (Booij et al. 2007). Hexachlorobenzene has been detected in shallow groundwater in China at depths of 4–8 m. Monitoring wells and drinking wells sampled in the Taihu Lake region of China in 2011 had an average hexachlorobenzene concentration of 16.53 ng/L and a range of 3.05–65.24 ng/L (Wu et al. 2014).

A study was conducted from 1974 to 1975 to collect and analyze surface water samples from sites of known hexachlorobenzene contamination along the Mississippi River near an industrial area in Geismar, Louisiana (EPA 1976a). The maximum hexachlorobenzene concentration detected in water was 90.3 ppb. A concentration of 2 ppb has been measured in the Mississippi River near Baton Rouge, Louisiana (Laska et al. 1976).

Industrial waste water samples contained hexachlorobenzene levels as high as 300 ppb (EPA 1976b; Schmitt et al. 1990). Effluent concentrations of hexachlorobenzene from four Canadian plants in the Great Lakes region ranged from 0.001 to 0.002 ppb (0.0015 ppb mean) (Oliver and Nicol 1982a).

Hexachlorobenzene concentrations were measured in water at an uncontrolled hazardous waste site near Bayou Baton Rouge, Louisiana (Davis and Morgan 1986). Surface water samples collected from a containment pond used for disposal of wastes from both rubber production and manufacture of chlorinated organics at the site contained up to 8,100,000 ppb hexachlorobenzene.

6.4.3 Sediment and Soil

Mean concentrations of hexachlorobenzene in lake sediments in the Great Lakes ranged from 0.2 to 97 ppb with the highest values measured in Lake Ontario. Deeper sediment layers (1-2 cm) had even higher concentrations of hexachlorobenzene (460 ppb) than surface (0-1 cm) samples (270 ppb), with the peak value corresponding to deposition in the years 1971–1976 declining to 270 ppb in 1976–1980 (Oliver and Nicol 1982a). A study of hexachlorobenzene concentration in Lake Erie sediments revealed a decrease of roughly 80%, with lake-wide average measured concentrations of 5.7 and 1.6 ng/g dry weight in 1971 and 1997, respectively (Marvin et al. 2004). In 1992, 2 cm deep bed sediment samples were collected from the South Platte River at Henderson, Colorado and Cache La Poudre River near Greeley, Colorado. The sediment contained 1.5 and <1 μ g/kg, dry weight, hexachlorobenzene, respectively. The authors concluded that this concentration was correlated to the hexachlorobenzene concentrations found in urban and agricultural lands in the South Platte River Basin (Tate and Heiny 1996). Outside of the United States, 12 sediment samples were collected in June 1993 near known discharges from municipalities and industries from Lake Ladoga, Russia. Hexachlorobenzene concentrations were 3.58 and 14.6 ng/g in 2 out of the 12 samples, and hexachlorobenzene was not detected in the remaining 10 samples (Ristola et al. 1996). In the levees of the Mississippi River near Baton Rouge, where the river water contained 2 ppb hexachlorobenzene, the soil contained 167 ppb (Laska et al. 1976).

Meijer et al. (2003b) studied hexachlorobenzene concentrations in background surface soils collected in 1998 from 191 locations around the world. The reported global mean background concentration was 0.68 ng/g dry weight. The lowest measured concentration, 0.010 ng/g dry weight, was found in samples from Bear Island in Norway, and the highest concentrations, 5.20 and 4.83 ng/g dry weight, were found in south Norway and Russia, respectively. This study also demonstrated how hexachlorobenzene concentration is directly correlated with soil organic matter (SOM) content. The highest hexachlorobenzene concentrations were measured in the northern hemisphere, in general, and in Europe, in particular. This corresponds with the location of soils with the highest SOM content, associated with forests, peat bogs, and grassland systems.

Yuan et al. (2014) evaluated hexachlorobenzene concentrations from soils collected in 2011 from the Tibetan Plateau. The soil samples were collected at locations with low soil organic carbon (SOC) content and >200 km from possible sources at high altitudes, where soil-air exchange processes are expected to greatly influence adsorption and re-volatilization of hexachlorobenzene. The average hexachlorobenzene concentration was 54.0 ng/kg. Hexachlorobenzene was detected in surface soil samples from agriculture

6. POTENTIAL FOR HUMAN EXPOSURE

soil used to grow rice, wheat, corn, bean, cotton, and vegetables in central China. The average concentration of hexachlorobenzene detected was 3.01 ng/g soil, based on a range of concentrations from not detected to 17.77 ng/g soil in 44 soil samples; the detection frequency was 86% (Zhou 2013). In Tianjin, China, hexachlorobenzene concentrations ranged from not detected to $1,924 \mu \text{g/kg}$ dry weight near the Tanggu Chemical Industrial District (Hou et al. 2013).

Hexachlorobenzene concentrations were measured in soil and sediment at several uncontrolled hazardous waste sites in several states (Davis and Morgan 1986). Hexachlorobenzene concentrations of up to 20,000 ppb were measured in soil at a scenic highway site near Bayou Baton Rouge, Louisiana, while concentrations in sediment of 39,500 ppb were measured from a bayou bank downstream of the site. Soil cores from a monitoring well (25–27 feet deep) were as high as 400,000 ppb, and as high as 90,000 ppb in soils collected at 40–41 feet deep (Davis and Morgan 1986). Soil and sediment collected from a disposal site near Sorrento, Louisiana contained 62,000 and 130,000 ppb hexachlorobenzene, respectively. Soil collected at a Crystal City, Texas pesticide disposal site was found to contain 20,000 ppb hexachlorobenzene. A maximum hexachlorobenzene concentration detected in soil at an industrial site of known contamination near Geismar, Louisiana was 53,130 ppb (Laseter et al. 1976).

Sediment concentrations of hexachlorobenzene vary widely from relatively unpolluted areas to those areas used extensively for disposal of hexachlorobenzene-containing wastes. Sediment hexachlorobenzene concentrations from San Luis Pass, located near industrial areas of West Galveston Bay, Texas ranged from 0.05 to 1.5 ppb (dry weight) with a mean of 0.49 ppb (Murray et al. 1981).

Hexachlorobenzene concentrations in marine sediment samples collected from an industrialized area of the harbor in Portland, Maine ranged from <0.03 to 0.37 ppb (mean 0.14 ppb) (Ray et al. 1983). Concentrations of hexachlorobenzene in sediment from the Niagara River watershed in the vicinity of several hazardous waste disposal areas ranged from 8,000 to 30,000 ppb (Elder et al. 1981). The average concentration of hexachlorobenzene in surficial sediments of the Kaohsiung coast (southwestern Taiwan), which receives wastewater from the largest industrial city in Taiwan (Kaohsiung City), ranged from 1.7 to 24.7 ng/g (Lee et al. 2000a). In Northern China, the average concentrations of hexachlorobenzene in surficial sediments of the Laizhou Bay ranged from below the limit of detection to 0.17 ng/g in marine sediment and from below the limit of detection to 14 ng/g in the adjacent river riverine sediment (Zhong et al. 2011). Concentrations in sediment from the Xiangjiang River in China ranged from 0.6 to 14.2 ng hexachlorobenzene/g sediment (Li et al. 2013). Surface sediments from Muroran Port, Japan were evaluated for hexachlorobenzene; concentrations ranged from 110 to 25,000 pg/g dry weight (Anezaki

and Nagahora 2014). Hexachlorobenzene was detected in Lenga Estuary, Chile at concentrations of 1–870 ng/g dry weight (Pozo et al. 2014).

6.4.4 Other Environmental Media

Concentrations of hexachlorobenzene have been detected in several species of fish and shellfish. Hexachlorobenzene concentrations were determined for several species of marine organisms collected from San Luis Pass near Galveston Bay, Texas (Murray et al. 1981). Mean concentrations of hexachlorobenzene at were 0.49 ppb wet weight for flounder (species unspecified), 0.65 ppb for longnose killifish (Fundulus similis), 0.88 ppb for brown shrimp (Penaeus aztecus), 9.6 ppb for blue crab (Callinectes sapidus), and 0.71 ppb for the dwarf squid (Lollingnucula brevis). Oysters (Crassotrea virginica) collected at the lower end of the Houston Ship Channel were found to contain hexachlorobenzene concentrations ranging from 0.31 to 1.41 ppb with a mean of 0.63 ± 0.39 ppb (Murray et al. 1980). The Sheboygan River in Wisconsin is another area of concern for contamination of organochlorine pesticides in fish due to the existence of wetlands, urban and developed land, woodland, and agricultural land surrounding this river. In addition, the lower segment of the river has a history of shipping, industrial and municipal activities, and dredging, including the existence of a landfill designated as a federal superfund site. Schrank et al. (1997) collected white suckers (Catostomus commersoni) from two sites of the Sheboygan River; one site was 1 to 2 km from the mouth of the river and the other was 50 km from the mouth of the river, which served as the reference site. The fish collected from both sites contained less than detectable residues of hexachlorobenzene along with other organochlorine compounds, thus indicating a minimized risk of exposure to hexachlorobenzene from this river. Hexachlorobenzene was detected in whole fish composites collected from bass and carp from 13 river sites collected in the Mobile, Apalachicola-Flint-Chattahoochee, Savannah, and Pee Dee River Basins at mean concentrations of 0.66 and 2.48, 0.81 and 2.38, 1.03 and 2.73, and 0.37 and 0.43 ng/g wet weight, respectively (Hinck et al. 2008).

DeVault (1985) reported concentrations of hexachlorobenzene in whole fish composites collected from the Great Lakes during 1980 and 1981 ranged from <0.002 to 3.47 mg/kg (<2–347 ppb). Hexachlorobenzene concentrations in fresh water trout (4–6+ years old) from the Great Lakes region ranged from 8 to 127 ppb with the highest concentration found in a fish collected near the discharge of the Niagara river into Lake Ontario (Oliver and Nichol 1982a). In another study of Great Lakes fish species, Newsome and Andrews (1993) reported hexachlorobenzene concentrations in fish fillet composites ranged from 0.22 ng/g (ppb) in bullhead to 9.05 ng/g (ppb) wet weight in trout in lake areas with open

