Aspects of formation and degradation of polychlorinated dibenzo-p-dioxins and dibenzofurans

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Abstract The research progress on the mechanisms of formation and degradation of polychlorinated dibenzo-p-dioxins (**PCDDs) and dibenzofurans** (**PCDFs) is reviewed. The primaty sources of PCDD/Fs are chemical, thermal, photochemical and enzymatic reactions. Most of the thennal sources result in emissions into air. The studies on the photodegradation and biodegradation offer promise for destruction of PCDD/Fs in environment.**

 $Keywords: polychlorinated dibenzo-p-dioxins, polychlorinated dibenzoformates, mechanism, formation, degradation.$

POLYCHLORISATED dibenzo-p-dioxins (PCDDs) and dibenzofurans (PCDFs) are two series of almost planar tricyclic aromatic compounds with very similar chemical properties. Different degrees and positions of chlorination on the aromatic ring structures can occur, leading to 75 PCDD and 135 PCDF isomers in total. The most toxic isomers are 17 congeners that are fully chlorinated in the 2, 3, 7, 8-positions. A draft of four- ear-long (1991-1994) US EPA reassessment **of** PCDD/Fs restated findings that PCDD/Fs is a probable human carcinogen. And it broke new ground in its discussion of noncancer health effects of the compounds such as disruption of the endocrine, reproductive, and immune systems^[1]. Due to environmental persistence of PCDD/Fs, these classes of compounds have ubiquitous distribution. The primary exposure of humans to these lipophilic compounds is via the food chain.

Many efforts have been made at various laboratories to present formation or degradation mechanisms and derive technologies for minimization of these compounds. Although extensive investigation on PCDD/ Fs in environment has not been canied out in China, some reports concerned about a few pollution areas revealed that PCDD/Fs released through chemical processes have resulted in serious environmental pollution in China^[2-9]. It is important to investigate the characteristic of mechanisms of formation PCDD/Fs in China and explore methods of solving the problem of PCDD/Fs pollution.

1 The natural production of PCDD/Fs

The production of chloromethane and other organochlorine compounds, when organic material is **burned, in association with the omnipresent chloride ion, led to the belief that PCDD/Fs could form during combustion** processes in nature. Laboratory studies have revealed that PCDD/Fs form in parts-per-billion amounts during the combustion of wood. It is estimated that about 59 kg of PCDDs is produced in Canadian forest fire annually^[10,11]. Thomas reported that about 0.2 kg toxicity equivalent (TEQ) is produced annually in the U.S. forest fires^[12]. Because there are 20 000 forest fires annually worldwide that burn 43 000 square kilometer, it is logical to assume that PCDD/Fs have been present in the environment for many centuries $^{[13]}$.

Another milestone observation is the enzymatic conversion of chlorophenola into PCDD/Fs in the parts-per-million range by horse-radish peroxidase enzyme (HRP). The equation is given below^[14].

PODS PCDFs

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This extraordinary revelation opens the door to the possibility that a source of environmental PCDD/Fs may be their completely natural formation from (natural) chlorophenols by soil and water microbes. Recent evidence shows that these reactions also occur under true environmental conditions in sewage sludge and peat $\text{bogs}^{[15,16]}$.

The formation of PCDDs from pentachlorophenol (PCP) in wood matrix and water by photolytic condensation under natural sunlight was demonstrated by Lamparski *et al*. $[17,18]$. The equation is shown below.

Kjeller *et al.* analyzed sediment core from the northwestern part of the Baltic ${Proper}^{\lceil 19 \rceil}$ and $2\,000$ herbage collected between 1861 and 1993 from a $\frac{1}{18}$ 1600 well characterized rural site in southeast Eng-
land^[20]. The results showed that PCDD/Fs concentration peaked in the 1960s and 1970s and have declined sinc well characterized rural site in southeast Eng- $\frac{8}{6}$ 1 200
land^[20]. The results showed that PCDD/Fs concentration peaked in the 1960s and 1970s and have $\frac{8}{5}$ 800 declined since then. Fig. 1 shows the levels of **3** ⁴⁰⁰ PCDD/Fs in a sediment core from Baltic Proper. The US EPA's draft of dioxin risk reassessment and the search of dioxin risk reassessment and the search in th stated that ancient human tissue sampling shows much lower PCDD/Fs levels than found today. **Average age (year)** Studies of sediment cores in the lakes near industri- **Fig. 1.** Total levels of tetra- to octa PCDD/Fs for each disk of were quite low till about 1920. These studies show tal PCDD. increases in PCDD/Fs concentrations beginning in

