Rates and Routes of Transport of PCBs in the Environment

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Polychlorinated biphenyls (PCBs) are one member of a class of chlorinated organic compounds which give rise to concern, because of their wide dispersal and persistence in the environment and tendency to accumulate in food chains, with possible adverse effects on animals at the top of the food webs, including man. In the past, attention has been concentrated on chlorinated hydrocarbon pesticides, such as DDT, dieldrin, heptachlor and HCH (hexachlorocyclohexane). More recently attention has been focused on PCBs and on chlorinated dibenzodioxins. Likely candidates for future attention are hexachlorobenzene, chlorinated dibenzofurans, and chlorinated phenols. In the past, such compounds have generally been studied individually. As the number of compounds giving rise to concern increases, there is an urgent need to establish uniform monitoring schemes and generalized models describing environmental transport and bioaccumulation which will be applicable to all compounds with these properties. Models such as the global monitoring scheme outlined by the SCEP study (1), and the global transport model outlined recently by Woodwell et al. (2), are needed in order to help identify sources of environmental contamination, to establish acceptable levels of discharge, and to estimate the effectiveness of different control strategies.

This paper summarizes the fragmentary knowledge available about production, uses, and losses of PCBs, and attempts to define the major routes of transport and reservoirs of PCBs in the environment. It discusses only production and uses within North America, and distribution of PCB residues in North America and adjacent seas (the eastern half of the North Pacific Ocean and the western half of the North Atlantic). It is expected that the picture will be generally similar for other areas where PCBs have been used extensively (i.e., industrialized regions, primarily in the North Temperate Zone), with minor differences depending on local patterns of use and disposal. However, references to studies made in Europe and Japan are made in this paper where they help to fill gaps in the North American picture.

Although we attempt to cover all the important environmental aspects of the use and distribution of PCBs, much of the discussion in this paper is extremely speculative. One of its main purposes is to point out the most important gaps in knowledge which must be filled before firm estimates can be made of present and future levels in the environment.

Production and Use in North America

The sole manufacturer of PCBs in North America is the Monsanto Company. Commercial mixtures are sold under the trade name Aroclor and are distinguished by numbers, in which the first two digits 12 specify polychlorinated biphenyls and the last two digits the approximate percentage of chlorine in the mixture (3). In November, 1971, Monsanto released figures for U.S. domestic sales of Aroclors during the period 1963–70, summarized here in Figs. 1 and 2, where they are divided according to category of use and grade of Aroclor.

Prior to Monsanto’s voluntary reduction of sales in September, 1970, approximately 60% of
sales were for closed-system electrical and heat transfer uses, 25% for plasticizer applications, 10% for hydraulic fluids and lubricants, and less than 5% for miscellaneous applications such as surface coatings, adhesives, printing inks, and pesticide extenders. Exports by Monsanto have averaged 13% of domestic sales over the period 1963 to 1970. Imports are thought to be small, comprised primarily of plasticizers in resins and adhesives, transformer oils and capacitor fluid in electrical equipment. Exports to Canada should be of the order of 7% of U.S. sales and are included in this model of North America.

Monsanto has restricted sales of Aroclors based on consideration of either the possibility of contamination of food products or its inability to control or monitor possible losses into the environment. The fraction of sales for use in confined systems, primarily electrical applications, will increase to approximately 90 percent in 1971 and to 100% in 1972. The higher chlorinated Aroclors, 1254 and above, comprised approximately 30% of sales prior to 1971. Their share of the reduced sales in 1971 is expected to fall to 22%. Aroclor 1242 is being replaced with Aroclor 1016 having a similar composition but with isomers containing 5 or more chlorine atoms removed.

The breakdown by grade of the current uses of PCB is: electrical capacitors, mainly Aroclor 1016 with limited usage of Aroclor 1221 and Aroclor 1254; electrical transformers, Aroclors 1242, 1254 and 1260; vacuum pumps, Aroclor 1248 and 1254; and gas transmission turbines, Aroclor 1221 and 1242. The breakdown by PCB grade of former uses include Aroclors 1232, 1242, 1248, 1254, and 1260 for hydraulic fluids; Aroclor 1242 for heat transfer systems; Aroclors 1248, 1254, 1260, 1262, and 1268 for plasticizers in synthetic resins; Aroclors 1221, 1232, 1242, 1248 and 1254 for adhesives; Aroclors 1221, 1232, 1242, 1248, 1254 and 1268 for plasticizers in rubbers; Aroclors 1242, 1254 and 1268 for wax extenders; Aroclors 1254 and 1260 for dedusting agents; Aroclor 1254 for pesticide extenders, inks, lubricants, and cutting oils; and Aroclor 1242 for carbonless reproducing paper.
Data on production outside the U.S. are not available. It has been estimated (4) that the Japanese production is 26 million pounds per year, 40–50% for capacitors, 15% for transformer oil, 10–15% for heat transfer fluid, 5% for plasticizers, 15% for carbonless duplicating paper, and 5—10% for export.

Routes into the Environment

As in the case of most industrial chemicals, loss figures for PCBs are practically non-existent. Possible routes into the environment include: (1) leaks from sealed transformers and heat exchangers; (2) leaks of PCB-containing fluids from hydraulic systems which are only partially sealed; (3) spills and losses in the manufacturing either of PCBs or PCB-containing fluids; (4) vaporization or leaching from PCB-containing formulations; (5) disposal of waste PCBs or PCB-containing fluids.

Examples of losses by each of these proposed routes include leaks from faulty heat exchangers in the contaminated Japanese rice oil (5) and contaminated chicken feed (Holly Farms) incidents; leakage of hydraulic fluid from an air compressor in Escambia Bay, Florida (6); the indirect evidence of possible losses during manufacture provided by the high level of PCBs in catfish in waters near Anniston, one of the two sites at which PCBs are manufactured in the U.S. (7); the leaching of PCBs from silos by sileage (7) and the use of waste electrical insulator containing PCBs as solvent in herbicide treatment of power rights-of-way near Martinsburgh, West Virginia (7). Other unreported disposal of scrap undoubtedly occurs into sewers (8). Statistical information on such losses is not available but could be generated by an accounting of PCB inventories and disposal by major users. To meet the problem of scrap disposal, Monsanto has set up a disposal system with a capacity of 10 million pounds per year for their customers. Within a year of announcement of the service, 500,000 pounds of waste PCBs had accumulated at the disposal site, where it was held in storage, pending the completion of an incinerator (9).