6. POTENTIAL FOR HUMAN EXPOSURE

fisheries. Zabik et al. (1995) reported that skin-off processing and selected cooking methods reduced hexachlorobenzene residues in chinook salmon and carp harvested from the Great Lakes. Concentrations of hexachlorobenzene averaged 0.017 ppm (17 ppb) and 0.011 ppm (11 ppb) (wet weight) in raw and cooked salmon fillets, respectively, and averaged 0.005 ppm (5 ppb) and 0.003 ppm (3 ppb) in skin-on and skin-off fillets, respectively. The average percentage loss of hexachlorobenzene from chinook salmon fillets by baking, charbroiling, and canning was 40%. Losses of hexachlorobenzene residues from carp fillets were slightly greater than 40% (Zabik et al. 1995). Walleye, siscowet, carp, and whitefish were collected for organochlorine pesticides analysis from Lake Superior along the Apostle Islands region during 1991 and 1992. Walleye and carp had hexachlorobenzene concentrations below the limit of quantification, while siscowet and whitefish measured concentrations of 3.2 and 2.8 ng/g wet weight of tissue, respectively (Gerstenberger et al. 1997). Organisms sampled during the summer of 1994 from the Keweenaw Peninsula of Lake Superior contained measured hexachlorobenzene concentrations ranging from 0.8 to 1.8 ng/g wet weight in smelts, 3.0–4.3 ng/g wet weight in herrings, 4.7–8.4 ng/g wet weight in bloaters, 1.1–4.1 ng/g wet weight sculpins, <0.1–0.2 ng/g wet weight in mysis, 0.8–1.4 ng/g wet weight in limnocalanus, 0.8 ng/g wet weight in amphipods, and 1.7–3.1 ng/g wet weight in lake trout (Kuchklick and Baker 1998). Grayling and lake trout were collected from four lakes (Feniak, Desperation, Schrader, and Elusive) from the foothills of the Brooks Range, Alaska. All lakes were oligotrophic, and the pH was neutral to slightly alkaline. Fifty-six grayling liver samples and 39 grayling muscle samples were analyzed for hexachlorobenzene concentration, and mean and median values were derived. In the 56 grayling liver samples, the mean and median concentrations were 0.65 and 0.48 ng/g dry weight, respectively. The 39 grayling muscle samples had mean and median concentrations of 0.33 and 0.22 ng/gdry weight, respectively. In lake trout, the mean and median concentrations in 33 liver samples were 1.15 and 0.87 ng/g dry weight, respectively, and in 34 muscle samples were 0.46 and 0.26 ng/g dry weight, respectively (Allen-Gil et al. 1997). Hexachlorobenzene concentrations in sea organisms from the Barents Sea were as follows (units=ng/g lipid weight): copepods (13.5), euphausids (16.5), amphipods (19.5), polar cod (39 ± 1.7) , and cod (65 ± 7.7) (Borgå et al. 2001). In 1991, the concentrations of hexa-

chlorobenzene in amphipods, isopods, and sculpins from the Bothnian Bay (Baltic Sea) were 340 (n=3), 370 (n=5), and 37 (n=3) ng/g dry weight, respectively (Strandberg et al. 2000). In a study of a set of 136 fish from 14 remote lakes in eight western U.S. National Parks/Preserves sampled between 2003 and 2005, the mean muscle tissue concentration of hexachlorobenzene was 0.26 ng/g whole weight and the range was 0.01-1.3 ng/g whole weight (Ackerman et al. 2008).

The bioaccumulative tendencies of hexachlorobenzene have made it a candidate for monitoring in the U.S. Fish and Wildlife Service National Pesticide Monitoring Program (Schmitt et al. 1990) and the

6. POTENTIAL FOR HUMAN EXPOSURE

National Study of Chemical Residues in Fish which was started in 1986 (NSCRF) (EPA 1992). Maximum hexachlorobenzene tissue concentrations (wet weight) detected in whole fish were 700, 130, 120, and 410 ppb in the 1976–1977, 1978–1979, 1980–1981, and 1984 sampling years, respectively. The geometric mean tissue concentration was 10 ppb for 1976–1977 and <10 ppb for all other sampling years (Schmitt et al. 1990). The highest hexachlorobenzene concentrations in the 1984 sampling period (410 ppb) were detected in whole fish from the Tombigbee River, Alabama in the vicinity of a pesticide production facility where concentrations during all sampling years had been the highest nationally. These national results from the ongoing study conducted by NSCRF show that hexachlorobenzene was detected at 46% of the 362 sites surveyed for fish tissue analysis. The mean hexachlorobenzene concentrations are listed in Table 6-2 (EPA 1992). The Freeport, Texas site is near a pesticide plant and the other four sites are close to a variety of chemical manufacturing plants. The Calcasieu River, Louisiana site is close to a Superfund site involving a variety of organic solvents (EPA 1992).

Hazardous waste dumping during the early 1940s and 1950s contaminated the Devil's Swamp, Louisiana with chlorinated hydrocarbons, which has greatly affected fish species. As fish is an important food source for the community surrounding this area, concentrations of contaminants are of great concern. Levels of hexachlorobenzene in various fish species collected from Devil's Swamp, Louisiana, was compared to those from a control site, Tunica's Swamp, Louisiana. Mean hexachlorobenzene concentrations calculated from concentrations of 13 different edible fish species tissues were 23.65 ng/g compared to 2.0 ng/g calculated from 10 different edible fish species from Tunica's Swamp (Tchounwou et al. 1998).

Hexachlorobenzene has been detected in tissues of various wildlife species throughout North America, but especially in wildlife indigenous to the Great Lakes region. Swift et al. (1993) reported mean concentrations of hexachlorobenzene of 0.02 ppm (20 ppb) wet weight (0.04 ppm [40 ppb] lipid weight basis) and 0 ppm wet weight (0.07 ppm [70 ppb] lipid weight basis) in mesenteric and subcutaneous fat and breast tissue, respectively, of common goldeneye waterfowl wintering in New York state. The detection limit in this study was 0.002 ppm. Gebauer and Weseloh (1993) reported that the geometric mean hexachlorobenzene concentrations of 0.4 and 0.9 μ g/kg (ppb) in muscle tissue in mallard ducks using a contaminated sediment site, and sewage lagoon site, respectively, as a resting and feeding area were significantly greater than levels found in ducks using a natural marsh area. Foley (1992) reported hexachlorobenzene residues in muscle tissues of several species of ducks and geese collected in New

Table 6-2.	Sites with the Five Highest Concentrations of Hexachlorobenzene in
	Fish

Whole-body hexachlorobenzene concentration (ppb; wet weight)	Type of sample (fish)	Location
913	Sea catfish	Brazos River, Freeport, Texas
202	Catfish	Bayou D'Inde, Sulfur, Louisiana
93.7	Carp	Mississippi River, St. Francisville, Louisiana
85.5	White sucker	Quinnipiac River, North Haven, Connecticut
75	Sea catfish	Calcasieu River, Moss Lake, Louisiana

^aFrom the EPA 1992 National Study of Chemical Residues in Fish

6. POTENTIAL FOR HUMAN EXPOSURE

York State in 1983–1984. Statewide residues were 64 ppb (wet weight) for buffleheads, 49 ppb for scaups, 26 ppb for mallards, 20 ppb for black ducks, 6 ppb for wood ducks, and 11 ppb for Canada geese. Hexachlorobenzene was detected at concentrations ranging from 0.002 to 0.022 mg/kg wet weight in liver samples of eight seabird species collected opportunistically from a fishery in the North Pacific Ocean in 1992 (Elliott 2005). Adult sea otters that had died along the coast of California were collected by the U.S. Fish and Wildlife Service and the California Department of Fish and Game. Hexachlorobenzene concentrations in liver, kidney, and brain tissues were 0.74–8, 0.28–2.6, and 0.28–0.74 ng/g wet weight, respectively (Nakata et al. 1998). The mean hexachlorobenzene concentration for 207 wild mink liver tissue samples, collected from 1991 to 1995 in the Northwest Territories, Canada from seven mink communities, ranged from 0.21 to 0.67 ng/g wet weight (Poole et al. 1998). Snail composite samples, without shells, were collected from two lakes (Feniak and Elusive) from the foothills of the Brooks Range, Alaska. All lakes were oligotrophic, and the pH was neutral to slightly alkaline. The mean concentration among these six samples was 0.15 ng/g dry weight with a median of 0.10 ng/g dry weight (Allen-Gil et al. 1997).

Hexachlorobenzene was detected in nestling bald eagle plasma collected in 2003 from four areas in southwestern British Columbia and one site in California at mean concentrations of 0.20, 0.26, 0.35, 0.31, and 0.08 μ g/kg wet weight, for Central Fraser Valley, Lower Fraser Valley, Nanaimo/Crofton area, Barkley Sound, and Santa Catalina Island, respectively (Cesh et al. 2008).

Hexachlorobenzene has also been detected in the eggs of various wildlife species in the Great Lakes region and Canada. Yamashita et al. (1992) reported hexachlorobenzene residues ranges of 8–36 ng/g (ppb) and 18–26 ng/g (ppb) on a wet weight basis in the eggs of the double-crested cormorant and the Caspian tern, respectively, collected during 1988 from the Great Lakes region. Somers et al. (1993) reported geometric mean concentrations of 0.013 µg/g (13 ppb) (wet weight) of hexachlorobenzene in double-crested cormorant eggs collected in southern Alberta, Canada. Elliott and Martin (1994) reported mean hexachlorobenzene concentrations in sharp-shinned hawk eggs in south central Ontario ranging from 0.010 to 0.051 mg/kg (10–51 ppb) from 1986 to 1989. Hexachlorobenzene concentrations in Cooper's hawk eggs ranged from 0.005 to 0.012 mg/kg (5–12 ppb) during the same period. Cobb et al. (1994) reported mean residues of 18.0 ng/g (18 ppb) in the chorio-allantoic membranes removed from great blue heron eggs collected from Puget Sound, Washington. Jarman et al. (1996) conducted an experiment with prairie falcon eggs that were collected from eyries in northern and central California between 1989 and 1991. Addled and unhatched eggs were frozen until chemical analysis. The following are the geometric mean concentrations (in mg/kg wet weight) of hexachlorobenzene at their respective

6. POTENTIAL FOR HUMAN EXPOSURE

sampling sites: Frog/Hand Nest 800; Pig Cyn 17; Crowley Tower 11; Willow Sp. 8.0; Goat Rock 10; Mt. Dome 20; and Mt. Diablo 81. Mean hexachlorobenzene residues in peregrine falcon eggs from Rankin Inlet (Hudson Bay, Canada) were $0.03 \ \mu g/g$ wet weight (n=2; range 0–0.15 $\ \mu g/g$ wet weight) between 1982 and 1986, and 0.030 $\ \mu g/g$ wet weight (n=20; range 0–0.165 $\ \mu g/g$ wet weight) between 1991 and 1994 (Braune et al. 1999).

Hexachlorobenzene residues were also detected in snapping turtle eggs collected from a wetland area on Lake Ontario. Residues in the snapping turtle clutches ranged from 43.9, 16.6, and 20.9 ng/g (ppb) (wet weight) to 494.7, 282.1, and 262.2 ng/g (ppb) (lipid weight) for the first five eggs that were layed, a composite sample of five eggs, and the last five eggs that were layed, respectively (Bishop et al. 1995). Bishop et al. (1996) conducted another study with snapping turtle eggs that were collected from nests at five locations from the Great Lakes Basin in 1990–1991. The eggs were analyzed for hexachlorobenzene and the results were compared to data collected from the same sites in the years 1981, 1984, 1988, and 1989. Based on the results, the hexachlorobenzene concentration was the highest in eggs from Cootes Paradise (170–350 ng/g) and lowest in eggs from Algonquin Park (3–20 ng/g). On the whole, the hexachlorobenzene mean concentration from the five sites in the Great Lakes Basin showed a decrease from the years 1984 to 1990 (Bishop et al. 1996).