a1 center of the U.S. have shown that PCDD/Fs the sediment core from Baltic Proper. \Box **, Total PCDF;** \blacksquare **, to-**

the 1920s and continuing till about $1970^{[1]}$. On these bases, it appears that the presence of PCDD/Fs in the environment occurs primarily as a result of anthropogenic practices.

2 Anthropogenic emissions of PCDD/Fs

PCDD/Fs are not produced intentionally, but are released into the environment in ultratrace amounts from various combustion processes and as a result of their occurrence as unwanted byproducts in various chlorinated chemical formulations.

 (i) Chemical processes. Many chemical reactions may result in the contamination of pesticides and technical products including chlorophenols , chlorophenoxy herbicides and polychlorinated biphenyls (PCBs). The production and use of these chemicals are nowadays banned or strictly regulated in most countries, but during the 1960s and 1970s these products were widely used and hence became a major source of PCDD/Fs contamination for the environment.

Chlorophenol has been used as preservative and biocide for many years. Although there were no accurate annual production figures for individual chlorophenols pesticides, the total production of **all** pesticides of this type was estimated to be 1.5×10^8 kg in 1980. Figures on PCP are available for the U.S. between 1972 and 1977. Average of annual production during this period was about 2×10^{7} kg. The PCDD/Fs content of commercial PCP mixtures was measured to be in the range of 205-1 070 ppm. Using these figures and the U.S. production estimates, it can be calculated that $4 \times 10^{3} - 21 \times 10^{3}$ kg of PCDD/ Fs was produced annually in the U.S. during PCP manufacture^[21].

Being contaminants of technical PCP and Na-PCP, it is estimated that the environment emission of

PCDD/Fs was about 240 kg PCDD/Fs or 1 kg TEQ annually from the 1970s to the 1980s in China. Na-PCP has been spread since the 1960s to control the spreading of snailborne schistosomiasis in the south of China. The analysis results of samples from these areas show that the levels of PCDD/Fs are significantly high^[7,8]. Bao reported findings about 10% of PCDD/Fs in the waste from reaction still for production of PCP by pyrolysis of hexachlorocyclohexane (HCH). It can be calculated that 1.5×10^4 kg PCDD/Fs or 90 kg TEQ is produced annually at the factory from 1990 to 1993.

Zheng et al. studied the mechanism of formation of PCDD/Fs from the pyrolysis of HCH. It was revealed that FeCl₃ plays an important part in formation of PCDD/Fs from the pyrolysis of HCH. The pattern of PCDD/Fs from products of the laboratory experiments is similar to the waste of the chemical factory.

Maximum production of 2,4,5-T was reached between 1960 and 1970, with the total of 48.2×10^6 kg produced in the world. The concentrations of 2, 3 , 7, 8-TCDD in 2,4,5-T have been determined in several laboratories. If we assume that the upper 2, 3, 7, 8-TCDD concentration limit was 100 ppm and the lower limit was 0.1 ppm, then the maximum amount of 2, 3, 7, 8-TCDD produced from 1960 to 1970 was about 4.8×10^3 kg. More than 11 million gallons of Agent Orange (1:1 w/w 2, 4-D and 2, 4, 5-T) were sprayed on jungles of Vietnam. Many people still suffered greatly while exposed to PCDD/Fs there.

PCBs production increased greatly at the end of the sixties, but by the end of the seventies, production decreased sharply due to environment concerns. The U.S. produced 58.9 \times 10⁶ kg commercial PCB mixture, Aroclors 1 248, 1 254 and 1 260, from 1968-1970, that was about half of the world total. It can be estimated that a total of 84-94 kg of PCDFs was produced from production of PCBs in the U.S. during this period^[21].