Another waste disposal problem is that of PCB-containing products including the PCB-impregnated paper in capacitors, caulking compounds, and carbonless duplicating paper. It has been postulated that the PCBs may be vaporized during incineration. Based on studies into the incineration of PCBs and pesticide residues, municipal incinerators meeting design guidelines of a residence time of 2 seconds at 2000°F are expected not to be a significant source of PCBs (9, 10). Poorly operated commercial and municipal incinerators, small domestic and apartment incinerators, and open-burning dumps may, however, be major sources of emission, but data are not presently available. Another problem is that of leaching from dumps. Again, data are scarce, but analysis (11) of stagnant water close to a sanitary landfill indicated levels below the detection limit of 4 ppb.

Quantitative Estimates of Rates of Loss into the Environment

It is unlikely that reliable quantitative estimates of past losses of PCBs into the environment will ever become available. However, in order to assess the biological significance of the PCBs now in the environment, it is necessary to make some numerical estimates, however rough. The following calculations are intended to provide order-of-magnitude estimates of rates of loss. Figure 3 shows qualitatively some of the many possible routes of loss into the environment.

Estimates of the rates of loss and disposal of Aroclors in 1970 will be based on rough guesses of the useful service life in different applications. Transformers are fairly permanent installations, and it is therefore estimated that only 10% of sales of transformer fluid is to replace oil that was scrapped and that the remaining 90% is for new units. The useful life of capacitors, especially those used in small items such as fluorescent light ballasts, is expected to be under a decade, and it is therefore estimated that the rate at which the capacitors are discarded, primarily into landfill dumps, is equal to half the rate of production. (Figure 1 indicates a doubling period for sales of about 10 years.) A similar figure is assumed for the scrapping of heat exchangers, but it is expected that the replacement of fluid decomposed under extreme thermal conditions would have led, in the past, to additional direct losses, primarily into sewers. It is estimated that the rate of vaporization of plasticizer amounts to 10 to 20% of
sales (12). Since many plastic objects have relatively short useful lives the rate of disposal of plasticizers into dumps is assumed to equal the residual 80 to 90% of sales. Hydraulic fluids and lubricants are rarely re-used, and it is therefore assumed that a major fraction of these fluids used for this purpose, together with those going into the miscellaneous applications, were scrapped at rates approximately equal to those of corresponding sales.

On the basis of these gross estimates, it appears that only about 20% of the 1970 sales in North America, some $7 \times 10^3$ tons, represented a net increase in the amount of PCBs in service, in transformers, heat exchangers and capacitors. The remainder is assumed to have been discharged into the environment, 1–2$\times 10^4$ tons by evaporation of plasticizers; 4–5$\times 10^3$ tons by leaks and disposal of hydraulic fluid and lubricant plus small amounts by disposal of heat transfer and trans-
Most of the PCBs discharged into the atmosphere will have been Aroclor 1248 to 1260 vaporized from plastic resins, augmented primarily by 1242 vaporized from burning dumps. The discharge into waters will be heavily weighted by the Aroclors used as hydraulic fluids and lubricants and is therefore likely to have included a mixture of Aroclors 1242 to 1260. The residual in dumps will have a large fraction of the Aroclor 1242 production.

Rates of Transport within the Environment

The modes of transport of the PCBs within the environment are complex. Vaporized PCBs will be partially adsorbed on particulates, transported with the prevailing winds, and deposited on land or water by particle sedimentation or rain-out. PCBs introduced into water streams may be adsorbed by the waterborne particulates or the benthos; the adsorbed PCBs will diffuse into the bottom sediment, redissolve in the water stream or be entrained with sediment eroded from the bottom surface.

The problem is further complicated by the assimilation, transport and degradation of PCBs by the biota. A major fraction of the PCBs discarded in dumps is encapsulated in sealed containers or plastic resins. The rate of loss from these will therefore be low until the confining material is degraded and the PCBs released. The PCBs will then slowly diffuse through the surrounding soil. In principle, the rates of transport may be calculated from knowledge of the physicochemical properties of the PCBs and the pertinent data on atmospheric conditions, particulate transport, hydraulic dispersion, bottom sediment transport, and biological degradation rates. At present, the data are too incomplete and the interactions between the different elements in the environment too complex to attempt a formulation of but the crudest of transport models. The similarity between the properties of the PCBs and DDT, summarized in Table 2, permits rough estimation of some of the otherwise unknown routes and rates of transport of PCBs from the corresponding information on DDT. In Fig. 4, a generalized model for the distribution and transport of the PCBs is outlined, without any indication of the transport down spatial gradients within the differ-

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**Table 1. Gross Estimates of Rates of Input and Accumulation of PCBs in North America in 1970.**

| Category of input               | Rates (tons/year) | PCB Grade |
|---------------------------------|-------------------|-----------|
| Vaporization of plasticizers    | 1–2×10⁴           | Mainly 1248 to 1260 |
| Vaporization during open-burning| 4×10⁴             | Mainly 1242 |
| Leaks and disposal of industrial fluids | 4–5×10⁴ | 1242 to 1260 |
| Destroyed by incineration and open burning | 3×10⁴ | Mainly 1242 |
| Disposal in dumps and landfills | 1.8×10⁴           | 1242–1260 |
| Accumulation in service         | 7×10⁴             | 1242–1254 |

| Reservoir                 | Accumulation (tons) |
|---------------------------|---------------------|
| Soil (excluding dumps)    | PCPs 1.5×10⁴, ΣDDT 3×10⁴ |
| Oceans (adjacent to North America) | 1.5×10⁴, 10⁴ |
| Fresh water (dissolved or in suspension) | 10⁴, ~10⁴ |
| Fresh water sediment      | 2×10⁴              |
| Biota                     | <10⁴, <10⁴         |

former oils, and 22×10³ tons by disposal in incinerators, dumps and sanitary landfills. Of the latter, we estimate (13) that 10 to 20% (3×10⁸ tons) were destroyed by burning, and 2% (4×10⁸ tons) were vaporized, mainly by open burning of wire scrap, auto components, and material in dumps.