Langlois and Langis (1995) reported that the concentration of hexachlorobenzene in the blubber of beluga whales from the St. Lawrence Estuary to Northern Quebec Province ranged from 0.22 to 0.93 mg/kg (220–930 ppb) wet weight. Consumption of blubber and organ meats from these whales by native peoples may constitute a potential health risk if these tissues are a significant part of the diet (Langlois and Langis 1995). In 1997, Gauthier et al. (1997) analyzed blubber biopsies from Northwestern Atlantic Balaenopterid whales summering in the Gulf of St. Lawrence. Samples were collected in the summer and fall of 1991 and 1992 from 21 minke, 15 fin, 6 blue, and 8 humpback whales. Hexachlorobenzene concentrations in the blubber of these whales were 101, 96, 110, and 177 ng/g lipid, respectively, and in all species, the concentrations were higher in males (140 ng/g lipid) than in females (103 ng/g lipid).

Becker et al. (1997) analyzed the blubber of 7 pilot whales, 5 harbor porpoises, 12 beluga whales, 2 northern fur seals, and 2 ringed seals that were obtained from the U.S. National Biomonitoring Specimen Bank. Hexachlorobenzene concentration ranges were 43–465 ng/g wet weight for pilot whales, 223–1,070 ng/g wet weight for harbor porpoises, 81.9–952 ng/g wet weight for beluga whales, 138– 741 ng/g wet weight for northern fur seals, and 125–156 ng/g wet weight for ringed seals (Becker et al. 1997). Elkin and Bethke (1995) reported that hexachlorobenzene was the most predominant organo-

6. POTENTIAL FOR HUMAN EXPOSURE

chlorine residue present in tissues of the caribou collected in the Northwest Territory of Canada. Residues ranged from a lipid corrected mean of 32.83 ng/g (ppb) in fat from Bathurst caribou to 129.41 ng/g (ppb) in Lake Harbor animals (Baffin Island). Consumption of meat and organs from these animals by native peoples, including native American populations of Inuit in Alaska, may constitute a potential human health risk if these tissues are a significant part of the diet.

The Florida Department of Environmental Protection and the Florida Marine Research Facility in St. Petersburg, Florida maintain archives of tissues obtained from dead Florida Manatees (*Trichechus manatus latirostris*). In 1996, Ames and Van Vleet (1996) obtained and analyzed 19 manatee samples from the Florida EPA. Of these samples, hexachlorobenzene was found at concentrations of 0.038 and 0.085 μ g/g in one blubber and one liver sample, respectively. The authors did not find a correlation between the contamination in manatees by hexachlorobenzene and other pesticides and the location in which the manatees were found dead; thus, the authors concluded that these manatees must have been contaminated elsewhere, especially since manatees are known to travel long distances.

Corsolini et al. (1999) analyzed chlorinated hydrocarbon concentrations in muscle and fat samples of the red fox collected from Sienna, Italy in 1992. Hexachlorobenzene mean concentrations in muscle and fat were 0.47 and 0.23 μ g/g lipid basis, respectively, and were the lowest of all of the chlorinated hydrocarbons that were tested (Corsolini et al. 1999).

Sitarska et al. (1995) collected tissue samples from 18 cows just after slaughtering, always from the same parts of the studied organs. Hexachlorobenzene mean concentrations were $1.394 \ \mu g/kg$ wet mass in the ovaries, $1.061 \ \mu g/kg$ wet mass in the mammary glands, and $0.550 \ \mu g/kg$ wet mass in the liver. Hexachlorobenzene was detected in the follicular fluid of cattle, sheep, goats, and pigs raised on local farms in Greece at mean concentrations of $1.77, 1.25, 1.63, \text{ and } 0.78 \ \text{ng/mL}$, respectively (Kamarianos et al. 2003).

Beyer (1996) subjected earthworms (*Lubricous terrestris*) to hexachlorobenzene in artificial soil. Over the course of 28 weeks, hexachlorobenzene concentrations in these earthworms ranged from 1.8 to 3 ppm. Beyer (1996) also conducted three 8-week experiments in which earthworm survival rates were 97, 99, and 100%, respectively.

Hexachlorobenzene was detected in composited milk samples collected through the U.S. Pasteurized Milk Network during 1990–1991 (Trotter and Dickerson 1993). The milk samples were collected at

6. POTENTIAL FOR HUMAN EXPOSURE

approximately 63 sites located in the United States, Puerto Rico, and the Panama Canal Zone. Hexachlorobenzene was detected in trace amounts in one sample collected in each of seven metropolitan areas (Cristobal, Panama Canal Zone; Kansas City, Missouri; Los Angeles, California; Memphis, Tennessee; Portland, Oregon; Spokane, Washington; Wichita, Kansas) and was detected at 0.001 ppm (1 ppb) in one sample from Rapid City, South Dakota.

Pesticide residue data from the FDA Adult Total Diet Study conducted from 1980 to 1982 were evaluated by Gartrell et al. (1986). These authors reported that hexachlorobenzene was detected in a wide variety of domestic foods: dairy products, meat, fish and poultry, oils and fats, and sugar and adjuncts. The highest mean concentrations of hexachlorobenzene were detected in oils and fats (0.9 ppb) and in meat, poultry, and fish (0.2 ppb). Concentrations of hexachlorobenzene in ready-to-eat foods were monitored for 10 years from 1982–1991 through the FDA's Revised Market Basket Survey. Hexachlorobenzene was detected in 618 samples of 81 different foods at a mean concentration of 0.0006 $\mu g/g$ (0.6 ppb) (KAN-DO Office and Pesticide Teams 1995). In food composites from six Canadian cities, the mean concentration of hexachlorobenzene in positive samples (4.8% of 913 total analysis) was 0.5 ng/g (Newsome et al. 2000). The U.S. Food and Drug Administration monitored domestic and imported apples and rice by collecting random samples for a period of 12 months beginning in October, 1993. Hexachlorobenzene was not determined to be in violation according to the concentration limits set for this compound in any of the domestic and imported apple and rice samples; however, it was found in 1 out of 612 imported rice samples (0.02 ppm), but this concentration does not violate any limit set by the EPA (Roy et al. 1997). The concentrations and occurrences of hexachlorobenzene residues in butter from Spain (n=36, 89% positive) and the rest of Europe (n=20, 70% positive) were 5.864 ± 3.171 ng/g wet weight and 3.022±3.964 ng/g wet weight, respectively (Badia-Vila et al. 2000).

The frequency of detection of hexachlorobenzene in the FDA Total Diet Study conducted in 1982–1984 was 9% (Gunderson 1988). Hexachlorobenzene intakes, in µg/kg body weight/day, estimated for these total diet analyses were 0.0020 and 0.0011 for 14–16-year-old males and 60–65-year-old females, respectively. In additional FDA Total Diet Studies, the frequency of detection of hexachlorobenzene residues declined to 7% in 1988 (FDA 1989), 5% in 1989 (FDA 1990), 4% in 1990 (FDA 1991), 2% in 1991 (FDA 1992), <2% on 1991–1993 (FDA 1994), and <2% in 1994 (FDA 1995). Hexachlorobenzene intakes (µg/kg body weight/day) estimated for the Total Diet Analyses also declined from intakes estimated in the 1982–1984 analysis and were 0.0011 and 0.0006 in 1988 (FDA 1989); 0.0009 and 0.0005 in 1989 (FDA 1990); 0.0005 and 0.0002 in 1990 (FDA 1991); and 0.0004 and 0.0002 in 1991 (FDA 1992) for 14–16-year-old males and 60–65-year-old women, respectively.

6. POTENTIAL FOR HUMAN EXPOSURE

Hexachlorobenzene levels in food products collected from FDA's Total Diet Study (TDS) market basket surveys are summarized in Tables 6-3 and 6-4. The TDS is FDA's ongoing market basket survey of core foods in the U.S. food supply to determine levels of various pesticide residues, contaminants, and nutrients in foods and to estimate exposures of these substances in representative diets of specific age and gender groups in the United States. For each market basket, food samples are collected from grocery stores and fast food restaurants in different cities, prepared table-ready (i.e., as they would be consumed), and composited for analysis.

Domestic samples of mixed feed rations were collected and analyzed by the FDA for pesticide surveillance during fiscal years 1989–1994. Hexachlorobenzene residue was detected in 1 of 457 samples in trace amounts (Lovell et al. 1996).

Burton and Bennett (1987) estimated a human body burden for hexachlorobenzene of 0.7 mg derived primarily from dietary intake of fatty foods ($0.2 \mu g/day$). Inhalation was estimated to contribute 100 times less than dietary intake ($0.002 \mu g/day$) and consumption of drinking water was also considered to contribute only negligible amounts of hexachlorobenzene ($0.06 \mu g/year$).

Hexachlorobenzene was detected in 98% of cheese samples obtained from of 61 commercially available brands in the Canary Islands (Almeida-Gonzalez et al. 2012). The average concentrations of hexachlorobenzene were 6.95±7.77 and 2.27±1.46 ng/g fat for conventional and organic cheeses, respectively.

An exploratory study of chemical exposure was conducted among Vietnamese, Bangladeshi, and local resident sportfish consumers in the Montreal region of the St. Lawrence River. The concentration ranges for the respective groups are as follows: 0.01-0.04; 0.01-0.02; and $0.01-0.07 \mu g/L$, indicating a positive correlation between local residents consuming sportfish from the St. Lawrence River and hexachlorobenzene concentrations (Kosatsky et al. 1999). Anderson et al. (1998) conducted a study to assess hexachlorobenzene contamination in human serum and urine samples from frequent consumers of sport fish from Lakes Michigan, Erie, and Huron. A telephone survey was conducted requesting fish eating habits with special attention to lake trout, brown trout, rainbow trout, or chinook or coho salmon, carp or catfish, and walleye or perch or smelt. After the survey, each angler was invited to give a serum sample. The minimum and maximum hexachlorobenzene concentrations of all 30 participating subjects were 0.02 and

Table 6-3. Hexachlorobenzene Levels in Food Items from the Food and DrugAdministration's Total Diet Study Market Baskets 1991–1993 Through2003–2004 Collected Between September 1991 and October 2003^a

