Aryl hydrocarbon receptor-interacting protein and pituitary adenomas: a population-based study on subjects exposed to dioxin after the Seveso, Italy, accident

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Abstract

Objective: The pathogenesis of sporadic pituitary tumors is unknown. Loss-of-function mutations of aryl hydrocarbon receptor-interacting protein (AIP) have been identified in patients with familial pituitary tumors. AIP is a chaperone protein with multifunction properties, including modulation of the transcriptional activity of the aryl hydrocarbon receptor, which mediates toxicological and carcinogenic dioxin effects.

Design: We investigated the incidence of pituitary tumors in the Seveso population exposed to 2,3,7,8-tetrachlorodibenzo-p-dioxin following an industrial accident in 1976.

Methods: Through the hospital discharge registration system of Lombardy Region, we identified incident cases of pituitary adenomas between 1976 and 1996 in the Seveso population, subdivided in zone A (n = 804), B (n = 5,941), and R (n = 38,624) according to high, intermediate, and low exposure to dioxin respectively, and in the surrounding non-contaminated area, as reference (n = 232,745).

Results: We identified 42 pituitary adenomas in the reference area, 1 prolactinoma in zone A (rate ratio (RR) 6.2; 95% CI 0.9–45.5, P = 0.07), 2 non-functioning pituitary tumors (NFP As) in zone B (RR 1.9; 95% CI 0.5–7.7, P = 0.39), and 3 prolactinomas and 2 NFP As in zone R (RR 0.7; 95% CI 0.3–1.8, P = 0.48).

Conclusions: The study is unique with regard to the availability of epidemiological and clinical data in an area of relatively pure dioxin exposure. The study indicates no statistically significant increase of incident pituitary tumors in this area, although the tendency toward a higher risk (three cases in zones A and B) of pituitary tumors in subjects exposed to high–intermediate dioxin concentrations in comparison with nonexposed population suggests the need for extended follow-up.

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Introduction

Pituitary tumors are benign lesions that account for substantial morbidity due to hormone over-production and/or intracranial mass effects. Despite the large number of mutational studies carried out in the past decades, the pathogenesis of these tumors remains largely unknown (1). Vierimaa et al. recently reported loss-of-function mutations of the gene encoding the aryl hydrocarbon receptor-interacting protein (AIP) in patients with familial pituitary tumors, suggesting that AIP may operate as a tumor suppressor gene determining pituitary tumor susceptibility (2). This hypothesis has been recently confirmed by in vitro data showing a reduced ability of mutant AIP to block cell proliferation (3).

Clinical characterization of patients with familial isolated pituitary adenomas associated with AIP mutations reported a relative predominance of somatotropinomas, with a low incidence of prolactinomas and nonfunctioning pituitary adenomas, an early age at diagnosis, and a particularly aggressive tumor behavior (3, 4). In contrast to the frequency of AIP mutations in familial isolated pituitary tumors, evaluated in ~15%, screening studies failed to detect AIP mutations in most sporadic pituitary adenomas (5).

AIP is a chaperone protein that interacts with several partners, including the aryl hydrocarbon receptor (AHR), an orphan nuclear receptor that is strongly activated by 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD or simply dioxin) and mediates dioxin toxicological properties (6). In particular, AIP modulates the function of AHR by both protecting AHR from ubiquitination or retaining AHR in the cytoplasm and preventing transcriptional activity (7, 8). Although these data
suggest that AIP inactivation might result in AHR signaling amplification, whether exposure to dioxin increases the risk of pituitary adenomas has never been determined. An industrial accident in Seveso, Italy, in 1976 exposed thousands of people to substantial quantities of dioxin. Mortality and morbidity findings during the 20–25-year period following the accident showed increased risk from cancers, particularly lymphoepithelioid neoplasms (9–11).

The aim of the study was to evaluate the impact of exposure to dioxin on the incidence of pituitary tumors in the Seveso population.

Materials and methods

Subjects

Immediately after the accident, three areas of high, intermediate, and low contamination (zone A, B, and R respectively) were delimited on the basis of soil dioxin measurements (Fig. 1A). A surrounding non-contaminated territory was adopted as reference area. Subjects were assigned to one of the contaminated zones or the reference area on the basis of their residence at the date of the accident. Improvements in analytical techniques allowed the measurement of TCDD levels in very small blood samples collected at the time of the accident among subjects chosen as the most highly exposed in zone A, B, and R (12, 13). Serum TCDD levels, measured on blood samples taken at the time of the accident (13), showed that contamination zones correctly represented three levels of decreasing exposure to dioxin (Table 1). Median levels in a random sample of subjects from the reference nonexposed area were 5.5 ppt, representing the typical value in the general population (14). A cross-sectional study conducted ~20 years after the accident on subjects from zones A and B showed persisting elevated levels of serum TCDD, as expected based on the long (>7 years) TCDD half-life (14, 15). All subjects living in the Seveso area were followed up, blind of their exposure status, to evaluate long-term health effects through mortality and cancer incidence studies (9, 11). Incident cases of pituitary adenoma between 1976 and 1996 were identified through the hospital discharge registration system of the Lombardy Region (where the study area is located). For each case, all relevant medical records were reviewed and diagnoses confirmed.