**Routes into the Environment**

The total rate of loss of PCBs is thus estimated to have been of the order of 1.5 to 2×10⁴ tons/year into the atmosphere, 4 to 5×10³ tons/year into fresh and coastal waters, and 1.8×10⁴ tons/year into dumps and landfills. The input into soils via the use of Aroclors as pesticide extenders is believed to have been small, less than 10 tons/year, on the basis of reports of purchases for this purpose. The unauthorized and unrecorded use of scrap PCBs as pesticide extenders is difficult to estimate but has probably been small. Direct discharge into the oceans, e.g. by dumping of hydraulic fluids and lubricants from ships, has probably been of relatively small magnitude, but may have been locally significant (49).
Table 2. Comparative Physical Properties of PCBs and DDT (Ref. 9, 23, 84, 85).

| Molecular weight | 1242 | 1248 | 1254 | 1260 | DDT |
|------------------|------|------|------|------|-----|
| Range*           | 154-358 | 222-358 | 290-392 | 324-460 | 352 |
| Average          | 262   | 288   | 324   | 370   | 352 |
| % Chlorine       | 42    | 48    | 54    | 60    | 50  |
| Solubilityb in H2O (at 20°C ppb) | 200 | 100 | 50 | 25 Est. | 0.7 |
| Vapor pressureb at 38°C (mm, Hg) | $10^{-3}$ | $3.7 \times 10^{-4}$ | $6 \times 10^{-5}$ | $2 \times 10^{-7}$ | $2 \times 10^{-10}$ |
| Vapor pressure at 20°C (mm, Hg) | $10^{-4}$ | $3 \times 10^{-4}$ | $3.6 \times 10^{-6}$ | $1.5 \times 10^{-7}$ |

* Based on constituents present in amounts of 1 percent or more.

b The high values of the solubilities and vapor pressures of the PCBs are a consequence of the heavy weighting given the lower chlorinated species.

e Extrapolated.

ent compartments of the environment. An attempt is made in the following sections to estimate fluxes between and within each compartment.

**Air Transport**

The estimated emission of 1500 to 2500 tons a year of PCBs, mainly by vaporization and open burning, is expected to be concentrated in urban areas. By analogy to DDT (17, 18) it is expected that most of the airborne PCBs will be adsorbed on particles. Since roughly three quarters of total suspended particulates (TSP) in urban areas are of non-agricultural origin (19), and the emission of TSP within cities is of the order of $2 \times 10^7$ tons per year (20), this gives rise to a rough estimate of PCB level on urban particulates of 50 to 80 ppm.

The half-life of particulates in the air will depend greatly on the size of particles to which the PCBs are attached and the extent of atmospheric precipitation. (19) It is estimated that most of the vaporized PCBs will be deposited within 2 or 3 days, mostly onto the land mass and coastal waters surrounding urban areas. The small amounts of PCB attached to fine particulates will reside in the atmosphere for extended periods and be transported to remote areas.

Based on the above assumptions, the rate of terrestrial net input of PCBs by serial fallout in North America will be a little less than the rate of vaporization, about 1000 to 2000 tons/year. The vapor pressure of Aroclor 1254, the dominant airborne contaminant, is close to that of DDE (21), and it is estimated, by analogy to DDE (17) that the likely half-life of PCBs in soil is of the order of five years. The PCBs removed from the soil

![Figure 4. Environmental transport model.](Image)
will become adsorbed to particulates, mostly of relatively short range, and be redeposited and vaporized one or more times until they reach the coast. By analogy to DDE (1, 2), the redistribution of PCBs by this mechanism is expected to have resulted in the redeposition into the ocean of one quarter of the PCBs that originally fall out in the terrestrial environment. The deposition pattern will be influenced by the size distribution of particles to which the PCBs are attached, with a major fraction ending in coastal waters and a small amount being transported into remote regions.

It is expected that a negligible fraction of the PCBs reaching the oceans will be revaporized; their concentration will be too low.

**Water Transport**

On account of the low water solubility and high specific gravity of PCBs, it is expected that most of the PCBs discharged into the environment will be resting as sludges or adsorbed in the sediment at the bottom of rivers or lakes near their point of discharge, and that transport in streams will be primarily by means of waterborne particles. Evidence for this postulate is indirect, consisting mainly of the strong adsorption of PCBs by surfaces (22), the observation of PCB concentrations as high as five times the solubility limit (6), and the evidence that chlorinated hydrocarbon pesticides are removed from contaminated lakes by adsorption in sediments and codeposition with algal bloom (24).

The transport of PCBs within rivers is as described previously, by solution and readsorption in the sediment and by sediment transport. The data on partition coefficient between sediments and water and on diffusivity in bottom sediment are not available so that the transport models (25) developed for rivers cannot yet be applied. The only measurements reported on PCBs in ground sediment are for Escambia Bay (6, 26). The PCB concentration in the sediment 10 miles downstream from a discontinued source of PCB loss was found to remain constant over a year suggesting that the PCBs are strongly adsorbed to sediment.

The amount of PCBs transported to the ocean in solution or suspension in river water is estimated as 200 tons/year (49), a small amount compared to the estimated input into rivers. Irrigation waters amount to about 1/15 of the total river flow and are expected to carry a correspondingly lower amount, about 15 tons/year, of PCBs to the terrestrial environment.

**Dredging and Dumping**

Dredging of inland rivers and harbors may lead to significant transfer of PCBs from contaminated sediments, especially when the dredge spoils are dumped at sea. According to the Council on Environmental Quality (27), some $13 \times 10^6$ tons of polluted dredged spoils are dumped at sea annually off the United States, 90% off the Gulf and Atlantic coasts. A high figure for PCB levels in polluted sediments is 30 ppm, the highest level recorded in the Escambia Bay incident away from the immediate vicinity of the discharge (26). If all the polluted dredge spoils were contaminated to this level, the total rate of transfer of PCBs to the oceans would be around 400 tons/year, mostly into the Atlantic Ocean and Gulf of Mexico. Based on the general level of contamination a more reasonable estimate of rate of transfer would be only 20 tons/year.