		Number		Minimum	Maximum	
Food	Number	≥LQ	Trace	Mean (ppm)	(ppm)	(ppm)
Milk, lowfat (2%), fluid	44	0	1	0.00000	0.0001	0.0001
Cheese, American, processed	44	0	8	0.00006	0.0001	0.0001
Cheese, cheddar, natural (sharp/mild)	44	0	5	0.00005	0.0003	0.0006
Beef, ground, regular, pan- cooked	44	0	11	0.00005	0.0001	0.0004
Beef roast, chuck, oven- roasted	44	0	2	0.00001	0.0001	0.0003
Pork chop, pan-cooked with oil	44	0	2	0.00001	0.0002	0.0003
Pork sausage (link/patty), oven-cooked	44	1	0	0.00005	0.0020	0.0020
Pork bacon oven cooked	44	0	2	0.00002	0.0003	0.0006
Lamb chop, pan-cooked with oil	44	4	18	0.00030	0.0001	0.0020
Frankfurter (beef/pork), boiled	44	0	8	0.00004	0.0001	0.0004
Tuna canned in oil, drained	40	0	1	0.00001	0.0005	0.0005
Peanut butter, creamy	44	0	1	0.00000	0.0002	0.0002
Peanuts, dry roasted, salted	44	0	1	0.00000	0.0002	0.0002
Bread rye	44	0	1	0.00001	0.0003	0.0003
Watermelon, raw/frozen	44	1	4	0.00006	0.0001	0.0010
Summer squash, fresh/frozen, boiled	44	0	5	0.00003	0.0001	0.0007
Squash, winter (Hubbard/acorn) Fresh/frozen broiled	44	0	3	0.00003	0.0001	0.0009
Potato chips	44	0	1	0.00000	0.0002	0.0002
Chili con carne with beans, canned	4	0	1	0.00003	0.0001	0.0001
Quarter-pound hamburger on bun, fast food	44	0	5	0.00001	0.0001	0.0002
Meatloaf, beef, homemade	44	0	9	0.00003	0.0001	0.0003
Butter salted	44	6	29	0.00087	0.0004	0.0070
Half & half cream	44	0	2	0.00000	0.0001	0.0001
Sandwich cookies with crème filling	44	0	1	0.00000	0.0001	0.0001
Cheese, Swiss natural	44	0	9	0.00009	0.0003	0.0005
Cream cheese	44	0	8	0.00007	0.0003	0.0005
Lunch meat (ham)	44	0	2	0.00001	0.0002	0.0004
Chicken nuggets, fast-food	44	0	1	0.00000	0.0001	0.0001
Chicken, fried (breast, leg, and thigh) fast food	40	1	0	0.00005	0.0020	0.0020
Haddock	20	0	6	0.00009	0.0002	0.0003

Table 6-3. Hexachlorobenzene Levels in Food Items from the Food and Drug
Administration's Total Diet Study Market Baskets 1991–1993 Through
2003–2004 Collected Between September 1991 and October 2003 ^a

		Numbe	r		Minimum	Maximum
Food	Numb	er ≥LQ	Trace	Mean (ppm)	(ppm)	(ppm)
Crackers butter type	44	0	3	0.00005	0.0004	0.0008
Quarter-pound cheeseburger on bun, fast food	44	0	2	0.00001	0.0001	0.0002
Fish sandwich on bun, fast- food	44	2	9	0.00015	0.0001	0.0020
Frankfurter on bun, fast-food	40	1	4	0.00007	0.0001	0.0020
Taco/tostada with beef and cheese, from Mexican carry out	44	0	3	0.00003	0.0003	0.0004
lce cream, regular, vanilla	44	0	2	0.00000	0.0001	0.0001
Sugar cookies	44	0	1	0.00001	0.0003	0.0003
Sour cream	44	0	8	0.00004	0.0001	0.0004
Olive/safflower oil	44	0	1	0.00002	0.0008	0.0008
Salmon, steaks/fillets, baked	24	11	12	0.00098	0.0003	0.0020
Baby food, squash	44	3	2	0.00008	0.0002	0.0010
Baby food, cereal, mixed, dry, prepared with water	44	0	5	0.00003	0.0002	0.0005

^aNumber \geq LQ: number of result(s) that were greater than the limit of quantification (LQ). Number of traces: number of result(s) that were greater than or equal to the limit of detection but less than the LQ. Statistics were calculated using value of 0 for results below the limit of detection.

Source: FDA 2006a

Table 6-4. Hexachlorobenzene Levels in Food Items from the Food and Drug Administration's Total Diet Study Market Baskets 2004-1 Through 2005-4 (Eight Market Baskets) Collected Between October 2003 and August 2005^a

		Number			Minimum	Maximum
Food	Number	≥LQ	Trace	Mean (ppm)	(ppm)	(ppm)
Cheese, American	8	0	1	0.00003	0.2002	0.0002
Beef, ground	8	1	1	0.00014	0.0001	0.0010
Lamb, chop	8	0	2	0.00003	0.0001	0.0001
Bologna	8	0	1	0.00004	0.0003	0.0003
Peanut butter, smooth	8	0	1	0.00001	0.0001	0.0001
Fruit-flavored sweetened cereal	8	0	1	0.00001	0.0001	0.0001
Summer squash, fresh/frozen	8	0	3	0.00005	0.0001	0.0001
Meatloaf	8	1	2	0.00004	0.0001	0.0001
Butter	8	0	2	0.00014	0.0002	0.0009
Half & half cream	8	0	1	0.00001	0.0001	0.0001
Cheese, Swiss, natural	8	0	1	0.00004	0.0003	0.0003
Cream cheese	8	0	1	0.00003	0.0002	0.0002
Sugar cookies	8	0	1	0.00001	0.0001	0.0001
Sour cream	8	0	1	0.00001	0.0001	0.0001
Salmon	8	0	6	0.00023	0.0002	0.0004
Baby food, squash	8	0	1	0.00001	0.0001	0.0001
Granola bar with raisins	8	0	1	0.00001	0.0001	0.0001

^aNumber \geq LQ: number of result(s) that were greater than the limit of quantification (LQ). Number of traces: number of result(s) that were greater than or equal to the limit of detection but less than the LQ. Statistics were calculated using value of 0 for results below the limit of detection.

Source: FDA 2006b

6. POTENTIAL FOR HUMAN EXPOSURE

0.2 ppb, respectively, with a median concentration of 0.1 ppb. Eight participants from Lake Michigan had minimum and maximum concentrations of 0.09 and 0.2 ppb, respectively, with a median concentration of 0.1 ppb, and 11 participants from Lake Huron and Lake Erie had respective minimum and maximum concentrations of 0.04 and 0.2 ppb and 0.02 and 0.2 ppb with median concentrations of 0.1 and 0.09 ppb. A comparison group from Arkansas (180 serum samples) had hexachlorobenzene concentrations ranging from not detected to 0.3 ppb with a median of 0.1 ppb (Anderson et al. 1998). This study illustrates that since the comparison group had the highest range in concentration, there is a wide spread of hexachlorobenzene contamination and any population may possibly be affected.

Sinkkonen et al. (1995) discovered concentrations of hexachlorobenzene in pine needles in the vicinity of a metal reclamation plant. Five sites sampled and analyzed for hexachlorobenzene in the wax of the needles and the rest of the needles in 1991 (0.257–0.731 ng/g; 0.758–3.170 ng/g), 1992 (0.142–0.692 ng/g; 0.255–1.785 ng/g), and 1993 (0 concentration found in the wax of the needles; 0.217–0.885 ng/g) show decreasing concentrations (Sinkkonen et al. 1995). In 1996, Sinkkonen et al. (1996) analyzed pine needles in and around the metal reclamation plant again. Composite samples from the years 1993, 1994, and 1995 had hexachlorobenzene concentrations of 6.9–8.3 ng/g in the wax of the needles and 0.2–2.2 ng/g in the rest of the needles, contrary to the decreasing trends found from the analysis conducted during 1991–1993.

6.5 GENERAL POPULATION AND OCCUPATIONAL EXPOSURE

Hexachlorobenzene is no longer produced (as an end-product) or used as a pesticide in the United States. Consequently, the current potential for exposure of the general population appears to be very limited. Members of the general population may be exposed to very low concentrations of hexachlorobenzene through ingestion of contaminated foodstuffs, particularly those with high lipid content such as meat, poultry, and fish. General population exposure to hexachlorobenzene via inhalation or dermal contact would be much less. Occupational exposure is possible for workers involved in the production of chlorinated hydrocarbons, which releases hexachlorobenzene as a byproduct.

Brock et al. (1998) investigated four families in Iowa and two families in North Carolina for exposure to several organochlorine pesticides by analyzing the pesticide levels in their serum. Among the farmers from Iowa, mean hexachlorobenzene concentrations in serum ranged from 0.12 to 0.26 ng/mL, and their spouses had mean levels ranging from 0.05 to 0.24 ng/mL. The two farmers from North Carolina had mean levels of 0.15 and <0.05 ng/mL, and their respective spouses had levels of 0.16 and 0.17 ng/mL. It

6. POTENTIAL FOR HUMAN EXPOSURE

was shown that in one family, the pesticide serum level in the spouse (0.17 ng/mL) was higher than that in the farmer (<0.05).

The concentration of hexachlorobenzene in 12 San Francisco, California firefighters' serum was evaluated by high resolution gas chromatography-high resolution mass spectrometry (Shaw et al. 2013). The samples were obtained within 24 hours of responding to a fire. All samples of serum contained hexachlorobenzene; the concentrations ranged from 8 to 46 ng/g lipid weight basis.

Glynn et al. (2000) studied the serum concentration of hexachlorobenzene in a group of 790 men ages 40-75 who were randomly selected from both rural and urban areas of Uppsala, Sweden. The mean concentration of hexachlorobenzene was 83.1±133.6 ng/g (ppb) lipid for this group. This study group had lower serum concentrations than males who had recent occupational exposure or high environmental exposure. Hagmar et al. (2001) examined blood samples from 110 men who consumed varying amounts of fish from the Baltic Sea. The median plasma level of hexachlorobenzene in the 43 Swedish and 67 Latvian adult males in this study group was 84 ng/g (ppb) lipid. In a study of intra-individual variation in serum levels of 39 Swedish men, mean hexachlorobenzene concentrations measured in 1991 and 2001 were 77 and 33 ng/g (ppb) lipid, respectively, displaying an average decrease of 53% after a decade (Hagmar et al. 2006). The average decrease in hexachlorobenzene concentration was associated with high fish consumption in 1991 as compared to 2001, but this only explained 12% of the variation. The serum concentrations of pregnant women from the Disko Bay area, Greenland were studied by Bjerregaard and Hansen (2000). The women in this study consume a high level of meat and blubber from marine animals. The concentration of hexachlorobenzene in plasma taken from these women between the years 1994 and 1996 was 1.2 ng/mL wet weight (range, 0.1–7.0 ng/mL wet weight). Hexachlorobenzene concentrations in serum in study groups of active smoking mothers, passive smoking mothers, and nonsmoking mothers from Germany were 0.87 ng/mL (range, 0.23–4.38 ng/mL), 0.55 ng/mL (range, <0.10–3.27 ng/mL), and 0.46 ng/mL (range, <0.10–2.73 ng/mL) (Lackman et al. 2000). The mean concentration of hexachlorobenzene in whole blood from infant children ranged from 0.13 to 0.23 ng/mL (Karmaus et al. 2001). The highest concentration was observed in children who were breastfed for over 12 weeks after birth.

Serum samples from French adults aged 18–74 years had a reported geometric mean hexachlorobenzene concentration of 24.3 ng/g lipid in The French National Nutrition and Health Study (Saoudi et al. 2014). The median concentration of hexachlorobenzene in serum samples from 101 subjects in the National Health System Patients Roster for Calatafimi, Italy was 18.63 ng/g lipids and the detection frequency was

6. POTENTIAL FOR HUMAN EXPOSURE

84.2% (Amodio, 2012). The average concentration of hexachlorobenzene in human serum samples from Bizerte, Tunisia was 49.1±29.6 (Ben Hassine et al. 2014).

