Effects of Dioxin Exposure on Thyroid Hormones of Populations Living in hot Spots of Dioxin Contamination in Vietnam

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Received date: July 24, 2015; Accepted date: August 17, 2015; Published date: August 25, 2015

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Abstract

Dioxin is still long persistent in environment. The effect of Dioxin on thyroid function is inconsistent. Da Nang and Bien Hoa airbases have been hot spots for persistent contamination since 1961. Inhabitant living around these sites has been exposed to dioxin for a long time. Serum dioxin level of people was much higher than those in other sites. This study evaluates serum dioxin level using DR CALUX and thyroid function including T3, T4, FT3, FT4, TSH measurement. T3 level was found association with serum dioxin level. There were not any relation between T4, FT3, FT4 and TSH and dioxin level.

Keyword:

Dioxin; Thyroid; Da Nang; Bien Hoa

Introduction

Dioxins [polychlorinated dibenzo-p-dioxins, dibenzofurans (PCDD/Fs)] are highly toxic environmental contaminants, which are lipophilic and resistant to biodegradation. These compounds enter the human body through the food chain and finally accumulate in adipose tissue [1]. A series of deleterious health effects are thought to be associated with dioxins exposure. From some studies on animals, as well as on some populations worldwide, dioxins were considered as endocrine disruptors, in which thyroid gland was one prominent target. In animal models, maternal exposure to TCDD induces elevated b-TSH and prenatal primary hypothyroidism. TCDD and other related compounds have been shown to accelerate thyroid hormone clearance by increasing metabolic enzyme activity and competing with plasma binding proteins [2-4]. The effects of dioxin on endocrine system may trigger other health outcomes such as infant growth retardation and developmental abnormalities, altered reproductive capacity [2,3]. Da Nang and Bien Hoa airbases were considered as hot spots of dioxin contamination in Vietnam. These former US airbases had served as sites for transportation and storage of great amount of herbicide for herbicide spraying program in southern Viet Nam that extended from 1962 to 1971. These two airbases were extremely contaminated with dioxins due to the tremendous amount of herbicide stored and spilled during mixing and loading there. Populations residing around these airbases were at high risk of exposure. Tai et al. [4] recently showed that levels of PCDDs/Fs in the breast milk of mothers residing near hot spots were threefold to fourfold higher than those in the breast milk of mothers living in unsprayed areas, and that infant daily dioxin intake (DDI) of Vietnamese infant in hot spots were estimated to be twofold to threefold higher than the recently documented values in US and Japanese infants [4]. In this study, we want to clarify the effects of dioxin exposure on thyroid function of general populations living in Da Nang and Bien Hoa airbases.

Materials and Methods

Subject

Subjects were recruited from general populations living around Da Nang and Bien Hoa airbases. In 2012, total 114 people, including 57 males and 57 females, were born before or during and after the period of herbicide spraying (1962-1971) participated in the survey. Demographic information of subjects was collected by an interview. A blood sample from each subject was collected in the morning of examination day. The serum was extracted from whole blood and stored at -20°C until analysis of hormones and dioxin levels.

We defined the body mass index (BMI) as weight (kg) divided by the square of height (m).

T3, T4, FT3, FT4, TSH measurement

The estimation of T3, T4, FT3, FT4, TSH were done by electrochemiluminescence method on Roche Elecsys 2010 instrument, using Elecsys and cobas e analyzers kit, with guidelines for the reference ranges of serum TSH, T3, T4, FT4, FT3 were as 0.27-4.3 µIU/ml, 1.27-3.07 nmol/l, 71.5-158 nmol/l, 12.7-20.8 pmol/l, 3.89-6.66 pmol/l, respectively. For the analysis purpose, the values that were below detection limits were set as half of detection limit.

Determine dioxin level in serum by DR CALUX

The DR CALUX-bioassay analysis was performed in Dioxin laboratory of Vietnamese Military Medical University, which had been certified for approval to perform DR CALUX analysis by Bio-Detection System, Amsterdam, Netherlands. Approximately 2 ml of serum was used for fat extraction by n-hexane (Sigma) and 2-propanol (Merk). Fat content was weighted before clean-up step, in which acid-labile matrix components were removed by passage through a silica (Sigma) column containing two layers: 20% and 33% (w/w) concentrated H2SO4 (Merk). This extract was dried and then diluted in dimethyl-sulphoxide (DMSO) (Sigma) before exposing to rat H4IE hepatoma (H4L1.1c4) cells (BioDetection System, Holland).
cells stably transfected with an AhR-controlled luciferase reporter gene construct (pGudluc1.1) and were grown confluent in 96-well view plates. Samples, TCDD standards (Bio-Detection System, Holland) and internal control were exposed in triplicate for 24 hours in the same plate, using DMSO (0.8% v/v) as a vehicle. After removal of the medium (Sigma), cells were washed twice with phosphate-buffered saline (Oxoid, Hampshire, UK). The cells were harvested in 30 μl cell lysis reagent (Luciferase Assay System; Promega, Leiden, The Netherlands). A linear standard curve was built and used to calculate dioxin level in samples, which finally expressed in a pg unit of total bioanalytical equivalent (BEQ)/g fat.