**Biota Flux**

The total quantity of PCBs stored in migratory animals is very small (see below), and their movements cannot account for a significant fraction of environmental transfer of PCBs. The movements with the greatest local significance are probably the migrations of seabirds from the northern to the southern hemisphere; these might account for the movement of about a ton/year (28). Human fishing is estimated to remove only 1–2 tons/year from the eastern North Pacific, the western North Atlantic and the Great Lakes (30).

**Separation and Transformation of PCB Isomers in the Environment**

There are three processes by which the compositions of PCB mixtures may change after release into the environment.

**Fractionation**

In general, the water solubilities and vapor pressures of PCB isomers decrease with increasing
Table 3. Percent Loss in Area of Seven Chromatogram Peaks of Aroclor 1254 After Steam Heating for 25 and 60 Minutes. [From Ref. 23]

| Peak | 25 Minutes | 60 Minutes |
|------|------------|------------|
| 1    | 66         | 83         |
| 2    | 41         | 74         |
| 3    | 22         | 73         |
| 4    | 40         | 54         |
| 5    | 14         | 51         |
| 6    | 0          | 15         |
| 7    | 0          | 33         |

chlorine content, although the decreases are not uniform (Table 2) (31). Hence the processes of evaporation, co-distillation and dissolution in water are expected to fractionate mixtures of PCBs, the lower isomers being much more mobile. This has been confirmed for co-distillation by the results of Freed (21, 23) (Table 3), which show that, after 60-min steam heating, the ratio of the first to the sixth of the seven major chromatographic fractions of Aroclor 1254 was reduced by a factor of 5. Thus environmental transport mechanisms involving these processes are expected to reduce the proportions of the lower isomers in environmental samples near to the point of release, and to augment them in samples from remoter areas.

Photolysis and Chemical Decomposition

Photolytic decomposition of certain PCB isomers has been reported by Risebrough et al. (32, 33), and Safe and Hutzinger (34, 35). The results of the latter suggest that in natural sunlight some higher isomers are more easily broken down than the lower isomers (35). Hence photolysis is expected to reduce the proportions of at least some of the higher isomers. Photolytic dechlorination is also expected to give rise to lower isomers, including some which may not be present in commercial mixtures (35).

Because of their high stability in industrial use, we assume that other forms of chemical (non-biological) decomposition of PCBs are very slow in the environment.

Metabolism and Excretion

Published studies of the dynamics of PCBs in rats and birds (36–39) indicate that the higher isomers are generally taken up and/or retained more efficiently. In each case the proportions of pentachloro- and lower isomers found in the animals' tissues were generally lower than those in the Aroclor mixture to which they had been exposed, but there appears to have been little differentiation of the higher isomers except for one hexachloro-isomer (36–38). It is not clear in each case whether the deficiencies of the lower isomers resulted from metabolism or differential excretion, but it is likely that both occurred. Because of their higher solubility in water (see above), the lower isomers would be expected to be excreted more easily than the higher isomers. In one study on rats, loss of PCBs from the body was markedly retarded by administration of carbon tetrachloride (36), suggesting interference with metabolism in the liver. In another study with rats, hexachloro-isomers were found in disproportionate amounts in the urine (37), suggesting that the tetra- and pentachloro-isomers were metabolized and the hexachloro-isomers excreted in part, while the heptachloro-isomers were differentially retained. Two studies of metabolism of PCBs in fish (40, 41) gave inconsistent results but showed no marked tendency for the higher isomers to accumulate differentially.

Evidence from Environmental Samples

Veith (42) has found that the proportion of higher isomers increases downstream in some Wisconsin rivers and that certain isomers characteristic of lower Aroclors disappear. Since the lower isomers are unlikely to be differentially retained in sediments (see above), this suggests that they are decomposed rapidly in the river environment. Evidence of biodegradation of pesticides under anaerobic conditions (43) suggests the possibility that the degradation of the missing lower isomers may occur primarily by microbial metabolism in the bottom sediment. We recommend that higher priority be given to this aspect of decomposition of PCBs.

A number of writers have reported that PCB samples extracted from animals generally match Aroclors 1254 or 1260, but that the lowest isomers are sometimes relatively deficient or missing, especially in animals high in food chains (38, 44). Since substantial quantities of Aroclors 1242 and
isomers and phenyls with birds. Mammals samples have sewage grams and/or isomers is comparison lowest to higher relatively ability. April 1248 Canada. (32, 38, 48), however, Relative Table 4. Relative Peak Heights in Gas-Liquid Chromatograms of Extracts from Sewage Sludges. [From Ref. 45]

| Relative retention time | Sludge A | Aroclor 1254 | Sludge B | Aroclor 1260 |
|-------------------------|----------|--------------|----------|--------------|
| 0.69                    | 38       | 30           | 39       | 3            |
| 0.81                    | 63       | 72           | 24       | 14           |
| 1.00                    | 51       | 59           | 17       | 4            |
| 1.21                    | 104      | 110          | 51       | 30           |
| 1.45                    | 100      | 100          | 100      | 100          |
| 1.71                    | 92       | 92           | 66       | 69           |
| 2.00                    | 32       | 26           | 70       | 70           |
| 2.34                    | 19       | 12           | 53       | 55           |
| 2.83                    | 21       | 19           | 84       | 87           |
| 3.33                    | 10       | 8            | 29       | 30           |
| 3.70                    | 5        | 4            | 25       | 23           |

1248 must have been released into the environment (see above), these observations suggest that a large proportion of the lower isomers (those with 4 or fewer chlorine atoms) is missing from the animal samples. However, few critical data have been published. In one set of data from sewage sludge (45) (Table 4), none of the lower isomers is clearly deficient. In published chromatograms for birds (38, 46, 47) (Table 5), only the lowest isomers (tetrachlorobiphenyls) and one higher isomer were markedly reduced in comparison to Aroclor 1254. In human adipose tissue (48), however, pentachloro- and hexachloro-isomers were reduced or missing.

The higher isomer reduced in the bird samples (38, 46) is probably that reported as deficient in samples from the Gulf of California and Quebec and found to be degraded by ultraviolet light (32, 33). With this one exception, we conclude tentatively that higher isomers (pentachlorobiphenyls and higher) are not significantly differentiated as they pass through food chains up to fish and birds. Mammals appear to be able to excrete and/or metabolize penta- and hexachloro-isomers.