In Western Australia, almost all (99%) of the plasma samples collected from pregnant women 2 weeks before delivery contained hexachlorobenzene at levels above the limit of detection (Reid et al. 2013). The range of hexachlorobenzene concentration in the 167 blood plasma samples was $0.005-2.00 \mu g/L$ and the average concentration was $0.08 \mu g/L$.

Nonoccupational exposure to hexachlorobenzene for residents of two U.S. cities (Jacksonville, Florida and Springfield, Massachusetts) were studied over three seasons; summer 1986, spring 1987, and winter 1988 (Whitmore et al. 1994). The study focused primarily on inhalation exposures with primary environmental monitoring consisting of 24-hour indoor, personal, and outdoor air. For the Jacksonville, Florida population, the estimated mean hexachlorobenzene concentration ranged from 0.3 to 1.3 ng/m³ for indoor air, from not detected to 0.2 ng/m³ for outdoor air, and from 0.4 to 0.9 ng/m³ for personal air. For the Springfield, Massachusetts population, mean exposures were much less. The estimated hexachlorobenzene concentrations ranged from not detected to <0.05 ng/m³ for personal air. The mean air exposure estimated for hexachlorobenzene in Jacksonville, Florida, was 10 ng/day, while dietary exposure ranged from 70 to 120 ng/day. The mean air exposure for Springfield, Massachusetts, was not detected in the personal air samplers, while the dietary exposure was 105 ng/day. In both the Jacksonville, Florida population, characterized as a high pesticide use area, and in the Springfield, Massachusetts population, characterized as a low pesticide use area, the dietary exposure to hexachlorobenzene was the predominant exposure pathway.

In the National Human Adipose Tissue Survey, hexachlorobenzene was found in 35 of 46 human adipose tissue samples from all regions of the United States at levels ranging from 12 to 1,300 ppb (EPA 1986c). In other studies of the general population, hexachlorobenzene has been found in human fat samples from residents of the Texas Gulf Coast at concentrations ranging from 18 to 35 ppb (Ansari et al. 1986). Kutz et al. (1991) summarized data on hexachlorobenzene residues in human adipose tissue collected in the United States from 1973 through 1983. The geometric mean concentrations increased slightly from 0.02 ppm (20 ppb) in 1973 to 0.05 ppm (50 ppb) in 1976, and then declined to 0.031 ppm (31 ppb) in 1983.

6. POTENTIAL FOR HUMAN EXPOSURE

Human breast adipose tissue samples from 36 females of 50–80 years in age were collected from the Yale-New Haven hospital in Connecticut. A correlation was made between breast adipose tissue and serum residues using Pearson's correlation coefficient. On a lipid adjusted basis, all 36 human adipose tissue samples were found to contain residues of hexachlorobenzene. The range of concentration was 2.5–33.3 ng/g (ppb) with a median of 17.7 ng/g (ppb) (Archibeque-Engle et al. 1997). A study conducted in British Columbia, Canada, Mes (1992) reported median and maximum hexachlorobenzene residues in biopsied fatty tissue of 18.8 and 87 ng/g (ppb), respectively. In a more recent study, small amounts of breast tissue were collected from 60 women undergoing breast surgery at Stanford University, California. The mean hexachlorobenzene concentration was 46 ng/g (ppb) fat with a minimum and maximum of 14 and 170 ng/g (ppb) fat, respectively (Petreas et al. 1998). Weistrand and Noren (1998) collected adipose tissue and liver samples from five Swedish men and two Swedish women. Hexachlorobenzene concentrations ranged from 12 to 129 ng/g (ppb) lipids with a mean of 56 ng/g (ppb) lipids in adipose tissue and 17 to 156 ng/g (ppb) lipids with a mean of 58 ng/g (ppb) lipids in the liver. Hexachlorobenzene levels in human adipose tissue from 64 mothers living in Veracruz, Mexico averaged 0.065 mg/kg (65 ppb) (range, 0.010–0.401 mg/kg) on a lipid adjusted basis (Waliszewski et al. 2000a). The mean concentrations of hexachlorobenzene in autopsy tissue samples from Greenlanders were 594 μ g/kg (ppb) lipid (range, 476–742 μ g/kg lipid), 588 μ g/kg (ppb) lipid (range, 156–1,890 μ g/kg lipid), $260 \ \mu g/kg \ (ppb) \ lipid \ (range, 175-387 \ \mu g/kg \ lipid), and 754 \ \mu g/kg \ (ppb) \ lipid \ (range, 603-943 \ \mu g/kg \ lipid)$ lipid) for subcutaneous fat, omental fat, brain, and liver tissues, respectively (Dewailly et al. 1999). Moreover, in a study evaluating hexachlorobenzene in 70-year-old Uppsala, Sweden citizens, plasma concentrations of hexachlorobenzene, ranging from 27.8 to 53.7 ng/g lipid, positively related to both visceral and subcutaneous adipose tissue content measured using abdominal magnetic resonance imaging (Roos et al. 2013).

In the National Health and Nutrition Examination Survey (NHANES II) conducted by CDC hexachlorobenzene levels in blood from the general population collected from 1976 to 1980 reported a median level for quantifiable positive results of 1.7 ppb, which did not vary among the four age groups studied (Murphy and Harvey 1985). Table 6-5 lists mean serum hexachlorobenzene concentrations (by whole weight and lipid-weight bases) in the U.S. population from NHANES IV (updated in 2015) using serum samples collected during the years 2003–2004 (CDC 2009, 2015). Mean concentrations of hexachlorobenzene were highest for adults \geq 20 years of age, slightly higher among females than males, and highest among Mexican Americans and lowest among non-Hispanic whites.

	rum hexachlorobenzene (95% Cl)	
Group	Whole weight (ng/g serum or ppb)	Lipid adjusted (ng/g lipid)
Age group		
All ages	0.092 (0.088–0.097)	15.2 (14.5–15.9)
12–19 years of age	0.065 (0.062–0.069)	13.3 (12.5–14.1)
≥20 years	0.097 (0.092–0.102)	15.5 (14.7–16.2)
Gender		
Males	0.090 (0.085–0.095)	14.5 (13.8–15.3)
Females	0.095 (0.089–0.100)	15.8 (15.0–16.6)
Race/ethnicity		
Non-Hispanic whites	0.094 (0.088–0.099)	15.1 (14.4–16.0)
Non-Hispanic blacks	0.081 (0.077–0.085)	14.5 (13.9–15.0)
Mexican Americans	0.098 (0.089–0.109)	16.2 (14.9–17.7)

Table 6-5. Geometric Mean Serum Hexachlorobenzene Concentrations (Whole Weight and Lipid Adjusted) for the Years 2003–2004 in the U.S. Population from the National Health and Nutrition Examination Survey

Source: CDC 2009, 2015

6. POTENTIAL FOR HUMAN EXPOSURE

Table 6-6 lists mean serum hexachlorobenzene concentrations (by whole weight and lipid-weight bases) in the U.S. population from NHANES IV (updated in 2015) using serum samples collected during the years 2007–2008 (CDC 2015). Grouped by race/ethnicity (non-Hispanic white, non-Hispanic black, Mexican-American), gender, and age, mean serum hexachlorobenzene levels were highest in the oldest age group of females (\geq 60 years of age) for each race/ethnicity, and higher among Mexican-American females than non-Hispanic white or black females.

Hexachlorobenzene levels in normal human blood serum samples were reported to be 2.2 ppb and were higher (4.6 ppb) in uremic serum samples (Rutten et al. 1988). In a study conducted in British Columbia, Canada, Mes (1992), reported median and maximum whole blood levels of 0.11 and 0.34 ng/g (ppb) in individuals from the general population. Hexachlorobenzene was detected in 100% of blood samples taken from young adults between the ages of 17 and 21 years of age from the Akwesasne Mohawk Nation on the St. Lawrence River at mean and maximum concentrations of 0.036 and 0.112 ppb, respectively (Gallo et al. 2011). Data indicated that Akwesasne young adults have sustained a lower overall exposure to hexachlorobenzene than similarly aged people from the general U.S. population, probably due to relatively lower legacy contamination and lower contributions to background body burden.

Butler et al. (2003) reported hexachlorobenzene concentrations in maternal and umbilical cord blood plasma for Inuit, Dene, Metis, Caucasian, and other non-Aboriginal participants living in the Northwest Territories and Nunavut areas of Arctic Canada. Overall, 523 women participated resulting in the collection of 386 maternal blood samples and 407 cord blood samples taken between May 1994 and June 1999. Hexachlorobenzene was detected in 100% of 385 maternal blood plasma samples with concentrations ranging from 0.02 to 4.51 μ g/L, and a mean value of 0.35 μ g/L. Hexachlorobenzene was detected in 97.5% of 400 cord blood plasma samples with concentrations ranging from not detected (detection limit was 0.02 μ g/L) to 1.01 μ g/L and a mean value of 0.10 μ g/L.

Although all uses of hexachlorobenzene as a pesticide in the United States were voluntarily canceled in 1984, occupational exposures may still occur among workers in the chlorinated solvent manufacturing industry, and workers currently involved in the manufacture and application of pesticides contaminated with hexachlorobenzene. Military or firefighting personnel who use pyrotechnic mixtures that release hexachlorobenzene and workers involved in the disposal of hexachlorobenzene contaminated materials, via combustion processes associated with municipal incinerators or those involved in the handling and treatment of wastes at hazardous waste sites, may be exposed to higher than background concentrations.

		Geometric mean serum hexachlorobenzene (± standard error)				
Group	Age (years)	Whole weight (ng/g serum or ppb)	Lipid adjusted (ng/g lipid)			
Non-Hispanic white male	12–19	0.062±0.003	12.4±0.8			
	20–39	0.062±0.003	9.45±0.46			
	40–59	0.074±0.003	10.1±0.3			
	≥60	0.075±0.003	11.8±0.5			
Non-Hispanic white female	12–19	0.033±0.001	7.06±0.29			
	20–39	0.047±0.002	7.94±0.23			
	40–59	0.071±0.005	10.8±0.7			
	≥60	0.102±0.004	15.1±0.7			
Non-Hispanic black male	12–19	0.046±0.004	9.67±0.79			
	20–39	0.051±0.003	9.11±0.58			
	40–59	0.061±0.003	9.58±0.43			
	≥60	0.062±0.003	10.3±0.6			
Non-Hispanic black female	12–19	0.034±0.003	7.20±0.57			
	20–39	0.035±0.001	7.04±0.3			
	40–59	0.059±0.004	9.90±0.58			
	≥60	0.090±0.006	15.1±1.2			
Mexican-American male	12–19	0.050±0.003	9.49±0.7			
	20–39	0.065±0.006	9.47±0.57			
	40–59	Not reported	20.6±8.3			
	≥60	0.089±0.011	12.1±0.8			
Mexican-American female	12–19	0.033±0.002	6.81±0.44			
	20–39	0.060±0.004	10.4±0.7			
	40–59	0.138±0.016	20.6±2.6			
	≥60	0.141±0.013	20.7±1.9			

Table 6-6. Geometric Mean Serum Hexachlorobenzene Concentrations (Whole Weight and Lipid Adjusted) for the Years 2007–2008 in the U.S. Population from the National Health and Nutrition Examination Survey

Source: CDC 2015

6. POTENTIAL FOR HUMAN EXPOSURE

NIOSH conducted a study in May, August, and November 2004 to investigate the extent of exposure to 30 U.S. magnesium workers during the processing of magnesium (NIOSH 2005). Hand wipes contained hexachlorobenzene concentrations of 0.14– 3.5μ g. The means of lipid-adjusted and whole-blood hexachlorobenzene from the 30 workers were 891 (range 253.0–6790.0) and 0.7 (range 0.2–3.4) ng/g (ppb) lipid, respectively. Full-shift air sampling results for hexachlorobenzene ranged from 0.096 μ g/m³ for a sample collected on a foundry operator to 5.3μ g/m³ for a maintenance helper working on a reactor. Bulk sample results indicated the presence of 250 μ g/g (ppb) of hexachlorobenzene in collected dust (NIOSH 2005). In a 10-year study (1976–1985), human adipose tissue samples and human milk from patients exposed to PCBs or pesticides in Ontario, Canada were analyzed for a variety of pesticides and industrial chemicals (Frank et al. 1988). Residues of hexachlorobenzene in milk ranged from a mean of 0.52 ppb (in whole milk) in 1983–1984 to a mean of 0.26 ppb (in whole milk) in 1985. The highest mean concentration (1.33 ppb) was observed in central Ontario residents. Urban residents had a higher mean concentration (0.57 ppb) as compared to rural residents with a mean of 0.27 ppb.

