Oculodermatological findings in workers with occupational exposure to polychlorinated biphenyls (PCBs)

A FISCHBEIN,1 JOAN N RIZZO,1 S J SOLOMON,2 AND MARY S WOLFF1

From the Environmental Sciences Laboratory,1 Department of Community Medicine and Department of Neoplastic Diseases,2 Mount Sinai School of Medicine of the City University of New York, New York, New York 10029, USA

ABSTRACT

Oculodermatological findings, such as hypersecretion of the Meibomian glands, swelling of the upper eyelids and hyperpigmentation of the conjunctivae are considered typical of “PCB poisoning.” They were common clinical manifestations of Yusho and Yu-cheng, two epidemics in Japan and Taiwan caused by the ingestion of rice cooking oil contaminated with polychlorinated biphenyls (PCBs) and polychlorinated dibenzofurans. To investigate the prevalence of such abnormalities in a population with long term occupational exposure to PCBs, a group of 326 workers employed in two capacitor manufacturing facilities were investigated in 1976, and 195 of these again in 1979. The median blood values of lower homologues of PCBs were 63 ppb (in plasma) in 1976 and 49 ppb (in serum) in 1979, and of the higher homologues 18 ppb and 17.5 ppb respectively. The prevalence of oculodermatological findings potentially related to the effects of PCBs were 9.4% and 13.3% at the two examinations. There was no significant association between such abnormalities and blood plasma/serum concentrations of PCBs. The observations in this work population exposed to PCBs differ from the Yusho and Yu-cheng experiences in that fewer clinical abnormalities were found. Suggestions are made that it may be inappropriate to extrapolate findings from the well known PCB poisoning episodes to exposures in occupational settings and that attention should be paid to the importance of polychlorinated dibenzofurans as an aetiological factor in human PCB poisoning.

Polychlorinated biphenyls (PCBs) are a group of complex chlorinated aromatic hydrocarbons that have been used industrially since the late 1920s.1 Because of their chemical stability, they were valuable for many industrial uses, such as additives to paints, surface coatings, and varnishes, and as a constituent in carbonless copy paper. Their principal use, however, has been as dielectric fluids in capacitors and transformers.

PCBs were manufactured in the United States under the trade name Aroclor (R), and the numbers ascribed to the trade name identified the degree of chlorination. For example, Aroclor 1254 and 1260 refer to 54% and 60% chlorine content respectively.

The manufacture of PCB containing carbonless copy paper was discontinued in the United States in 1971. In 1972 the Food and Drug Administration (FDA) established regulatory guidelines for the control of human exposure to PCBs after the compounds were shown to have contaminated food from packaging materials.2 The manufacturing of all PCBs was discontinued in the United States in 1977 and their use further restricted.3

Despite the widespread industrial use of PCBs since the 1930s, it was not until 1966 that they were first recognised as a major environmental pollutant,4,5 and subsequently identified in the air, soil, water. Bioaccumulation has also been described in vegetation, fish, and milk products.6

PCBs are characterised by a low degree of biodegradability and tend to persist in the environment.

Levels of PCBs have been well documented in people of industrialised nations.4–7 Since PCBs are lipophilic compounds, they tend to accumulate in tissues or organs according to adipose tissue content.
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Information available from numerous animal studies indicates a wide range of PCB related toxic effects on the liver, lipid metabolism, and both the immune and reproductive systems.\(^8\)\(^9\) PCBs produce hepatomas and hepatocellular carcinomas in experimental animals.\(^10\)\(^11\) Differences between species in their response to PCBs have also been substantiated.\(^8\)\(^12\)

Dermatological abnormalities have been frequently reported in animals exposed to PCBs. The pilosebaceous unit is the most commonly affected target organ, and acneiform lesions (chloracne) are considered typical sequelae from exposure to PCBs and related compounds. Enlargement of the Meibomian glands and swelling of the eyelids are also frequent manifestations associated with experimental PCB poisoning.