The lack of the lower isomers in samples from relatively contaminated areas could be accounted for either by metabolism or by their greater mobility. However, Risebrough and Berger (33) found that lower isomers were relatively deficient also in samples from fish in a remote lake in northern Canada. This suggests that differential metabolism is the primary mechanism in the environmental differentiation of isomers, outweighing the effects of differential photolysis and differential mobility, which would be expected to have reduced the proportion of higher isomers in remote samples.

Conclusions and Comments on Separation of PCB Isomers in the Environment

We conclude that most PCB isomers with four or fewer chlorine atoms have been degraded in the environment, possibly by microbial action. Decomposition of penta- and hexachloro-isomers appears to occur in birds and mammals, but this will have affected only a negligible fraction of the PCBs in the environment. Accordingly, we assume (3) that some 75% of the Aroclor 1242 released into the environment, 60% of Aroclor 1248, 20% of Aroclor 1254 and 5% of Aroclor 1260, have disappeared.

The above discussion has been confined to a general comparison of lower and higher isomers, because of the scarcity of data on relative proportions of individual isomers in environmental samples. In fact, the physical and chemical properties and biodegradability of PCB isomers depend on the positions of substitution as well as the total chlorine content. More precise data on the

| Peak No. | Relative retention time* | Eagle carcass | Aroclor 1254 |
|---------|-------------------------|---------------|--------------|
| 1       | 0.39                    | 7.5           | 6.8          |
| 2       | 0.49                    | 34.8          | 100.0        |
| 3       | 0.53                    | 55.0          | 42.3         |
| 4       | 0.58                    | 15.7          | 21.0         |
| 5       | 0.70                    | 68.6          | 191.5        |
| 6       | 0.79                    | 28.0          | 32.2         |
| 7       | 0.83                    | 111.9         | 127.5        |
| 8       | 0.97                    | 68.7          | 66.5         |
| 9       | 1.04                    | 91.6          | 105.2        |
| 10      | 1.23                    | 92.5          | 139.5        |
| 11      | 1.46                    | 100.0         | 100.0        |
| 12      | 1.72                    | 62.2          | 75.5         |
| 13      | 2.01                    | 28.7          | 23.6         |
| 14      | 2.32                    | 18.4          | 42.6         |
| 15      | 2.86                    | 32.6          | 38.8         |
| 16      | 3.38                    | 15.4          | 22.2         |

* p,p'-DDE = 1.00

Table 5. Relative Peak Areas (Peak 11 = 100) for Chromatogram of Extract of Eagle Carcass and Aroclor 1254. [From Ref. 47]
exact quantities of individual isomers in environmental samples would permit more precise conclusions about the sources and routes of contamination. We urge that numerical characterization of individual peaks should be published whenever possible.

**Cumulative Input into the Environment**

From an extrapolation of the sales curve in Fig. 1 to zero production in 1930, it is estimated that the cumulative sales in North America over the period 1930–1970 were $5 \times 10^5$ tons. Assuming that the proportions of sales for different uses were similar throughout the period, the cumulative losses may be estimated as about $3 \times 10^4$ tons into the air, $6 \times 10^4$ tons into fresh and coastal waters, and $3 \times 10^6$ tons into dumps and landfills. Using the results of the previous section, it may be estimated that roughly one-third of the PCBs released into the air and one-half of those released into water have now been degraded. It is difficult to estimate the extent of degradation in dumps, because some of the PCBs there may still be in sealed containers, but we assume in any case that leaching from dumps into fresh waters has so far been negligible.

Of the PCBs released into the air, we have assumed that most would have been adsorbed onto particulates soon after volatilization and would therefore have fallen out relatively close to the source. By analogy with DDT residues, PCBs in soil would be expected to have a half-life of the order of 5 years (2, 17). Again using the analogy with DDT and its metabolites, we may estimate that roughly one-quarter (1) of the total (i.e., $5 \times 10^6$ tons) will have been transferred into the sea, mainly into the Atlantic Ocean, and the remainder ($1.5 \times 10^4$ tons) distributed over terrestrial North America, with a negligible fraction having fallen into fresh waters. Other inputs to the sea are difficult to estimate but have probably been of the order of $10^4$ tons (49). The remainder ($2 \times 10^4$ tons) is assumed to have accumulated in lakes and rivers.

**PCB Levels and $\Sigma$ DDT/PCB Ratios in the Environment**

In this section we compare the above estimates of cumulative input into the environment with measurements of PCB levels in physical and biological samples. The purposes of this comparison are: (1) to use the observed distribution of PCBs in the environment to check the consistency of our

![Figure 5. Ocean sub-model.](image)
models of environmental transport; (2) to define the types and locations of input to the environment which are responsible for the most serious contamination of the biota.

### Uptake and Biological Magnification of PCBs in Food Chains

A large fraction of the reported measurements of PCB concentrations in environmental samples have been from animals, especially fish and birds high in marine food chains. Figure 5 outlines schematically a model for the transport of PCBs within a simple marine ecosystem. Laboratory experiments (6, 41, 55) indicate that aquatic invertebrates and fish can accumulate PCBs to levels between $3 \times 10^3$ and $7 \times 10^4$ times higher than those in the ambient water. It is not yet clear whether concentrations of PCBs are consistently increased as they pass up food chains from invertebrates to fish and from fish to fish (57). However, PCB levels are clearly magnified within food chains involving birds and mammals by a factor on the order of 10 to 100 at each step (32, 44, 56, 57, 58). Hence, in the long food chains characteristic of marine systems (59), the levels in the top predators may be $10^9$ times higher than those in the ambient water (Tables 6 and 7). In extreme cases, such as the starved guillemots in Table 6, or the eagle fat in Table 7, the concentration factors may be as high as $10^9$ or $10^8$. Because of the extreme variability observed both within and between samples, it is difficult to use measured levels in animals as a direct measure of levels in the environment. However, it is possible to compare levels in the same species or in ecologically equivalent species to define spatial gradients in contamination (29, 44).