Statistical analysis: we used median of Dioxin level (69.16 BEQ) as cut off to divide the subjects into two groups, high category and low category. In general analysis, data were expressed as frequency with percentage for categorical variables. For comparison of difference in percentage between groups, OR, CI, χ2 was performed. All comparisons were considered under a significance level 0.05. All data were analyzed using STATA version 12.0.

### Results

| High | Low | Total |
|------|-----|-------|
| n (%) | n (%) | n (%) |
| Mean BEQ | 106.38 (35.34) | 38.67 (18.4) |
| Range | 70.58 - 209.54 | 5.40 - 67.75 |

| Age group | n (% |
|-----------|------|
| 20 - ≤ 30 | 2 (50) |
| 30 - ≤ 40 | 3 (50) |
| 40 - ≤ 50 | 16 (48.8) |
| 50 - ≤ 60 | 26 (54.17) |
| >60 | 10 (43.48) |
| Total | 57 (50) |

| Subject | High | Low | Total |
|---------|------|-----|-------|
| TSH | 8 (14.04%) | 10 (17.54) | 18 (15.79) |
| Abnormal | 49 (85.96) | 47 (82.46) | 96 (84.21) |
| Total | 57 (100) | 57 (100) | 114 (100) |

| Subject | n (%) |
|---------|-------|
| OR=0.767, 95% CI: 0.279-2.111, χ2=0.2639, p=0.607 |

| Subject | n (%) |
|---------|-------|
| OR=2.407, 95% CI: 1.222-5.163, χ2=0.519, p=0.023 |

| Subject | High | Low | Total |
|---------|------|-----|-------|
| >69.16 BEQ/g | 57 (100) | 114 (100) |
| n (%) | n (%) | n (%) |
| Age group | | |
| 20 - ≤ 30 | 2 (50) |
| 30 - ≤ 40 | 3 (50) |
| 40 - ≤ 50 | 16 (48.8) |
| 50 - ≤ 60 | 26 (54.17) |
| >60 | 10 (43.48) |
| Total | 57 (50) |

| Subject | High | Low | Total |
|---------|------|-----|-------|
| OR=1.638, 95% CI: 0.782-3.434, χ2=0.4298, p=0.1898 |

| Subject | High | Low | Total |
|---------|------|-----|-------|
| FT3 | 25 (43.86) | 32 (56.14) | 57 (50) |
| Abnormal | 32 (56.14) | 25 (43.86) | 57 (50) |
| Total | 57 (100) | 57 (100) | 114 (100) |

| Subject | High | Low | Total |
|---------|------|-----|-------|
| OR=1.838, 95% CI: 0.782-3.434, χ2=0.4298, p=0.1898 |

| Subject | High | Low | Total |
|---------|------|-----|-------|
| T4 | 16 (28.07) | 18 (31.58) | 34 (29.82) |
| Abnormal | 41 (71.93) | 39 (68) | 80 (70.18) |
| Total | 57 (100) | 57 (100) | 114 (100) |

Table 1: Demographic characteristics by BEQ category.

The subjects were divided into two groups basing on level of dioxins in serum blood. The cut-off value was set equal to absolute median of dioxin concentration (69.19pg BEQ/g fat). Demographic characteristics of the participants in both groups were presented in Table 1. Distribution of age, gender, BMI between high and low groups was not different (p>0.05). The mean dioxin level in high group was approximately 3 times higher than that in low group (106.38 vs 38.67 pg BEQ/g fat).

Effect of Dioxin on thyroid function was showed in Table 2, number of abnormal of TSH, T3, T4, FT3, FT4 in High BEQ category were higher than that in Low category. But number abnormal of TSH, T4, FT3, and FT4 were not significantly difference between two groups. The odd ratios of all thyroid hormone exception of T3 did not increase or decrease significantly between groups. Odd ratio of T3 showed significant association between high category and low category (OR=2.407, CI: 1.222-5.163, χ2=0.519, p=0.023).
Table 2: Abnormal thyroid hormone levels by BEQ category.

| OR         | 95% CI              | p-value |
|------------|---------------------|---------|
| FT4 Normal | 1.183, 95% CI: 0.53-2.642, \( \chi^2=0.0588, p=0.682 \) |         |
| Abnormal   | 1.166, 95% CI: 0.393-3.462, \( \chi^2=0.0768, p=0.782 \) |         |

Overall, results of studies that examined TCDD and thyroid function provide little and inconsistent evidence of long-term TCDD effects on the adult thyroid. Our result was not consistent with those of Ott who found increases in mean total T4 and TBG with TCDD levels [5], Chevrier [3] and Calvert [6]. However, we found that abnormal T3 level significant relation to high category that agreed with observation of Johnson on human [7].