Human health effects associated with exposure to PCBs have not been well defined. In the 1930s and 1940s several investigators of occupationally exposed individuals reported multiple skin lesions characterised by chloracne, and occasionally associated with systemic effects.\(^13\)\(^14\) It should be emphasised, however, that in many of the early reports exposures may well have been not only to PCBs but to a combination of PCBs and other halogenated hydrocarbons, particularly chlorinated naphthalenes.\(^15\)

Yusho and Yu-cheng

A clinical syndrome of “PCB poisoning” was first described in detail in connection with two large scale outbreaks of PCB poisoning that were due to the ingestion of contaminated rice cooking oil. The first epidemic (“Yusho: Japanese “oil disease”) occurred in 1968 in Japan, and the second (Yu-cheng: Chinese “oil disease”) in 1979 in Taiwan.\(^16\)\(^17\) Chloracne, which is characterised by comedos, pustules, and straw coloured cysts was frequently observed. Oculodermatological signs and symptoms were important diagnostic criteria for these diseases and included hypersecretion of the Meibomian glands, swelling of the eyelids, and hyperpigmentation of the conjunctivae. These abnormalities were found in 60–85% of the affected individuals.\(^18\)\(^–\)\(^20\)

It is important to note that polychlorinated dibenzofurans (PCDFs) and polychlorinated quaterphenyls (PCQs) were identified both in cooking oils and in patients with the Japanese and Taiwanese PCB poisoning and have been indicated as possible aetiological factors in the PCB poisoning.\(^21\)\(^–\)\(^24\)

Materials and methods

To evaluate the health effects of long term occupational exposure to PCBs, a cross sectional study was conducted in March 1976 of 326 workers employed in two capacitor manufacturing facilities.\(^25\) The use of PCB was discontinued in the manufacturing process in July 1977. A follow up examination was conducted 45 months later, in December 1979. Workers in PCB related occupations were invited to take part in the examinations on a voluntary basis. The most important criterion for inclusion in the study was long term employment of 20 years or more, although some individuals with shorter durations of employment who came to the examination were also included. The exposure of all individuals to PCBs occurred during employment in the production of capacitors, which were filled with dielectric fluid containing PCBs.

There was a potential for exposure through skin absorption, inhalation, and, to a lesser degree, ingestion of fluid containing PCBs. The workers handled heated and partially volatilised dielectric fluids. A wide range of laboratory and clinical tests was performed on the study population, and various questionnaires were administered to determine occupational and medical histories, and to elicit symptoms potentially related to the toxic effects of PCBs. Special designed forms were used to record oculodermatological findings. PCBs were measured in plasma and serum by gas chromatography (electron capture detection) according to methods previously described.\(^26\) PCB residues were calculated as Aroclor 1248 (for peaks with retention times less than p,p'-DDE) and designated as lower homologues of PCBs (LPCB). Peaks with retention times greater than p,p'-DDE were calculated as Aroclor 1254, and regarded as higher homologues of PCBs (HPCB). A clinical biochemistry screen was also performed and included liver function tests (SGOT, SGPT, Alk phos, \(\gamma\)-GTP, LDH, and bilirubin) and serum lipids (triglycerides and cholesterol).

PCB levels were available in 289 of the 326 (88.7%) subjects in 1976. To avoid potentially confounding factors in the analysis of the data, 43 subjects with a history of excessive alcohol consumption or of taking medications with potential effects on the liver and serum lipids were excluded from the analysis. The study sample thus consisted of 246 individuals in 1976, 106 women aged 23 to 60 (median age 49), and 140 men aged 21 to 67 (median age 40). Although there was a significant difference in age between the sexes (Wilcoxon two sample tests, \(s = 1569.5, p < 0.001\)), this pattern was not observed for any of the other independent variables of interest. The individuals were subdivided into two groups based on the presence or absence of oculodermatological abnormalities considered to be related to PCBs. Twenty three indi-
The median LPCB values of the examined workers were 63 ppb in 1976 and 49 ppb in 1979, whereas the HPCB levels were 18 ppb and 17.5 ppb. Although the mean HPCB levels of the exposed workers did not decrease from the time of the initial study \(t = -0.46, p > 0.05\), there was a significant reduction in LPCB \(t = -2.66, p < 0.008\).