### ΣDDT/PCB Ratios

Another approach (32, 56) utilizes the similarity in properties between the persistent PCBs and the persistent metabolites of DDT, primarily DDE. (In the remainder of this section, DDT and its metabolites DDE and DDD are grouped together as ΣDDT). Experiments suggest that PCBs and ΣDDT are accumulated by aquatic animals to an extremely similar degree from ambient water (41), and measurements in the same areas indicate that the ratios ΣDDT/PCB are very similar in animals from different levels in the food chain (Tables 6 and 7) (32, 56, 75). Hence the ratio ΣDDT/PCB in environmental samples, even in animals at the top of food chains, can be used to infer the ratio in the substrate, at least to order of magnitude (32, 56). Then the extensive information about the environmental distribution and transport of ΣDDT (2, 17) can be used to make inferences about the distribution and transport of PCBs.

### Air

The only numerical measurements available of PCB levels in air are some estimates of levels in total suspended particulates (TSP) in four U.S. cities between 1968 and 1970 (E.P.A., unpublished data). The mean level of PCBs was about 50 ppm, with levels in one city substantially lower than others.

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Table 7. PCB levels in Swedish marine organisms 1965–68 [From Ref. 44]. Figures given are the mean (ppm in extractable fat), with range and sample size in parentheses; ΣDDT is the total of DDT and its metabolites.

|                     | Baltic Sea |                             | Stockholm Archipelago |                             |
|---------------------|------------|------------------------------|-----------------------|------------------------------|
|                     | Mean       | Range                        | ΣDDT/PCB              | Mean                        | Range                        | ΣDDT/PCB              |
| Mussel              | 4.3        | (1.9–8.6)                    | (40)                  | 5.2                         | (3.4–7.0)                    | (15)                  |
| Herring             | 6.8        | (0.5–23)                     | (18)                  | 5.1                         | (3.3–8.5)                    | (4)                   |
| Seal                | 35.        | (16–44)                      | (3)                   | 30.                         | (16–56)                      | (3)                   |
| Guillemot eggs      | 250.       | (140–360)                    | (9)                   | 9400.                       |                              | (1)                   |
| Heron               |            |                              |                       | 14000.                      | (8400–17000)                 | (4)                   |
| White-tailed Eagle: |            |                              |                       | 910.                        | (490–1500)                   | (3)                   |
| breast muscle       |            |                              |                       | 540.                        | (250–800)                    | (5)                   |
| brain               |            |                              |                       |                              |                              |                       |
| eggs                |            |                              |                       |                              |                              |                       |
higher. This is at least consistent with our earlier estimate based on rates of transport in the environment.

Risebrough et al. (60) were unable to detect PCBs in a sample of marine air off the California coast. This suggests that long-range (intercontinental) transport of PCBs in air may be negligible.

Land

We have estimated the cumulative net input into terrestrial North America as of the order of $1.5 \times 10^6$ tons. For comparison, the total quantity of $\Sigma$DDT remaining in terrestrial North America is of the order of $3 \times 10^4$ tons, as estimated by two methods: total application minus estimated total losses to the sea (1); mean levels of 2–3 ppm in a wide variety of treated soils, and lesser amounts in untreated soils (2, 17, 61). Hence the continental mean $\Sigma$DDT/PCB ratio should be of the order of 20. However, PCBs are expected to be distributed primarily around urban areas and downwind from them, whereas $\Sigma$DDT is still concentrated in the agricultural and forested areas where it was applied (61).

A nation-wide monitoring scheme using starlings (Sturnus vulgaris) and woodcocks (Philohela minor) is in progress. Early results show mean PCB levels in pooled samples of fat from woodcock wings from eastern states in the range 4–7 ppm (62). $\Sigma$DDT levels from these pools are not yet available, but fat from woodcocks in Canada contains 6–130 ppm, depending on the local history of application (29). Woodcocks feed on earthworms and should be a good indicator for soil concentrations.

Other data from terrestrial birds may be biased towards low $\Sigma$DDT/PCB ratios, because the birds were obtained in industrialized regions. The median $\Sigma$DDT/PCB ratio in a small sample of terrestrial birds from California was 4 (32). Terrestrial birds of prey in five European countries have very high levels of PCBs, with $\Sigma$DDT/PCB ratios in the range 0.3–5 (63–67).

In human food, market basket surveys in the United States suggest a $\Sigma$DDT/PCB ratio of about 4 (68). Ratios in human milk (69, 70) and human blood plasma (71) are in the same range. However, except for sporadic instances of contamination, most of the human intake of PCBs appears to be in fish (68), whereas $\Sigma$DDT is much more widely distributed in the diet (72). If the fish and shellfish portions of the diet are excluded, the $\Sigma$DDT/PCB ratio would probably be greater than 10. Similar data have been reported from Sweden, where the $\Sigma$DDT/PCB ratio in total diet samples and in human milk is about 7 (73).

Fresh Waters

Measurements of PCB levels in water are available only for relatively polluted rivers and bays and are summarized in note (50). Table 8 summarizes PCB levels and $\Sigma$DDT/PCB ratios in freshwater fish and birds. PCB levels are both absolutely and relatively high in industrial rivers and in the Great Lakes; within the Great Lakes there is a gradient in $\Sigma$DDT/PCB ratios from west to east. Levels of PCBs are lower in the three samples from lakes in less industrialized areas listed at the bottom of Table 8. These data are consistent with our conclusion that most PCBs in fresh water systems result from local discharges by industries, but that there is also widespread aerial fallout of small quantities.