The Chinese commercial PCBs, known as PCB₃ and PCB₅, were found to contain PCDD/Fs at about 217 -417 ng TEQ/g. Approximately 10×10^6 kg commercial PCBs was produced from 1965 to 1974 in China. Most of these PCBs products were used as dielectric fluid in electronic capacitors, and a small part of which was used as additives in paints. The electronic capacitors containing PCBs have been considered as solid wastes since the 1980s.

(ii) Pulp bleaching. During Dioxin's6 in Japan Rappe reported findings of PCDD/Fs from samples of crab hepatopancreas and sediments collected outside a pulp mill in Sweden. The levels of 2, 3, 7, 8-TCDD and 2, 3, 7, 8-TCDF were at least ten times higher in the samples collected outside the mill than in the background samples. Changing bleaching processes and wastewater treatment enabled the industry to reduce the amount of PCDD/Fs by more than 90% between 1988 and 1992 in the U. S. A. A 98% decrease in PCDD/Fs originating from bleaching operations over the period from 1988 to 1993 for the entire Canadian chemical pulping sector was observed $[25]$.

Non-wood plant fibers in paper making nowadays in China are dominant. Bleaching of non-wood plant fibers in China usually requires only a single step of hypochlorite bleaching process. The levels of PCDD/ Fs in the bleached pulp from five pulp mills are about 30 pg TEQ/g dry pulp¹⁶¹.
(iii) Metal processing. It was reported that PCDD/Fs could be identified in dust from a steel mill

in Sweden at levels of 800 pg TEQ/g dust in 1986. A study in a pilot plant reported collection of smoke containing 110 ng TEQ/m^3 , where scrap metal was melted and recycled using batchwise charging. The Swedish EPA considers the production of iron and steel to be the major source of PCDD/Fs in Sweden^[15]. It is estimated that 50-150 g TEQ PCDD/Fs was produced annually during metal processing in the U. $S^[26]$. PCDD/Fs are emitted by the burning of scrap metal such as copper cable coated with PVC plastic insulation. In fly ash from a copper wire and aluminum scrap recovery facility, PCDD/Fs concentrations are 50⁻⁴⁶⁰ ppm^[27].
(**iv**) Combustion processes. The emissions of PCDD/Fs from incineration processes were first ob-

served by Olie et $al.$ ^[28]. Since then, numerous measurements have been performed in a variety of combustion sources, and trace quantities of PCDD/Fs emissions have been found in most combustion sources. Municipal solid waste incinerators (MSWI) have been suggested **as** most serious PCDD/Fs emission sources judging from the high PCDD/Fs concentrations in flue gas and flue gas discharged from incinera-

tors, and therefore, have been the subject of major research efforts to reduce PCDD/Fs pollution. The U. S. EPA's dioxin reassessment showed that waste combustion accounted for about 95% of all the known PCDD/Fs emission. Medical and municipal waste combustion dominates combustion sources. The U.S. PCDD/Fs emissions from combustion sources are shown in table $1^{[12]}$.

A better understanding of the mechanism of PCDD/Fs formation provides a better guidance for devising technical control measures and improving the performance of practical combustion equipment in order to reduce PCDD/Fs emissions. Many incinerators reduce the emission of PCDD/Fs to a level below 0.1 ng TEQ/ m^3 which is the guideline in many European countries^[15].

Table 1 The USA annual PUDD/ is emissions from combustion sources $\{g \}$ FLU/year)					
Medical waste	Municipal waste	Petroleum	Industrial wood	Residential	Coal combustion
incineration	incineration	combustion	purning	wood burning	
5 000	2 000	90	150	40	

Table 1 The USA annual PCDD/Fs emissions from combustion sources (**TF.Q/year)**

In spite of 20-year research and obtained insights in the reactivity of fly ash towards PCDD/Fs formation, the mechanism of formation of these compounds remains unclear^[30]. Recent discussions of the possible mechanisms of PCDD/Fs formation have focused on two hypotheses: $| \cdot \rangle$ Dickson et al. have proposed that PCDD/Fs are formed from chloroaromatic precursors such as chlorophenols^[31-33] and chlorobenzenes^[34] by reactions which have been shown to occur by heterogeneous catalysis on the surface of fly ash particles at **250-400T** . These products could be formed by Ullman-type surface reaction. The precursors might be already present in the fuel, or they could be formed in the higher temperature postcombustion zone by multistep reactions, including aromatization of aliphatic cornpounds and subsequent chlorination by molecular chlorine^[35]. A source of Cl for the organic precursors may be provided through the metal catalyzed Deacon reaction:

