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DIOXIN CANCER RISK — EXAMPLE OF HORMESIS?

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A recent case-control study implied an inverse correlation between the measured body burden of dioxins (polychlorinated dibenzo-p-dioxins and polychlorinated dibenzofurans, PCDD/F) and the risk of soft tissue sarcoma in normal population exposed to dioxins mainly via food. The surprising result could not be explained by biases or confounding. There is no a priori confounding by occupational chemicals in a random sample from general population, but exposures to other lipid soluble chemicals with similar sources might be expected to associate with that of dioxins. One such group is polychlorinated biphenyls (PCB). Therefore three most relevant dioxin-like PCB compounds PCB 77, PCB 126, and PCB 169 were now analyzed from the same patients. Cases were 110 soft-tissue sarcoma patients undergoing surgery for their disease, and referents were 227 patients operated for appendicitis. Dioxin and PCB concentrations were analyzed from subcutaneous fat samples by high-resolution gas chromatography–mass spectrometry and TCDD equivalent concentrations (WHO-TEq) were calculated by using toxicity equivalency factors of WHO. The highest risk of sarcoma was found in the septile with the lowest body burden of sum WHO-TEq, and the differences of septiles 2 and 6 from septile 1 were statistically significant. If soft sarcoma risk is true at high occupational levels of dioxins, the provocative result suggests that a possibility of a J-shaped dose-response curve should be taken into consideration and studied further. This is also supported by the similar J-shaped dose responses in animal studies.
INTRODUCTION

There has been a continuous search for a “physiological” function of the AH receptor (dioxin receptor) (Poellinger, 2000). In this discussion the fact that induction of metabolism of foreign substances via AH receptor is a physiological function, has been less often emphasized. A physiological function means implicitly that at some level of activation, the action of AH receptor is useful as a response to environmental stimuli. In fact it would be strange that a highly conserved gene (Hahn, 2002) would have been maintained, unless the function is useful or even a necessity for life.

On the other hand it is clear that dioxin-like compounds cause a plethora of adverse effects at high dose levels (for a review, see Pohjanvirta and Tuomisto, 1994). Most of these are AH receptor-dependent and they include various developmental effects (Abbot and Birnbaum, 1989, 1990, Alaluusua et al., 1993, Kattainen et al., 2001, Mably et al., 1992), thymic atrophy and immunological effects (Kerkvliet et al., 1990), various hyperplastic/metaplastic effects (Poland and Knutson 1982) and cancer (Kociba et al., 1978). In humans several effects have been seen after high accidental or occupational doses, e.g. chloracne and pigmentation (Sweeney and Mocarelli 2000, Geusau et al., 2001), several developmental effects such as growth retardation, hyperpigmentation, neurobehavioural changes, and alterations of sexual development after intrauterine exposure (Rogan et al., 1988, Masuda et al., 1996, Feeley and Brouwer 2000) and tooth deformities after accidental exposure during childhood (Alaluusua et al., 2004). A modest increase in total cancer is suggested by occupational cohort studies (Kogevinas 2000). Increases of gastrointestinal and hematopoietic malignancies have been seen after a TCDD release accident in Seveso, Italy (Bertazzi et al., 1997, 2001). Soft-tissue sarcoma was increased toward the high end of occupational exposures in a large industrial cohort (Fingerhut et al., 1991, Steenland et al., 1999).

Previous cancer studies suffer from poor exposure assessment. In most studies exposure information is based on indirect methods such as work histories, sometimes supported by chemical analyses in part of the studied population and modeling, or in many cases on questionnaire information. Therefore we recently undertook a major attempt of studying soft tissue sarcoma risk in general population in correlation to dioxin concentrations (Tuomisto et al., 2004). Sarcoma patients coming to surgery for their tumor were analyzed for their dioxin concentrations, and in a case-control setting appendicitis patients were studied as references. A surgical patient control group was needed, because dioxin analysis was
from a subcutaneous fat sample. Among the general population in Finland the variation in dioxin concentrations is very large, because the dominant source of dioxins is fish, and fish consumption varies widely in the population. In this study no positive correlation was seen between the dioxin concentrations and soft tissue sarcoma risk, on the contrary, sarcoma risk was highest among those having the lowest dioxin level.

Now three PCB congeners contributing most to the dioxin equivalents have been analyzed in addition to the 17 PCDD/F congeners, and we are able to present data on these congeners added to the total burden of dioxins.

**MATERIALS AND METHODS**

A detailed description of the methods was given in the previous paper (Tuomisto et al., 2004). Therefore only the most pertinent details are shortly described here.