Statistical analysis

Rate ratios (RR) and 95% confidence intervals (CI) were estimated with Poisson regression techniques controlling for age, gender, and calendar period, assuming a 10-year latency for dioxin effects. The surrounding non-contaminated area was used as reference. The results reported here refer to subjects, aged 0–74 years, resident in the accident area at the time of the accident (July 10, 1976).

Results

Characteristics of pituitary adenoma cases diagnosed in the Seveso population are summarized in Table 1. Assuming a 10-year latency for dioxin effects after the 1976 accident, eight tumors were identified between 1986 and 1996. In particular, one case of microprolactinoma was diagnosed in a 33-year-old woman living in the most contaminated zone (zone A, 804 subjects) at the time of the accident. Interestingly, soon after the accident she developed chloracne, the typical skin disorder due to exposure to dioxin-like compounds. In 1997, this subject was involved in a study designed to investigate health status and serum dioxin levels in chloracne cases (15), which showed that her serum TCDD was still very elevated (231 ppt). Two nonfunctioning pituitary adenoma (NFPA) were diagnosed among subjects present in zone B (n = 5,941) and five (three prolactinomas and two NFPA) in zone R (n = 38,624). Age-, sex-, and period-adjusted relative risks (RRs) indicated no statistically significant increase of the accident. Improvements in analytical techniques allowed the measurement of TCDD levels in very small blood samples collected at the time of the accident among subjects chosen as the most highly exposed in zone A, B, and R (12, 13). Serum TCDD levels, measured on blood samples taken at the time of the accident (13), showed that contamination zones correctly represented three levels of decreasing exposure to dioxin (Table 1). Median levels in a random sample of subjects from the reference nonexposed area were 5.5 ppt, representing the typical value in the general population (14). A cross-sectional study conducted ~20 years after the accident on subjects from zones A and B showed persisting elevated levels of serum TCDD, as expected based on the long (>7 years) TCDD half-life (14, 15). All subjects living in the Seveso area were followed up, blind of their exposure status, to evaluate long-term health effects through mortality and cancer incidence studies (9, 11). Incident cases of pituitary adenoma between 1976 and 1996 were identified through the hospital discharge registration system of the Lombardy Region (where the study area is located). For each case, all relevant medical records were reviewed and diagnoses confirmed.

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in pituitary adenoma incidence in subjects exposed to high and intermediate concentrations of dioxin in comparison with the nonexposed reference population (42 cases per 232,745 population). However, a tendency toward a higher risk of pituitary tumors in zones A and B with RRs of 6.2 (95% CI 0.9–45.5, \( P = 0.07 \)) and 1.9 (95% CI 0.5–7.7, \( P = 0.39 \)) respectively, was observed (Fig. 1B). The incidence of pituitary tumors in the least contaminated area (zone R) did not differ from the reference population (RR = 0.7; 95% CI 0.3–1.8, \( P = 0.48 \)).

### Discussion

This study first reports the incidence of pituitary tumors in an area of relatively pure exposure to TCDD. TCDD is the most toxic member of the large family of polychlorodibenzodioxins, which was recognized as a strong toxicant and carcinogen in experimental animals. Studies of military and occupational cohorts have led to the classification of TCDD as a human carcinogen (16). The Seveso accident in 1976 caused severe TCDD exposure to a population comprising of both genders and all ages, with little or no interference by other contaminants. In the early post-accident period, several health outcomes, such as spontaneous abortion, cytogenetic abnormalities, congenital malformations, liver function, immunological and neurological impairment, and chloracne, were evaluated (11). Cohort studies that investigated the long-term impact of the accident recorded elevated risk of lymphatic and hematopoietic neoplasm, increased cardiovascular mortality in the first years after the event, and suggestive increases in diabetes and chronic lung diseases (9–11).