Plasma hexachlorobenzene concentrations in a Louisiana population living in a hexachlorobenzenecontaminated area averaged 3.6 ± 4.3 ppb. The highest plasma level (345 ppb) was detected in a waste disposal facility worker, while the highest plasma level in the general population was 23 ppb (Burns and Miller 1975). Workers at a carbon tetrachloride and perchloroethylene production facility had plasma hexachlorobenzene levels of up to 223 ppb. Hexachlorobenzene blood levels were determined over a 4-year period in men employed in the manufacture of chlorinated solvents (Currier et al. 1980). Blood levels ranged from 5 to 1,121 ppb (310.7 ppb mean) in 1974, 30–990 ppb (311.5 mean) in 1975, 3– 600 ppb (159.9 mean) in 1976, and 22–467 ppb (170.3 mean) in 1977. The hexachlorobenzene blood levels were strongly associated with the number of years worked in the chlorinated solvents plant, but they were poorly correlated with airborne hexachlorobenzene concentrations ranging from <1 to 13 ppb or wipe samples from work areas ranging from 0.03 to 124 µg/100 m².

Vegetable sprayers applying hexachlorobenzene-contaminated dimethyltetrachloroterephthalate (DCPA) had plasma levels of hexachlorobenzene ranging from 0 to 310 ppb (mean 40±63 ppb), accompanied by elevated levels of delta-aminolevulinic acid, but no health related adverse effects (Burns et al. 1974). Elevated urinary uroporphyrin and coproporphyrin were found in 1 of 54 men occupationally exposed to hexachlorobenzene (Morley et al. 1973).

6. POTENTIAL FOR HUMAN EXPOSURE

Workers at a new hazardous waste incinerator in Constanti, Spain had mean plasma levels of hexachlorobenzene at 152 µg/kg lipid (range, 19.4–854.0 µg/kg lipid) (Domingo et al. 2001). Employees at a hazardous waste incinerator in Constanti, Spain working in the plant, laboratory, and administration had mean plasma levels of hexachlorobenzene of 134 and 84, 182 and 179, and 223 and 179 µg/kg lipid in 1999 and 2000, respectively (Schuhmacher et al. 2002). Residents (n=608) living near an electrochemical factory in Catalonia, Spain had mean serum concentrations of hexachlorobenzene as follows (units ng/mL): general population, male (50.2), female (48.0), all (48.9); nonfactory workers, male (39.8), female (47.9), all (46.3); and living with a worker of the factory, yes (46.8), no (46.8) (Ballester et al. 2000).

Individuals employed in industries that manufacture or process hexachlorobenzene or products containing hexachlorobenzene may be exposed to the highest concentrations. The National Occupational Exposure Survey (NOES), conducted by NIOSH from 1981 to 1983, estimated that 1,038 workers employed at 10 facilities were potentially exposed to hexachlorobenzene in the United States (NOES 1990). The NOES database does not contain information on the frequency, concentration, or duration of exposure; the survey provides only estimates of workers potentially exposed to chemicals in the workplace.

6.6 EXPOSURES OF CHILDREN

This section focuses on exposures from conception to maturity at 18 years in humans. Differences from adults in susceptibility to hazardous substances are discussed in Section 3.7, Children's Susceptibility.

Children are not small adults. A child's exposure may differ from an adult's exposure in many ways. Children drink more fluids, eat more food, breathe more air per kilogram of body weight, and have a larger skin surface in proportion to their body volume. A child's diet often differs from that of adults. The developing human's source of nutrition changes with age: from placental nourishment to breast milk or formula to the diet of older children who eat more of certain types of foods than adults. A child's behavior and lifestyle also influence exposure. Children crawl on the floor, put things in their mouths, sometimes eat inappropriate things (such as dirt or paint chips), and spend more time outdoors. Children also are closer to the ground, and they do not use the judgment of adults to avoid hazards (NRC 1993).

Children are particularly susceptible to hexachlorobenzene by ingestion of breast milk. Table 6-7 summarizes the concentrations of hexachlorobenzene in human breast milk taken from women living in different regions of the world. Hexachlorobenzene was detected in human milk samples collected from

Table 6-7. Mean Levels of Hexachlorobenzene in Breast Milk

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Location of study	Concentration ^a	Reference
Australia (Victoria, 1985/1986)	0.005 ^b	Monheit and Luke 1990
Australia (Victoria)	0.41	Quinsey et al. 1995
Bavaria	0.019	Raab et al. 2013
Brazil (Porto Alegre)	0.02	Beretta and Dick 1994
Canada (1967–1992)	0.002–0.00044 ^b	Craan and Haines 1998
Quebec	0.002–0.00040 ^b	
Ontario	0.002–0.00048 ^b	
Canada (Arctic Quebec)		Dewailly et al. 1993
Inuit women	0.136	
Caucasian women	0.028	
Canada (multiple locations, 1992)	0.0044 ^b	Newsome et al. 1995
	0.015°	
Canada	0.026	Mes et al. 1993
Czech Republic		Schoula et al. 1996
Prague	0.639	
Kladno	0.570	
Uherske Hradiste	0.482	
France (multiple locations)	0.147	Bordet et al. 1993
Ghana (Akomadan)	0.04	Ntow 2001
Italy		Larsen et al. 1994
Rome	0.25	
Pavia	0.12	
Milan	0.20	
Mexico (Veracruz)	0.047	Waliszewski et al. 1996
Mexico (Guerrero)	0.013±0.014	Chavez-Almazan et al. 2014
New Zealand		Bates et al. 1994
Auckland, urban	0.020	
Northland, rural	0.021	
Christchurch, urban	0.030	
Canterbury, rural	0.063	
New Zealand	0.006763	Mannetje et al. 2014
Slovak Republic (Bratislava)	0.339	Prachar et al. 1993
Spain (Madrid)	1.0	Conde et al. 1993
Industrialized area	1.74	
Sweden (1972–1980)	0.110–220	Norén and Meironyté 2000
Sweden (1972–1980)	0.110–220	Norén and Meironyté 2000

Table 6-7. Mean Levels of Hexachlorobenzene in Breast Milk

Location of study	Concentration ^a	Reference
Thailand (Northern Thailand)	0.0051 ^b	Stuetz et al. 2001
United States (Arkansas)	0.03	Mattison et al. 1992
United States (Hawaii, 1979–1980)	0.046±0.049	Takei et al. 1983

^aµg/g on lipid basis. ^bWhole milk sample. ^cMilk fat sample.

6. POTENTIAL FOR HUMAN EXPOSURE

286

54 residents of Hawaii during 1979–1980 (Takei et al. 1983). The incidence of detection of hexachlorobenzene in the sampled population was 100% and the mean concentration of positive detections was 46±49 ppb (ng/g lipid basis) with residues ranging from 18 to 38 ppb (ng/g lipid basis). The authors state that the levels of hexachlorobenzene in human milk from residents of Hawaii are consistent with levels detected in an earlier human milk study conducted on women on the mainland United States. Schecter et al. (1998) found that hexachlorobenzene residues in the breast milk of a mother with nursing twins decreased from 10.7 ng/g (ppb) lipid to not detectable in 30 months. Thus, nursing infants are particularly susceptible to hexachlorobenzene poisoning due to the mother's decrease in body burden and the infant's intake (Schecter et al. 1998). In human milk samples collected from 2002 to 2007 from women residing in the agricultural region of Salinas, California and the urban San Francisco Bay Area, California, median concentrations of hexachlorobenzene in breast milk were 191 and 223 pg/g (0.191 and 0.223 ppb), respectively (Weldon et al. 2011). A study of organochlorine pesticide concentrations in human milk sampled throughout Canada during 1992 found that hexachlorobenzene was present in all 497 milk samples at mean concentrations of 0.44 ng/g (ppb) in whole milk and 14.5 ng/g (ppb) in milk fat (Newsome et al. 1995). A comparison by Canada Health of human milk contamination in whole milk from 1967 to 1992 was conducted in Canada, Quebec and Ontario. This study showed a decrease in hexachlorobenzene concentration in Canada (mean concentration; 2–0.44 ng/g), Quebec (median concentration; 2–0.40 ng/g), and Ontario (median concentration; 2–0.48 ng/g) (Craan and Haines 1998). Newsome et al. (1995) reported that concentrations of hexachlorobenzene were higher in women from the Great Lakes Basin area as compared to the rest of Canada. Concentrations of hexachlorobenzene were higher in the breast milk of women who consumed more than 100 g of fish weekly. Hexachlorobenzene has also been detected in the breast milk of women from China. For example, samples from women from Beijing had hexachlorobenzene concentrations ranging from 0.40 to 3.79 µg/kg whole milk (Song et al. 2013).

Children may also be exposed to chemicals via ingestion of contaminated foods. Hexachlorobenzene residues have been detected in 76% of samples analyzed as part of the National Human Adipose Tissue Survey (FY82) (EPA 1986c). These hexachlorobenzene residues are most likely the result of consumption of low levels of hexachlorobenzene in food, with calculated yearly intakes of 68, 22, and 5 µg for adults, toddlers, and infants, respectively (EPA 1986b). Yess et al. (1993) evaluated hexachlorobenzene residues from 1985 to 1991 detected in the Total Diet Studies of infant and adult foods that are consumed by infants and young children. These authors reported maximum hexachlorobenzene residues detected in various food groups as follows: combination meat dinners—pork (0.4 ppb), beef (0.3 ppb), chicken/turkey/vegetable (0.3 ppb), beef and vegetable (0.1 ppb); vegetables

6. POTENTIAL FOR HUMAN EXPOSURE

and fruits—pears (1.0 ppb), apples (0.4 ppb), and carrots (0.2 ppb); milk products—canned evaporated milk (0.5 ppb), whole milk (0.2 ppb), and low-fat (2%) milk (0.1 ppb); and peanut butter (5.0 ppb).