Based on our assumptions of mean concentrations in fresh waters (50), we estimate that the water of the Great Lakes contains a quantity of PCBs of the order of 100 tons; the total quantity

### Table 8. PCB Levels and $\Sigma$DDT/PCB ratios in North American Fresh Water Vertebrates.

| Area                     | Mean PCB level in fish (ppm) | $\Sigma$DDT/PCB ratio in fish and/or fish-eating birds | Reference |
|--------------------------|-------------------------------|------------------------------------------------------|-----------|
| U.S.: industrialized rivers | 1–213                         | 0.005–0.35                                           | 55        |
| Lake Ontario             | 19                            | 0.05–0.2                                             | 55, 74    |
| Lake Michigan            | 20                            | 0.4–1                                                | 55        |
| Lake Huron (Georgian Bay)| —                             | 0.8                                                  | 75        |
| Lake Superior            | —                             | 2.4                                                  | 75        |
| Ontario:                 |                                |                                                      |           |
| Lake Nipigon             | 0.1                           | 1–4                                                  | 75        |
| U.S. & Canada:           |                                |                                                      |           |
| Prairie Lakes            | <0.1                          | 2                                                    | 76        |
| N. Quebec:               |                                |                                                      |           |
| Lake Minto               | 0.1                           | 1                                                    | 33        |
in other lakes would be negligible. The quantity carried in river water would be only of the order of 3 tons. It is difficult to estimate the total quantity in the biota, but since the Great Lakes' fisheries probably remove less than a ton of PCBs per year (30), the total quantity in fish is probably only of the order of 10 tons. These quantities are negligible in comparison to the estimated cumulative input of $2 \times 10^4$ tons after allowing for degradation. Hence we conclude that almost all the PCBs that have been released into fresh waters are adsorbed onto bottom sediments. There are no data to indicate the rate at which they are being released into water and into the biota or being biodegraded.

It is expected that PCB levels in ground water will be negligible, since any PCBs in water percolating through the soil should be adsorbed onto soil particles (77).

The Sea

According to our estimates, the total input of PCBs into the seas around North America has been of the order of $1.5 \times 10^6$ tons, mostly into the Atlantic Ocean, in part by localized discharge and in part by aerial fallout. For comparison, the total input of $\Sigma$DDT into the oceans has been estimated as of the order of $5 \times 10^6$ tons (1), of which a disproportionate quantity, probably more than $10^6$ tons, would have fallen into the North Atlantic Ocean, because of the heavy use of DDT in North America (17). Thus, on these arguments, the total quantity of $\Sigma$DDT in the oceans would exceed that of PCBs by several times, especially in the eastern North Pacific. However, $\Sigma$DDT has been introduced over large areas (1, 2), so that it should be more uniformly distributed than the PCBs, the discharges of which are localized near industrialized coasts.

Available measurements of PCB levels and $\Sigma$DDT/PCB ratios, summarized in Table 9, are generally consistent with these predictions, although the pattern is less clear than that in Table 8, in part because of the wide variability between species (cf. Tables 6 and 7). The highest PCB levels and the lowest $\Sigma$DDT/PCB ratios have generally been in areas close to industrial activity. At least in the Pacific, PCB levels decrease and $\Sigma$DDT/PCB ratios increase with increasing distance offshore. Data on marine birds summarized by Keith and Gruchy (29) show that these offshore gradients exist in both the Atlantic and Pacific, and show further that PCB levels in the Atlantic seabirds are 5–10 times higher than those in the same or ecologically equivalent species in the Pacific. The correlation of high PCB levels with industrial activity has been demonstrated most clearly in Europe, where it has been observed along the coasts of Sweden (65, 81), the Netherlands (38) and Great Britain (82).

The only major discrepancy from this general picture is the observation (57) of extremely low $\Sigma$DDT/PCB ratios (of the order of 0.01) in zooplankton over a wide area in the North Atlantic Ocean. $\Sigma$DDT/PCB ratios in fish in the same area were of the order of 0.2 to 0.5. It is almost inconceivable that aerial fallout of PCBs could have exceeded that of $\Sigma$DDT by such large factors over such a large area; hence to explain the discrepancy, it is necessary to assume either selective removal of $\Sigma$DDT or a local source of PCBs. A likely local source is dumping or leakage from ships. PCBs introduced into the ocean in this way would frequently be mixed with oil, would tend to mix with surface slicks (83), and hence would be peculiarly subject to uptake by plankton. Further study is necessary to confirm this suggested explanation or otherwise explain the anomalous low $\Sigma$DDT/PCB ratio in the Atlantic Ocean.

Table 9. PCB Levels and $\Sigma$DDT/PCB ratios in Marine Vertebrates.

| Area                  | Mean PCB level in fish (ppm) | $\Sigma$DDT/PCB ratio in fish, fish-eating birds and/or mammals | Reference |
|-----------------------|------------------------------|-----------------------------------------------------------------|-----------|
| Long Island Sound     | 1.2                          | 0.08–0.18                                                       | 78        |
| Bay of Fundy          | 0.5                          | 0.4                                                             | 29, 79, 80|
| Atlantic Ocean        | 0.1                          | 0.2–0.5                                                         | 57        |
| Puget Sound           | 0.16                         | 1.1                                                             | 56        |
| San Francisco Bay     | 0.1–1.2                      | 1–3                                                             | 32, 56    |
| Californian Coast     | 0.02–1                       | 5                                                               | 32, 56    |
| Gulf of California    | —                            | 10                                                              | 32        |
| Gulf of Panama        | —                            | 1–2                                                             | 32        |
| Pacific Ocean (Galapagos & Hawaii) | 0.03          | >10                                                             | 56        |
Assuming the standing crop of fish in the eastern North Pacific to be of the order of 10^8 tons, and that in the western North Atlantic to be of the order of 5 × 10^7 tons (derived from 59), we may estimate the corresponding loads of PCBs to be of the order of 10 and 15 tons respectively. Assuming the standing crop of plankton to be five times larger than that of the fish (1, 2), the corresponding load of PCBs might be of the order of 100 tons. In the case of ΣDDT, it has been estimated that the total load in marine plants may be several times larger than that in marine animals (2); this may be true for PCBs also, but we know of no measurements to support it. In any case it seems unlikely that more than a few × 10^7 tons of PCBs are stored in the marine biota of the area considered, a very small fraction of the total quantity introduced. This fraction may be significantly greater than the corresponding fraction for ΣDDT (1), because of the extraordinarily high levels of PCBs in the zooplankton.

We know of no measurements of PCB levels in sea water. However, as in the case of fresh water, it seems likely that most of the PCBs would be attached to sediments or to floating particles. Observations on sediments (6, 26) are in accord with this.

As for ΣDDT, the deep oceans represent an ultimate sink for PCB residues. There is no information from which to estimate the rate of transfer of either to the deep oceans.