**Study population**

Most sarcoma patients in southern Finland are treated by the multidisciplinary sarcoma group of Helsinki University Hospital, and the rest in the University Hospitals of Kuopio, Turku, or Tampere. All patients over 15 years of age referred to these hospitals for operative treatment of STS between June 1997 and December 1999 were eligible as cases. Patients over 15 years of age, operated due to an appendicitis diagnosis in any study hospital were eligible as controls. They were collected from the same catchment area as the STS patients. Southern Finland was divided into 15 areas, and one hospital performing appendectomy operations was recruited to the study from each area (in Helsinki, two hospitals).

Informed consent was obtained from all patients in writing before the operation. The study was duly approved by the Ethics Committees. The total number of patients recruited during the fieldwork was 972, and after exclusion of some patients for technical reasons (see Tuomisto et al., 2004), 954 patients (148 cases and 806 controls) were available for matching.

The cases and controls were individually matched for area and age at the end of the fieldwork. Area was defined as the area of residence of the patient using the 15 areas above. The age was determined at the day of operation. Maximum allowed age difference between cases and controls was ±3 years, if case was <38.0 years old, and ±6 years, if case was >38.0 years old. The control closest by age was matched to the case. Cases with fewer controls had a priority over cases with more controls. The number of controls per case was limited to three. For 110 cases 227 matching controls could be found in the pool. Thirty-nine cases had one control, 25 cases had two, and 46 cases had three controls.

**Exposure assessment**

A subcutaneous fat sample of the matched 337 patients, obtained during an appendectomy or sarcoma operation, was analyzed for the 17 toxic polychlorinated dibenzo-p-dioxins and dibenzofurans (PCDD/Fs) and three dioxin-like non-ortho polychlorinated biphenyls: PCB 77 (3,3′,4,4′-tetrachlorobiphenyl), PCB 126 (3,3′,4,4′,5-pentachlorobiphenyl), and PCB 169 (3,3′,4,4′,5,5′-hexachlorobiphenyl). Measurements were done by gas chromatography – mass
spectrometry (Vartiainen et al., 1997) at the Laboratory of Chemistry, which is an accredited testing laboratory (T077) for the analysis of dioxins in human samples (current standard: EN ISO/IEC 17025) and has successfully participated in WHO/Euro intercalibrations. The concentrations were summed up after the value of each congener was multiplied by its relative toxic potency (toxic equivalency factor, TEF). The TEF values according to WHO (Van den Berg et al., 1998) were used, resulting in toxic equivalent concentrations (WHO-TEq) comprising 17 PCDD/Fs and 3 PCBs. All analytical work was performed blind so that the chemistry laboratory did not know the diagnosis of the patient. Strict quality assurance measures were undertaken.

Patients were also asked detailed questionnaire information about socio-economic and lifestyle factors and chemical exposures. Of the matched subjects, 84 cases (76 %) and 185 controls (81 %) have also questionnaire information.

**Statistical analyses**

Conditional logistic regression analysis was performed with SAS PHREG procedure. Odds ratios were estimated for each septile of WHO-TEq. All analyses were adjusted for sex. Several variables collected with the questionnaire were used as confounders in the analysis one by one. Non-binary variables were analyzed as quartiles.

Exposure to the following chemicals was asked as a binary variable: solvents, solvent-based paints, formaldehyde, insecticides, fungicides/herbicides, wood preservatives, strong detergents, heavy metals, and other chemicals.
RESULTS

Contribution of PCB-TEqs to total WHO-TEq was similar in all age groups, about 20% on average (Fig. 1). On the other hand, individual variation was large, and PCB contribution was from 1% to 57%. The range of total TEq values was from 4.6 to 197.8 ng/kg (in fat). There was high correlation between PCB-TEq and PCDD/F-TEq (Pearson correlation coefficient 0.84) and between PCB-TEq and total WHO-TEq (r=0.91).

The STS risk was higher in the lowest septile than in other septiles (Fig. 2), and the difference was significant in the second and the sixth septile. The odds ratios (compared with the lowest septile) varied from 0.15 (95% CI 0.04 – 0.6) to 0.39 (95% CI 0.12 – 1.27). When the analysis was performed with WHO-TEq quintiles instead of septiles, the odds ratios were 1, 0.44 (0.18-1.09), 0.73 (0.29-1.84), 0.40 (0.14-1.13), and 0.49 (0.16-1.49). The analysis was also calculated using total WHO-TEq concentration as a continuous linear variable showing a decreasing trend (OR 0.86 for an interquartile TEq increase of 33.32 ng/kg [WHO-TEq in fat], 95% CI 0.60 – 1.24). When the three PCB-compounds were analyzed separately, the continuous linear variables were: PCB 77, 1.08 (0.93-1.26); PCB 126, 0.92 (0.7-1.22); PCB 169, 0.93 (0.68-1.27), and the sum TEq of all three PCBs 0.92 (0.69-1.23).