Hexachlorobenzene intakes, in µg/kg body weight/day, estimated for these total diet analyses (1982– 1984) were 0.0015 for 6–11-month-old infants. A follow-up study found a decrease in intakes that were estimated in 1982–1984. Hexachlorobenzene intakes (µg/kg body weight/day) were estimated to be 0.0016 in 1988 (FDA 1989); 0.0007 in 1989 (FDA 1990); 0.0004 in 1990 (FDA 1991); and 0.0003 in 1991 (FDA 1992) for 6–11-month-old infants.

Hexachlorobenzene plasma concentrations were studied for a population of 4-year-old children born between 1997 and 1999 in an urban area under the influence of hexachlorobenzene emissions from a chloro-alkali plant and in a rural area where hexachlorobenzene was present at relatively low levels (Carrizo et al. 2008). In the urban area of Ribera d'Ebre, Catalonia, Spain, the measured average hexachlorobenzene plasma concentration was 1.4 ng/mL, with a range of 0.17–5.8 ng/mL. The measured average plasma concentration in the rural area of Menorca Island in the Mediterranean Sea was 0.42 ng/mL, with a range of 0.067–2.1 ng/mL. Higher concentrations of hexachlorobenzene were measured in children who were breastfed as opposed to formula fed, showing that a major portion of the compound was incorporated into the serum during the lactation period. The average concentrations were 1.5 and 0.99 ng/mL for Ribera d'Ebre and 0.47 and 0.23 ng/mL for Menorca for breastfed and formulafed children, respectively (Carrizo et al. 2008). Dallaire et al. (2002) studied umbilical cord plasma of newborns collected between 1993 and 2000 from the Lower North Shore of the St. Lawrence River in Quebec, Canada, and found a 69% decrease in the mean hexachlorobenzene concentration in that time span, falling from 35.5 µg/kg in 1993 to 11.6 µg/kg in 2000. In a study of umbilical cord blood samples taken from Inuit infants born in Nunavik, Quebec between 1994 and 2001, the mean hexachlorobenzene concentration decreased an average of 6.6% per year during that time span (Dallaire et al. 2003). Hexachlorobenzene was detected in 100% of the 251 cord blood samples collected (Dallaire et al. 2003).

Although inhalation exposures of hexachlorobenzene in children have not been studied, it is anticipated that exposure by this route will not be significant in outdoor environments. The Henry's law constant of hexachlorobenzene is 5.8×10^{-4} atm-m³/mol (Ten Hulscher et al. 1992), indicating that this compound will volatilize, especially in moist soils with low organic content. Hexachlorobenzene's high log K_{oc} of 6.08 (EPA 1981), however, indicates that volatilization from soil surfaces will be attenuated. Considering that hexachlorobenzene concentrations in the environment are extremely low, exposure of children by inhalation is expected to be insignificant. After a hexachlorobenzene spill, inhalation

6. POTENTIAL FOR HUMAN EXPOSURE

exposure may be important before environmental equilibrium is attained. Under these conditions, high concentrations of hexachlorobenzene would be found in the atmosphere, due to hexachlorobenzene's calculated vapor density of 10. This situation, however, is not expected to occur since hexachlorobenzene is no longer produced or used commercially and is only found as an impurity in pesticides and as a byproduct of chlorinated hydrocarbons.

The EPA issued a warning regarding pesticides and advised that potential exposure of pesticides to young children via dermal absorption and ingestion was more important than inhalation routes (Jantunen et al. 1997).

6.7 POPULATIONS WITH POTENTIALLY HIGH EXPOSURES

In addition to those individuals occupationally exposed to hexachlorobenzene identified in Section 6.5, several groups within the general population may receive potentially higher exposures to hexachlorobenzene. These groups within the general population include individuals living near facilities where hexachlorobenzene is produced as a byproduct, individuals living near the current or former NPL hazardous waste sites where this compound is present, recreational and subsistence fishermen who consume higher amounts of fish than the general population, and native populations (including Native American populations such as the Inuit of Alaska) who may be exposed to higher levels of hexachlorobenzene associated with dietary intakes of caribou and other game species.

Recreational and subsistence fishermen who consume appreciably higher amounts of locally caught fish from contaminated water bodies may be exposed to higher levels of hexachlorobenzene associated with dietary intake than members of the general population (EPA 1995a). Hexachlorobenzene contamination in fish and shellfish has triggered the issuance of several human health advisories. As of September 1994, hexachlorobenzene was identified as the causative pollutant in fish consumption advisories in Louisiana and Ohio. This information is summarized in Table 6-8. EPA has identified hexachlorobenzene as a target analyte and recommended that this chemical be monitored in fish and shellfish tissue samples collected as part of state toxics monitoring programs. EPA recommends that residue data obtained from these monitoring programs should then be used by the states to conduct risk assessments to determine the need for issuing fish and shellfish consumption advisories for the protection of the general public as well as recreational and subsistence fishermen (EPA 1997a).

State	Waterbody	Extent	Species
Louisiana	Devil's Swamp Lake and Bayou Baton Rouge	7 miles	All fish
	Calcasieu and Cameron parishes	6 miles	All fish and shellfish
Ohio	Tuscarawas River	Turkey Foot Road. (SR 619), Barberton, to South Broadway Street (SR 416), New Philadelphia (Tuscarawas County)	Carp-common, bass- smallmouth, bullhead- yellow, bass-rock, catfish-channel, bass- largemouth

Table 6-8. Fish Consumption Advisories^a

^aFrom EPA 1997a National Listing of Fish Consumption Advisories

6. POTENTIAL FOR HUMAN EXPOSURE

Native American populations such as the Inuit of Alaska or other subsistence hunters living in high latitude areas of the United States and Canada may be exposed to hexachlorobenzene residues in caribou, beluga whales, polar bears, seals, and other game species. Significantly higher concentrations of hexachlorobenzene (mean 136 ng/g [ppb]) were reported in breast milk of Inuit mothers from eastern Canada (Quebec Province) as compared with residues of 28 ng/g (ppb) in Caucasian mothers (Dewailly et al. 1993). By analogy, it is possible that Inuit populations in western North America (Alaska) may receive potentially higher hexachlorobenzene exposures from their dietary habits. In a follow-up study by Dewailly et al. (1999) 26 subcutaneous fat samples, 41 omental fat samples, 17 brain samples, and 26 liver samples were collected in November 1992 to Mid-October 1994 from Inuit Greenlanders. Mean hexachlorobenzene concentrations were 594, 588, 260, and 754 μ g/kg lipid basis, respectively. A comparison of these data clearly suggest an increase in Inuit population's hexachlorobenzene levels from dietary habits. Maternal body burden and lactational transfer of hexachlorobenzene can increase tissue levels in the neonate (Ando et al. 1985; Frank et al. 1988).

6.8 ADEQUACY OF THE DATABASE

Section 104(i)(5) of CERCLA, as amended, directs the Administrator of ATSDR (in consultation with the Administrator of EPA and agencies and programs of the Public Health Service) to assess whether adequate information on the health effects of hexachlorobenzene is available. Where adequate information is not available, ATSDR, in conjunction with NTP, is required to assure the initiation of a program of research designed to determine the health effects (and techniques for developing methods to determine such health effects) of hexachlorobenzene.

The following categories of possible data needs have been identified by a joint team of scientists from ATSDR, NTP, and EPA. They are defined as substance-specific informational needs that if met would reduce the uncertainties of human health assessment. This definition should not be interpreted to mean that all data needs discussed in this section must be filled. In the future, the identified data needs will be evaluated and prioritized, and a substance-specific research agenda will be proposed.

6.8.1 Identification of Data Needs

Physical and Chemical Properties. The physical and chemical properties of hexachlorobenzene are sufficiently well documented to permit estimation of the compound's environmental fate (EPA 1981; Hansch et al. 1995; Haynes and Lide 2010; ten Hulscher et al. 1992; O'Neil et al. 2006; Verschueren 2001). No further information is needed.

6. POTENTIAL FOR HUMAN EXPOSURE

Production, Import/Export, Use, Release, and Disposal. According to the Emergency Planning and Community Right-to-Know Act of 1986, 42 U.S.C. Section 11023, industries are required to submit substance release and off-site transfer information to the EPA. The TRI, which contains this information for 2013, became available in October of 2014. This database is updated yearly and should provide a list of industrial production facilities and emissions.

Hexachlorobenzene is not currently manufactured as a commercial end-product in the United States and has not been commercially produced since the late 1970s (EPA 1986b). However, hexachlorobenzene currently is produced as a by-product or impurity in the manufacture of chlorinated solvents, other chlorinated compounds, and several currently registered pesticides (Bailey 2001; EPA 1986b; IARC 1979).

The total amount of hexachlorobenzene released as a byproduct in the production of all chlorinated solvents was estimated as 0.3 kg/year in the mid-1990s (Bailey 2001), while hexachlorobenzene released through use of eight major pesticides containing hexachlorobenzene accounted for 1,270 kg/year (Bailey 2001). Current production estimates for hexachlorobenzene as a by-product or impurity are not available. Current, quantitative estimates of production of hexachlorobenzene from all sources are needed to evaluate potential exposures and risks to human health.

There are no current commercial uses of hexachlorobenzene in the United States, although the compound was used as a fungicide until 1984 when the last registered use as a pesticide was voluntarily cancelled (Beyer 1996). Prior to the registration cancellations, hexachlorobenzene was registered as a seed protecting for use on several grains (principally wheat) and field crops (EPA 1986b). Hexachlorobenzene was also used in pyrotechnic and ordinance materials and in synthetic rubber production (EPA 1986b). Impurities of hexachlorobenzene in currently registered pesticides (picloram, PNCB, chlorothalonil, Dacthal[®], atrazine, simazine, and pentachlorophenol) (Bailey 2001; EPA 1986b, 1993) appear to be a continuing source of hexachlorobenzene exposure for the general population. Currently, data regarding the release of hexachlorobenzene into the environment as an impurity via the use of other pesticides is not available.

No current information was located on import/export volumes for hexachlorobenzene, although import/export volumes for hexachlorobenzene/DDT combined are available (NTDB 1995).

6. POTENTIAL FOR HUMAN EXPOSURE

Hexachlorobenzene is listed as a hazardous waste. It is regulated under the Clean Water Effluent Guidelines as stated in Title 40, Section 400–475, of the Code of Federal Regulations and the Resource Conservation and Recovery Act (RCRA) (see Chapter 8). Past disposal methods have included incineration, landfills, discharges to municipal sewage treatment plants, and emissions to the atmosphere. The recommended method of disposal for hexachlorobenzene is incineration (Clayton and Clayton 1981; EPA 1988a, 1989a; HSBC 2012; Lamb et al. 1994). No further information on disposal practices is needed; however, estimates on the volume of hexachlorobenzene disposed of annually and the disposal method used are needed to assess exposure pathways.