$$
2HCl + \frac{1}{2}O_2 \rightarrow H_2O + Cl_2
$$

Chlorophenols and chlorobenzenes can also be produced from heterogeneous combustion reactions of C_2 aliphatics[M1. ii) Stieglitz *et al.* have suggested a **de** novo synthesis of PCDD/Fs from active carbon particulates by gas-solid and solid-solid reactions with air, moisture and inorganic chlorides, catalyzed by Cu (\prod) ions^[37,38].

Dickson *et al.* demonstrated that PCDDs are formed from the model precursor compound PCP much faster than from reactions **of** particulate carbon in the heterogenous phase under identical conditions. In the postcombustion and heat exchanger sections of an incinerator fast reactions of chlorophenols and chlorobenzenes precursors leading to PCDD/Fs formation are expected to predominate. In pollution control equipment slower processes such **as de** now, synthesis from particulate carbon may contribute to the levels and distribution of PCDD/Fs congeners and other chlorinated organic compounds in incinerator emission^[35].

De novo synthesis led to the proposal of particulate carbon-present in all fly ashes after extraction-as most likely precursor. If true, questions then **arise** : what is the nature of this carbon, and how is it mobilized from the solid carbon matrix to form aromatic moieties? The previous work by Stromberg *et* al. suggested that CO₂ could act as a precursor to chlorobenzenes under proper conditions via a Fisher-Tropsch type mechanism^[40,41]. However experiments with ¹³C-labeled CO and CO₂ reacting with fly ash showed that neither is an effective precursor to $PCDD/Fs^{[42]}$.

Froese et *al.* showed that heterogeneous combustion reactions **of** acetylene produce chlorobenzenes, chlorophenols and $P CDD/Fs^{39}$. This experiment hints that there is relationship between the two postulations^[39]. A simplified schematic of conceivable *de novo* and precursor PCDD/Fs formation pathway is summarized in figure 2.

Fig. 2 . **Potential** PCDD/ **Fs formation mutes in incineration.**

3 PCDD/Fs **emissions** in **some countries**

Table 2 shows PCDD/Fs emissions estimates from the known sources.

These figures are based on limited data and must be considered as very rough estimates.

4 Studies on the degradation of PCDD/Fs

(i) Photodegradation of PCDD/Fs. i) Photolysis of PCDD/Fs in solvents. It is known that PCDD/Fs decompose in organic solution if they are irradiated by UV-light of 250 -315 nm. Grosby et al. found that 2 , 3 , 7 , 8 -TCDD undergoes rapid photolysis in methanol under UV light^[43]. Buser demonstrated that dechlorination of higher chlorinated PCDD/Fs to lower chlorinated ones appears to be the major reaction pathway. fie products of OCDD photodegradation suggested a preferential loss of the chlorine atoms in lateral $(2-$, $3-$, $7-$, $8-$) positions on the dibenzo-p-dioxin ring^[44]. Experiments showed that the rate of photolysis of OCDD is greatly affected by the surrounding solvent medium. The **measured** first order rate indicates that photolysis occurs more rapidly in solvents that are good H donors^[45]. Zheng et al. reported that photolysis of PCDDs in carbon tetrachloride with UV light produces the corresponding chlorobenzenes. C-0 bond rather than C-C1 bond cleavage is found to be a major pathway for photodegradation of PCDDs^[46]. Photolysis of PCDDs in chlorform by a 300 W medium pressure mercury Vapor lamp showed that half-lives of the compounds are less than 5 $min^{[47]}$.