The decrease in odds ratios was not abolished by any of the studied confounders (sex, age, education, body mass index, fish consumption, smoking, alcohol consumption, town size, and chemical exposure) (Fig. 3). On the other hand, including confounders in the analysis tended to cause the odds ratios to decrease in more linear fashion than in the basic result, especially to straighten the kink of the second septile. At the same time, however, the statistical significance of decreased odds ratio disappeared in septile 2 (which was the most unstable septile) in most analyses and in septile 6 in some analyses. A continuous linear analysis including fish as a confounder showed a decrease (OR 0.83 for an interquartile TEq increase, 95% CI 0.53 – 1.30).
DISCUSSION

The present results indicate that the previous analysis comprising only 17 PCDD/F congeners does not essentially change, when the non-ortho PCB congeners, which contribute most to the PCB-TEq are also included in the sum analysis. The result adds to confidence that previously reported result between dioxin-TEQ and decreased cancer risk of cancer is not confounded by the levels of PCB. In the previous study each dioxin congener was also analyzed separately, and a general trend was that the congeners contributing most to WHO-TEq showed a result similar to the main analysis of sum TEq. In the present study PCB 126 (which contributes most to PCB-Teq), and PCB 169 indicated even alone the same trend of decreasing risk. It should be noted that the concentrations of all these compounds correlate with each other, since their sources are in part similar.

The contemporary body burden is considered a good measure for lifelong exposure to dioxins and PCB compounds. The half-life of TCDD is 7 to 8 years (Poiger and Schlatter, 1986, Pirkle et al., 1989, Flesch-Janys et al., 1996) and this means at a constant intake a cumulation of about 40 years before reaching steady state. Because exposure is mainly from food (Liem et al., 2000), fairly stable exposure can be assumed on an average, and considering the long half-life, the present concentration means integration of exposures during at least 10-20 years, in most cases probably over the whole lifetime. Although different from congener to congener, elimination half-lives of all dioxins and PCBs are very long, and such integration can be assumed to hold for WHO-TEqs as well.

Present exposure in Finland is mainly from Baltic fish, up to over 80% (Kiviranta et al. 2001), although during previous decades also meat and dairy products contributed to some extent (Hallikainen and Vartiainen, 1997). Because fish consumption varies from person to person, this could be assumed
to be one of the major differences causing variation in dioxin and PCB levels in different people. An obvious hypothesis then could be that fish consumption might decrease cancer risk, and cause the decreasing trend observed. However, if fish consumption was added to the analysis as a confounder, the main result did not change. Interestingly the analysis improved in the sense that the decrease associated with increasing dioxin TEqs became more regular and linear. These issues require further scrutinizing.

In animal studies, TCDD is a clear carcinogen in several species at high doses (Kociba et al., 1978, Pitot et al., 1987, NTP 1982, Goodman and Sauer, 1992). However, a tendency of less tumors or altered hepatic foci (indicating a carcinogenic process) were seen in animals given low doses of TCDD (ca. 1 ng / kg body weight / day), than were present in the controls in several studies (Kociba et al., 1978).
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1978, Pitot et al., 1987, Viluksela et al., 2000). This dose would lead to a TCDD concentration in rats of approximately 100 – 500 ng/kg (TCDD in fat) (Kociba et al., 1978, Viluksela et al., 2000). This matches with the higher end of the human concentrations in the present study (Fig. 4).

Andersen and Conolly (1998) have suggested a mechanistic model to explain the U-shaped dose-response in liver tumors described by Pitot et al. (1987). The model assumes that some (but not all) of the mutated cells are sensitive to the mitoinhibitory effect of TCDD, resulting in a decrease in the number of mutated cells at low doses. Further studies to scrutinize such mechanisms are urgently needed.

It may also be worthwhile to note that the non-induced activity of some CYP enzymes, notably CYP1A1, is very low (Whitlock 1999). A slight induction could be expected to facilitate the metabolism of many xenobiotics including carcinogenic polycyclic hydrocarbons. Depending on the relative activity of the next stage of metabolism, and possible interactions of dioxins with phase II conjugation reactions (cf. Nguyen et al., 2003), clearance of such compounds could then be increased and their chances of acting as cancer initiators would be decreased.

In conclusion, the previously reported increased risk of STS at the lowest levels of WHO-TEq was maintained when three most important dioxin-like PCBs were included in the analysis. The provocative result within the present population levels of dioxin exposure challenges the present carcinogenicity estimates based on linear extrapolation from high to low doses. The existence of a true J-shape dose-response curve in carcinogenesis by dioxin-like compounds requires more scrutiny, but there are biologically plausible reasons to consider this possibility.

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