Environmental Fate. Hexachlorobenzene released to the environment partitions to several environmental compartments (air, water, soil and sediment, and biological organisms). Hexachlorobenzene partitions to the atmosphere from soil surfaces through volatilization (Nash and Gish 1989). The remainder is adsorbed strongly to soil where it persists for extended periods (half-life of months to years) due to its resistance to biodegradation (Beall 1976; Beck and Hansen 1974; Isensee et al. 1976). Leaching of hexachlorobenzene into groundwater is not expected to occur very rapidly under most circumstances due to the compound's high sorption characteristics (Swann et al. 1983). Yuan et al. (1999) have reported that hexachlorobenzene can be dechlorinated to dichlorobenzenes under anaerobic conditions in the laboratory using sewage sludge as inoculum.

Once in the atmosphere, hexachlorobenzene exists in both the vapor and particulate phase; however, the vapor phase predominates (Ballschmiter and Wittlinger 1991; Bidleman et al. 1989; Lane et al. 1992). Degradation of hexachlorobenzene in the atmosphere is quite slow (1.69 years) (Brubaker and Hites 1998). Since hexachlorobenzene is hydrophobic, wet deposition will not be an important loss process. In cold high latitude zones, dry deposition of hexachlorobenzene aerosols is encouraged (Ballschmiter and Wittlinger 1991; Lane et al. 1992; Wania and Mackay 1993). Atmospheric transport of hexachlorobenzene is a major mechanism for global translocation of this compound (Eisenreich et al. 1981; Kelly et al. 1991). Long-range global transport of hexachlorobenzene released anywhere in the world can occur via atmospheric or oceanic systems (Ballschmiter and Wittlinger 1991; Wania and MacKay 1993).

Hexachlorobenzene released to water will volatilize, adsorb to sediments, or bioaccumulate in fish and other aquatic organisms (Bishop et al. 1995; EPA 1992; Kelly et al. 1991; Langlois and Langis 1995; Oliver and Nichol 1982a; Quemerais et al. 1994; Rostad et al. 1993; Schmitt et al. 1990; Zabik et al. 1995). Hydrolysis and biodegradation are not significant processes in water. Information on biodegra-dation of hexachlorobenzene under anaerobic conditions in a laboratory study exists (Yuan et al. 1999),

6. POTENTIAL FOR HUMAN EXPOSURE

but degradation data under field conditions were not found. Further information on these processes, including degradation products, are needed to determine potential mechanisms and sources of hexachlorobenzene releases from soils and the potential for the compound and its degradation products to contaminate groundwater.

Both bioaccumulation and biomagnification of hexachlorobenzene were reported to occur in an aquatic laboratory microcosm system (Burkhard et al. 1997; Isensee et al. 1976); however, data by Russell et al. (1995) suggests that hexachlorobenzene bioaccumulates, but is not biomagnified in certain fish populations in Lake Erie. In terrestrial ecosystems, hexachlorobenzene can also be accumulated in several agricultural plant species in the roots and parts of the plants closest to the soil (Kraaij and Connell 1997; Scheunert et al. 1983; Schroll et al. 1994; Smelt and Leistra 1974). In lichens, a high latitude forage food for caribou, hexachlorobenzene was found to be bioconcentrated 8,800,000–17,000,000 times the concentration in the atmosphere (Muir et al. 1993). Although the issue of biomagnification in some ecosystems needs to be clarified, there are adequate data on the bioconcentration of hexachlorobenzene in both aquatic and terrestrial ecosystems.

Bioavailability from Environmental Media. Hexachlorobenzene can be absorbed following inhalation of contaminated workplace air (Burns et al. 1974; Currier et al. 1980; Richter et al. 1994). Since hexachlorobenzene is moderately volatile, inhalation may not be a major concern except at hazardous waste sites or in industrial settings. Hexachlorobenzene can be absorbed following ingestion of contaminated food or water. Exposure to hexachlorobenzene through ingestion of food contaminated with low levels of the compound is probably the greatest source of exposure for the general population. Exposure to hexachlorobenzene through ingestion of contaminated drinking water is not expected to be an important source of concern since the compound is not very soluble in water. Although there are no quantitative data on the human absorption of orally administered hexachlorobenzene, gastrointestinal absorption has been demonstrated for rats (Albro and Thomas 1974; Ingebrigtsen and Nafstad 1983; Ingebrigtsen et al. 1981). The lymphatic system has also been shown to play an important part in the absorption of hexachlorobenzene in the intestines. Hexachlorobenzene is absorbed by the lymphatic system in the region of the duodenum and jejuno-ileum and is deposited in the adipose tissue, bypassing the portal circulation (Iatropoulos et al. 1975). Since hexachlorobenzene is tightly bound to soil particles, ingestion of hexachlorobenzene-contaminated soil, particularly by children, may also be an important route of exposure near production and processing facilities or near hazardous waste disposal sites. No information was available regarding absorption of hexachlorobenzene following dermal contact. Information regarding the bioavailability of hexachlorobenzene from both ingestion of soil-bound hexa-

6. POTENTIAL FOR HUMAN EXPOSURE

chlorobenzene particularly in children and from dermal contact with contaminated soils are needed, particularly in assessing health risks to populations living near hazardous waste sites.

Food Chain Bioaccumulation. Like many of the other organochlorine pesticides, hexachlorobenzene is lipophilic and has a high bioaccumulation potential. Hexachlorobenzene is bioaccumulated in fish and other aquatic organisms (Bishop et al. 1995; EPA 1992; Langlois and Langis 1995; Murray et al. 1980, 1981; Oliver and Nichol 1982a; Schmitt et al. 1990; Zabik et al. 1995) as well as waterfowl (Cobb et al. 1994; Foley 1992; Gebauer and Weseloh 1993; Somers et al. 1993; Swift et al. 1993; Yamashita et al. 1992). Hexachlorobenzene is bioaccumulated in aquatic food chains with virtually no degradation of the compound by the exposed organisms (Isensee et al. 1976). The results of a laboratory aquatic ecosystem study suggest that bioaccumulation as well as biomagnification of hexachlorobenzene occurs (Isensee et al. 1976). In terrestrial ecosystems, several agricultural crops have been found to accumulate hexachlorobenzene in their roots and in portions growing closest to soil level (Ecker and Horak 1994; Scheunert et al. 1983; Schroll et al. 1994; Smelt and Leistra 1974). The edible root portion of carrots accumulated the highest hexachlorobenzene concentration with a BCF of 19 (Smelt and Leistra 1974). Lichens, a primary forage for caribou, were also shown to bioaccumulate hexachlorobenzene (Muir et al. 1993). A field study on a terrestrial ecosystem suggested that hexachlorobenzene was biomagnified through various trophic levels of the food web (Hebert et al. 1994). Further studies are needed to resolve whether hexachlorobenzene is biomagnified in both aquatic and terrestrial ecosystems.

Exposure Levels in Environmental Media. Reliable monitoring data for the levels of hexachlorobenzene in contaminated media at hazardous waste sites are needed so that the information obtained on levels of hexachlorobenzene in the environment can be used in combination with the known body burden of hexachlorobenzene to assess the potential risk of adverse health effects in populations living in the vicinity of hazardous waste sites.

Environmental monitoring data are available for hexachlorobenzene in air (Currier et al. 1980; Davis and Morgan 1986; Eisenreich et al. 1981; EPA 1975b, Hoff et al. 1996; Lee et al. 2000b; Mann et al. 1974; Poissant et al. 1997; Tiernan et al. 1985), water (Chan et al. 1994; Davis and Morgan 1986; EPA 1976a; Hoff et al. 1996; Laska et al. 1976; Oliver and Nichol 1982a; Quemerais et al. 1994), soil (Elder et al. 1981; Laseter et al. 1976; Laska et al. 1976), and sediment (Davis and Morgan 1986; Elder et al. 1981; Murray et al. 1981; Oliver and Nichol 1982a; Ray et al. 1983; Rostad et al. 1999). Current information on hexachlorobenzene concentrations in groundwater is needed. Human intake estimates for exposure from environmental media are available (Whitmore et al. 1994), but are limited. In general, while

6. POTENTIAL FOR HUMAN EXPOSURE

monitoring data are available for most environmental media, much of the information on environmental levels was collected during the 1970s through the mid 1980s. More recent monitoring data from all environmental media would provide more accurate information for estimating human and animal intakes.

Exposure Levels in Humans. Hexachlorobenzene has been detected in human adipose tissue (Ansari et al. 1986; EPA 1986c; Frank et al. 1988; Kutz et al. 1991; Mes 1992), blood (Burns and Miller 1975; Burns et al. 1974; CDC 2009, 2015; Currier et al. 1980; Murphy and Harvey 1985; Rutten et al. 1988), and milk (Craan and Haines 1998; Frank et al. 1988; Newsome et al. 1995; Schecter et al. 1998; Takei et al. 1983). Studies exist that relate occupational exposure to blood levels of hexachlorobenzene (Burns et al. 1974; Currier et al. 1980). Studies to compare the steady-state intake of hexachlorobenzene as measured by urinary and fecal excretion as it relates to blood levels in occupationally exposed workers would be particularly useful. Since hexachlorobenzene has been detected in both urine and feces, a study of this nature could be conducted. These studies might also address possible individual differences in the metabolism of this compound.

This information is necessary for assessing the need to conduct health studies on these populations.

Exposures of Children. Exposure pathways for children have been well documented in breast milk (Newsome et al. 1995; Schecter et al. 1998) and diet (FDA 1992; Yess et al. 1993). Data documenting body burdens for children are needed. Data addressing exposure to children who live, play, or attend school near NPL sites, industrial sites, such as chlorinated hydrocarbon production factories, or on farmlands where hexachlorobenzene is being released as an impurity of another pesticide would allow for a better assessment of hexachlorobenzene exposure. As hexachlorobenzene is released due to the use of other pesticides on foods, an evaluation of possible hexachlorobenzene residues in children's food substances would further enhance the ability to construct a complete picture of exposure. Studies revealing contamination of drinking water or groundwater would also prove essential in this assessment. Attention should also focus on the use of tap water as a contaminant source when used to prepare infant formulas from condensed or powdered forms. As children are often bound to pick up soil off the ground and maybe even put this soil in their mouths, studies regarding exposure to children through soil would be helpful. Information concerning childhood-specific means to decrease exposure would be useful.

Child health data needs relating to susceptibility are discussed in Section 3.12.2, Identification of Data Needs: Children's Susceptibility.

6. POTENTIAL FOR HUMAN EXPOSURE

Exposure Registries. No exposure registries for hexachlorobenzene were located. This substance is not currently one of the compounds for which a sub-registry has been established in the National Exposure Registry. The substance will be considered in the future when chemical selection is made for sub-registries to be established. The information that is amassed in the National Exposure Registry facilitates the epidemiological research needed to assess adverse health outcomes that may be related to exposure to this substance.

Currently, Inuit communities appear to have potentially higher exposures to hexachlorobenzene and should be further monitored (Dewailly et al. 1993, 1999).

6.8.2 Ongoing Studies

No NIH or EPA ongoing studies regarding the potential human exposures to hexachlorobenzene were identified.