The table shows the frequency of oculodermatological abnormalities in the examined population. At the first examination, 29 workers (9.4%) had at least one oculodermatological sign and five individuals had more than one abnormality. Seventeen (5.5%) had injected conjunctiva, seven (2.3%) had eye discharge, five (1.6%) had swelling of the upper eyelid, and four (1.3%) had enlarged Meibomian glands.

In the follow up study performed in 1979, 195 of the original 326 workers returned for re-examination. Fourteen were excluded from the analysis due to “other” abnormalities as in the table.

Of the remaining 181, 24 (13.3%) exhibited abnormal oculodermatological findings. Fifteen (8.3%) had injected conjunctiva, three (1.7%) showed discharge, four (2.2%) eyelid oedema, and two (1.1%) had enlarged Meibomian glands. None had abnormal pigmentation of the conjunctiva, and of those with injected conjunctiva, only one had this abnormality in 1976. Eleven individuals categorised as “injected” at the second examination had been classified as normal in 1976, and one had eyelid oedema in 1976. Two workers who were excluded from consideration in 1976 because of “other” or pupillary abnormality had injected conjunctiva in 1979.

Significant differences were not found for PCB levels, liver function tests, serum lipids, or systolic and diastolic blood pressures between those with and without oculodermatological signs. All mean levels were within the normal laboratory range. The low prevalence of abnormal clinical biochemistry tests have been summarised previously.\(^{25}\)

**Discussion**

In the major outbreaks of PCB poisoning from ingestion of contaminated oil in Japan and Taiwan, most (60%–85%) exposed individuals developed oculodermatological signs,\(^{19,20}\) characterised by swelling of the upper eyelid, enlargement of the Meibomian glands with eye discharge, hyperpigmentation, and injection of the conjunctiva. At the time these findings were regarded as typical PCB related abnormalities.

In the present investigation we examined a group of workers with long term occupational exposure to PCBs with 30–60% chlorination, and significant...
absorption of PCBs was reflected in the high serum PCB levels. By contrast, the PCB levels in patients with Yusho and Yu-cheng were generally lower than those found in the present study, although methodological variations may account for some of the differences.27

Our observations show a discrepancy in the clinical effects reported for the Yusho and Yu-cheng patients and our occupationally exposed study group—that is, despite higher PCB levels among the workers, the prevalence of oculodermatological signs was much lower than in the typical Yusho and Yu-cheng patients. This is consistent with our previously reported findings concerning dermatological abnormalities, including chloracne.28

In evaluating these differences it should be recognised that several investigators have identified polychlorinated dibenzofurans and polychlorinated quaterphenyls in the contaminated cooking oils, as well as in the patients with Yusho and Yu-cheng.21242930 It has also been shown that PCDFs are relatively more toxic than PCBs,3132 Moreover, it has been further suggested that these contaminants may have played a crucial part in the appearance of the toxic clinical signs of PCB poisoning.30 Different toxic responses observed in animal studies have also been attributed to contamination of some PCB mixtures with PCDFs and perhaps chlorinated naphthalenes.31

At present, we do not have conclusive evidence concerning PCDFs in the dielectric fluid to which our study subjects were exposed. Elsewhere, analyses of PCBs of varying degrees of chlorination have shown levels of PCDFs less than 1/1000 as compared with those found in Yusho and Yu-cheng.33

Another possible explanation for the apparent discrepancy in clinical findings may relate to the route of exposure. Our subjects were exposed in PCBs in an occupational setting, in which long term exposure to volatilised PCBs occurred as well as exposure by skin contact; by contrast, the intoxication in Japan and Taiwan resulted from the ingestion of contaminated oil over a relatively short time. Our study suggests that it may be inappropriate (with the present knowledge of PCB related compounds) to extrapolate from Yusho and Yu-cheng any clinical effects that might be related to PCB exposure in an occupational environment. Based on our investigation, severe clinical effects are relatively rare and differ considerably in degree from the Japanese and Taiwanese experiences. The current lack of corroborative data from analytical chemistry to support the presence of PCDFs and PCQs in the dielectric fluid and in the exposed workers prevents us at this time from reaching final conclusions about cause-effect relations.

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