Several recent studies reported that patients with familial pituitary adenomas frequently carry loss-of-function mutations of AIP, a chaperone protein that, among several other functions, modulates AHR transcriptional activity (2). In the present study, we tested the hypothesis that the direct activation of AHR by carcinogens might result in increased risk of pituitary adenoma development. The study indicates no statistically significant increase in the prevalence of the disease due to the massive exposure to TCDD. We are confident that the study was not affected by major selection bias, although the small population size for this particular tumor type is an obvious study limitation. First, zone categorization received support from results on blood dioxin measurements, which clearly indicated that there was a definite dose gradient \( A > B > R \) (13, 14). Interestingly, the patient with prolactinoma in zone A was one of the 182 subjects who developed chloracne, the typical skin disorder due to halogenated hydrocarbon compounds, which was associated with very intense exposures to dioxin in the contaminated zones.

PRL-oma, prolactin secreting adenoma; NFPA, nonfunctioning pituitary adenoma.

* Tetrachlorodibenzo-p-dioxin (parts per trillion, ppt) measured in serum samples collected between 1977 and 1978 from subjects present at the accident in the contaminated zones.

<table>
<thead>
<tr>
<th>Exposure zone</th>
<th>Median serum dioxin levels (ppt)*</th>
<th>Gender</th>
<th>Age at diagnosis (years)</th>
<th>Year of diagnosis</th>
<th>Tumor type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zone A N = 804</td>
<td>447</td>
<td>Female</td>
<td>33</td>
<td>1993</td>
<td>PRL-oma</td>
</tr>
<tr>
<td>Zone B N = 5941</td>
<td>94</td>
<td>Female</td>
<td>55</td>
<td>1993</td>
<td>NFPA</td>
</tr>
<tr>
<td>Zone R N = 38 624</td>
<td>48</td>
<td>Male</td>
<td>47</td>
<td>1986</td>
<td>PRL-oma</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Female</td>
<td>22</td>
<td>1996</td>
<td>PRL-oma</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Female</td>
<td>16</td>
<td>1990</td>
<td>PRL-oma</td>
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<td>48</td>
<td>1990</td>
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<td></td>
<td></td>
<td>Male</td>
<td>71</td>
<td>1994</td>
<td>NFPA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male</td>
<td>47</td>
<td>1995</td>
<td>PRL-oma</td>
</tr>
</tbody>
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doubling of incidence of pituitary adenomas in 1991 in comparison with the previous data in 1958 (20). It is unknown whether this rise in incidence was due to access to better diagnostic techniques, increased medical awareness, environmental toxins and disruptors, or a combination of these factors. The present results do not provide evidence for a major role of specific environmental carcinogens, such as dioxin, polycyclic aromatic hydrocarbons, and polychlorinated biphenyls, in increased incidence of pituitary tumors. However, our study showed a tendency toward an elevated risk of developing pituitary tumors in the most polluted zone A and in the zone of intermediate pollution (zone B) in comparison with the nonexposed reference population. This trend (although based on only three cases) deserves some considerations. First, it is likely that this tendency did not reach statistical significance due to the limited number of exposed subjects and the low incidence of pituitary tumors in the general population. Moreover, the study was a longitudinal study conducted ~20 years after the accident, a latency that was appropriate for the development of diseases with high proliferative potential, such as lung, digestive, lymphatic, and hematopoietic cancers (9). However, considering the indolent nature of pituitary adenomas, this period of observation might still be inadequate. Extended follow-up of pituitary tumor incidence in the Seveso cohort is ongoing.

Pituitary tumors observed in the polluted zones were prolactinomas and NFPAs, while it is well established that AIP mutations are present in families with somatotroph adenomas, in families with both somatotroph and lactotroph adenomas and rarely in families with other types of pituitary tumors (2–4). This difference may be due to either the lack of correlation between exposure to dioxin and pituitary adenoma development or, alternatively, to the multiplicity of proteins, other than AHR, that interact with AIP. In particular, it has been recently demonstrated that naturally occurring AIP mutations lead to disrupted interaction between AIP and phosphodiesterase-4A5, resulting in deregulation of cAMP degradation (3, 21). Since cAMP is a proliferative signal for somatotrophs and not for the other pituitary cell lineages (22), it is tempting to speculate that impairment of phosphodiesterase-4A5 activity might be one of the crucial events for somatotrophs proliferation in familial cases of acromegaly with AIP mutations.

In conclusion, in a unique investigation based on a health surveillance program in an area of relatively pure exposure to dioxin, we did not find statistically significant increase in the prevalence of pituitary tumors. However, the tendency toward a higher risk of pituitary tumors in subjects exposed to high–intermediate dioxin concentrations in comparison with the nonexposed population strongly indicates the need for extended follow-up.

**Declaration of interest**

The authors declare that there is no conflict of interest that would prejudice the impartiality of this scientific work.

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