Summary and Conclusions

According to the estimates made in this paper, the PCBs released into the North American environment in the past are now concentrated in three major compartments in the environment: (a) buried in landfill dumps (roughly 3 × 10^4 tons, without allowing for degradation); (b) attached to sediments in rivers and the Great Lakes (roughly 2 × 10^4 tons); (c) attached to sediments on the continental shelf (roughly 10^4 tons). A further substantial quantity (of the order of 2 × 10^4 tons) has been widely distributed over the land and sea by aerial fallout and by disposal from ships. All the numerical estimates are expected to be valid to order of magnitude only.

Transfer of PCBs within the environment is expected to take place by the following main routes: (a) volatilization, aerial transport on particulates, and fallout; (b) leaching from dumps; (c) sediment transport in rivers and in the shallow sea; (d) sedimentation in the ocean. Uptake and transport by the biota is probably a quantitatively unimportant route of transfer of PCBs but is of major biological significance. Virtually no evidence is available on the rate of transport by any of these routes, but it is expected that all are very slow. In particular, PCBs in sediments in rivers and lakes are likely to move downstream and augment those in the shallow sea for a long period into the future.

As a result of Monsanto's restrictions on distribution, several inputs into the environment are likely to have been sharply reduced. As existing products containing PCBs are scrapped, the remaining inputs into the North American environment are expected to decline gradually over a period of the order of ten years.

As in the case of DDT and its metabolites, the total quantities of PCBs accumulated by the biota are an extremely small fraction (less than one percent) of those in the environment. This raises the possibility that the dissemination of relatively small fractions of the PCB production into sensitive areas may be primarily responsible for locally high levels of contamination of the biota. Evaluation of the long-term effects of the accumulation of PCBs and of the change in use patterns and production will require the development of environmental transport models more sophisticated than those currently in use, together with the requisite data.

The following aspects of environmental transport of PCBs should have high priority for future research:

1. Properties and Rates. Data are needed on the solubilities and vapor pressures of individual isomers; on the partition coefficients between water and sediments, water and fats; on the rate of photochemical decomposition and aerobic and anaerobic biodegradation; diffusion coefficients in sediment.

2. Routes. The magnitude of leaching from dumps and aerial transport of PCBs must be established by direct measurements of concentrations in ground waters, airborne
particulates, and rainfall. Data are needed on the magnitude of industrial losses in the past.

3. **Transport Models.** Data are needed on the transport of PCBs in contaminated sediments in rivers and shallow seas, including the dynamics of uptake from shallow sediment and losses to the ocean floor.

4. **Levels.** Additional data, including quantitative evaluation of the fractionation of isomers, are needed for fresh waters, soils, terrestrial plants and animals.

5. **Bioaccumulation model.** Inasmuch as the period since the first commercial application of PCBs is less than the lifetime of a number of species, including man, and the use and discharge pattern has been variable, there is a need to project the long-range impact of the PCBs on the different elements of the ecosystem.

Emphasis should be on coordinated approaches to sampling and modelling rather than a statistical accumulation of data.

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13. Approximately 190×10⁶ tons of municipal waste and an equivalent amount of industrial solid waste are disposed of annually in the U.S.A., mostly in dumps. Some 4.7×10⁶ tons of industrial solid waste are dumped at sea (27). Of the remainder, 18×10⁶ tons are burned in municipal incinerators, 14×10⁶ tons in commercial incinerators, and 14×10⁶ tons in open-burning dumps (14). Emission of PCBs from these sources may be gauged indirectly from measurements of the emission of combustible matter and polynuclear aromatics. From measurements reported by A. D.
Little (15) and Hein and Engdahl (16) it is estimated that 0.25 percent of solid waste is emitted by municipal incinerators and up to 20 percent by open burning. If it is assumed that the fraction of PCBs in the organic matter evolved from these sources is equal to the fraction of PCBs in the solid waste, the rate of emission of PCBs will be of the order of two tons per year from municipal incinerators and 200 tons/year from open burning dumps. Commercial incinerators and burning of automobile components each contribute a total quantity of polynuclear aromatic compounds (a rough gauge of incomplete burning) similar to that emitted by open burning (14). The estimate of 200 tons/year of PCBs from the burning of commercial waste and auto components is based on the relative magnitude of combustible pollutants from these different sources, weighted by the estimate of the fraction of total PCB production disposed of in open dumps. These estimates are highly speculative but are considered sufficient to discount the burning of wastes as the major source of PCB emissions.

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Dumping of polluted dredge spoils was estimated in the text to contribute no more than 20 tons/year. Accidental leaks from coastal industries might be of the order of 10^6 tons/year, based on measurements in sediments following the Escambia Bay incident (6, 26). Deliberate dumping at sea is estimated to account for 2-3 percent of industrial solid wastes in the U.S.A. (13): if it accounts for the same fraction of industrial disposal of scrap PCBs, this would amount to less than 100 tons/year, after allowing for degradation. Losses from ships may amount to a few X 10^4 tons/year, based on the likely contribution (5-10%) of shipping to total uses of hydraulic and lubricating fluids. Losses from aircraft are assumed to be small. Most of these figures are probably somewhat high, so we estimate the total input to the oceans from these sources as less than 10^6 tons/year. Only some 10-20 percent of this total is likely to have entered the Pacific Ocean.

50. Only scattered data on concentrations of PCBs in water have been published. Water entering a Swedish water works contained 0.5 ppt (parts per trillion) (86). Industrial rivers in Maine, Massachusetts, Wisconsin, and Japan contained levels ranging from 0.2 to 2.8 ppb (42, 51-53). Measured levels in Green Bay and Milwaukee Harbor, Wisconsin, were in the range 20-70 ppt (42, 52). Other estimates can be derived indirectly from ZDDT/PCB ratios in aquatic fauna (see text). In Lake Michigan, where ::DDT levels are of the order of 1-3 ppt (54), ZDDT/PCB ratios in fish are of the order of 0.4-1 (55). In San Francisco Bay, the ZDDT/PCB ratio in the biota is 1-2 (32, 56), and the level of ZDDT in the San Joaquin River, the main input to the bay, is about 100 ppt (32). For the rough calculations of storage and outflow made in this paper, we use these data as a basis to adopt 0.5 ppt as a plausible background level for unpolluted fresh waters, 5 ppt for the Great Lakes, 50 ppt for slightly polluted rivers and polluted bays, and 500 ppt for highly polluted rivers.

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