The reasons why we observed the significant association between the risk of having thyroid dysfunction with T3 but not T4, FT4 and TSH were still not explained clearly in previous studies. Chevrier J. reported weak association between T3 and TCDD, with \( p=0.13 \). There are a lot of cause effects of thyroid function and made differences between results of the studies. In this case, the models of previous studies only measurement toxicity of 17 congeners of Dioxin. The substances were part of contribution to TEF. In our study, TEF was measurement based on DR CALUX technique. It can estimate toxicity of all substance of dioxin and dioxin – like compounds, including PCDDs, PCDFs, PCBs. On the other hand, Pavuk [8] used cut-off point (94 ppt for low and high category) higher so much than that in our study, 69.16 BEQ/g (ppt). They might cause of difference among the results.

Some human studies have investigated associations between serum TCDD concentrations and thyroid hormone levels in adults, and results have been inconsistent. Ott et al. reported positive associations between T4 and whole-blood TCDD concentrations measured in 131 trichlorophenol production plant workers more than 45 years after an accidental exposure to TCDD. Chloracne status and TCDD levels estimated by back-extrapolation to the time of the accident were also positively associated with T4 levels. There were no associations with TSH [5]. Other studies have similar result: high serum TCDD concentrations (\( \geq 1,860 \) pg/g lipids) measured more than 15 years after the last exposure were associated with an elevated free T4 index among 2,7,8 trichlorophenol plant workers relative to 257 referents with lower exposure (mean serum TCDD concentration=7 pg/g lipids), although no clear exposure-response relationship was demonstrated. No association was found between TCDD and total T4 or TSH [6]. Result of the study in Australia show that back extrapolated serum TCDD concentrations (but not con-current levels) were negatively correlated with triiodothyronine T3 but not T4 [7]. In a large (n=1,009 exposed and 1,429 referents) prospective study of US Air Force veterans who sprayed TCDD-contaminated Agent Orange in Vietnam between 1962 and 1971, Pavuk et al. reported positive associations between serum TCDD concentrations in 1982–1997 and concomitantly measured TSH levels (but not total T4 levels, free T4 index, or T3% uptake) [8]. Chevrier [3] reported an inverse association between serum TCDD concentrations measured shortly after the Seveso trichlorophenol plant explosion and total T4 (but not free T4, free T3, or TSH) levels measured approximately 20 (but not 30) years later among women who were premenarche at the time of the accident. However, the result showed no clear associations between 1996 serum TCDD concentrations and any 1996 or 2008 thyroid hormone measures, suggesting an effect of initial exposure rather than of later body burden [3].

### Discussions

There has been doubt that the orange agent could increase the thyroid hormone clearance in animal models. The present study investigated the hypothesis that exposure to dioxins could associate with the dysfunctions in thyroid hormones in human by conducting a cohort study in Bien Hoa and Da Nang, Vietnam. Our findings in this study suggested that the dioxins could be associated with the dysfunction of thyroid hormone in terms of T3 plasma concentration, but not for TSH, FT3, T4 and FT4.

Overall, results of studies that examined TCDD and thyroid function provide little and inconsistent evidence of long-term TCDD effects on the adult thyroid. Our result was not consistent with those of Ott who found increases in mean total T4 and TBG with TCDD levels [5], Chevrier [3] and Calvert [6]. However, we found that abnormal T3 level significant relation to high category that agreed with observation of Johnson on human [7].

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Although few data are available on TCDD’s potential for thyroid hormone disruption in women, a number of studies of other polychlorinated dibenzodioxins, dibenzofurans, and biphenyls whose dioxin-like activity was evaluated using toxic equivalents (TEQs) have been conducted, with similarly inconsistent results [9-13].

A recent study on approximately 225,000 veterans of the Vietnam era found that those who served in Vietnam or were otherwise exposed to defoliants had a 2.5-fold to threefold higher prevalence of the diagnosis of Graves’ disease, compared to Veterans who served elsewhere [14] non-significant increase in mean total T4 and a significant increase in mean FTI was found in exposed US workers, but the risk of abnormally high FTI was not increased [6]. Some report dioxin decrease total T3 serum level. Another result see changes in total T3 and total T4 [15,16]. Meanwhile, other results did not show any effects of dioxin on thyroid function [17-19].

One of limitation of this study was the lack of a control cohort including people who lived in assumed unexposed areas. This drawback, however, could be mitigated by comparing the risk of having thyroid dysfunction between the groups of having low and high serum dioxin level. Furthermore, the serum dioxin levels served as a reliable measurement for the level of dioxin exposure. Thus, the validity of the study was still guaranteed.

In conclusion, the findings in our study indicated that the dioxins exposure was associated with thyroid functions in term of T3. Further studies with larger sample size, however, should be conducted to confirm the results.

### Acknowledgements

The authors thank Mrs. Nguyen Thi Hien, President of VAVA Da Nang for their valuable help during field surveys; medical staff in the 17 Hospital in Da Nang and Medical staff in Public Health department of Bien Hoa city.

Funding sources was supported by Project from Ministry of Natural Resources and Environment.
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