The mechanism of photolytic degradation of PCDD/Fs in water is not clear. The photolysis of 2, 3, 7, 8-TCDD appears to be considerably less efficient in water than in hexane at 313 nm^[48]. However. phdolysis **rates** of 2, 7-DCDF, 2, 3 , 7 , 8-TCDF and 1 , 2, 7, 8-TCDF in aqueous solution irradiated at 300 **nm are** 4-15 times faster than in organic solutions, and in the cases of 2, 7-DCDF and 1, 2, **7, 8-** TCDF, 20 times faster than in 60% acetonitrile/water. Dechlorination may be a major process in the pho-

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todecomposition of higher chlorinated PCDFs in hydrocarbon solvents but not in water^[49]. Several studies have shown that the photolysis rates of both PCDDs and PCDFs may be significantly enhanced in natural water, a fact usually attributed to the presence of naturally occurring sensitizers^[50]. Although the identity of the sensitizers is unknown, dissolved fulvic and humic acids and proteins have been shown to sensitize the photodegradation of organic contaminants in aqueous solutions.

ii) Photolysis of PCDD/Fs at the surface of solids. Strong public concern posed by PCDD/Fs has spurred research efforts to develop treatments to eliminate these hazards. Several successful techniques that have been reported for contaminated soil sites involve difficult and expensive procedures. The needs for practical, lower cost techniques still exist. The facts that photolysis of OCDD on soil by UV irradiation produces 2, 3 , 7 , 8 -TCDD suggested a preferential loss of the chlorine atoms in ortho $(1-, 4-, 6-, 9-)$ positions^[51,52]. Many studies were made to estimate the theoretical maximum rate for soil decontamination. Mathematic analysis of these studies showed that the average decontamination rate is 3.3 ng TCDD/ h $~cm^{2[53]}$. Since photolysis can occur only in a shallow zone at the surface of soil in the depth of 0.06-0.13 mm, PCDD/Fs molecules must first be brought into solution and then transported to the surface by solvent related processes. Substantial photodegradation $(60\% -85\%)$ of TCDD throughout 6 cm deep soil column was found after 60 d sunlight exposure^[54]. Yan *et al*. reported the effect of cocontaminants in soil on the photodegradation of PCDDs. The cocontaminants could be quenchers or sensitizers and change the efficiency of **phototransformation[55.561** .

Hosoya *et al.* used octadecylsilylated silica gel (C_{18}) as a reaction medium for the photolysis of 2, 3,7,8-TCDD in 10% 2-propanol/water. The compound decomposed completely in 5 min after being exposed to a 450 W UV lamp. It is demonstrated that complete C_{18} trapping with continuous photodecomposition of TCDD contained in an aqueous alcohol waste is possible^[55]. Schuler *et al.* investigated the photodegradation of PCDD/Fs dissolved in cuticular wax from laurel cherry leaves. Application of the results to an exiting model of atomspheric deposition of aerial contaminants demonstrated that photodegradation is an important process in the fate of PCDD/Fs on vegetation and in the use of vegetation as a tool for **air** monitoring of PCDD/ $\text{Fs}^{[57]}$.

iii) Advanced oxidation processes. Advanced oxidation processes have already found a widespread application in the treatment of industrial wastewater, ground water and landfill leachate. Zhang *et a1* . used $TiO₂$ to catalyze the photolysis of PCDDs in an aqueous system. The results showed that PCDDs tested in the experiment were destroyed by $84\% - 91\%$ within 4 h^[58]. The quantum yield of OCDD photodegradation is increased about fourfold when 0.3% H₂O₂ has been incorporated into the solution^[56]. Vollmuth *et a1* . expected a better way to destroy PCDD/Fs contaminants in seepage water by UV/ozone treatment, but there is still a long way to go^[59]. Sommer *et al.* investigate the photochemical decomposition of PCDD/Fs on fly ash particles from municipal solid waste with medium and low pressure mercury lamps in an oxygen/ ozon atmosphere. The maximum value of the degradation rates is about $70\%^{60}$.
(ii) Biodegradation of PCDD/Fs. Bioremediation strategies for PCDD/Fs contaminated environ-

ments would be a highly desirable approach, as it may be considerably less costly than physical-chemical processes, and as it can be used in *situ.* However very few bacterial or fungal species are known to degrade PCDD/Fs. The half-life of TCDD in soil in the absence of UV light is approximately 10 years, indicating the general lack of alternative degradation pathway. Du *et al.* demonstrated that 8 bacteria strains screened from contaminated soil could degrade 2-chlorinated dibenzo-p-dioxin well, 1 of the 8 bacteria strains could degrade 2, 3, $7-T_3CDD^[62]$. The fate of PCDD/Fs was studied in anaerobic microcosms, using PCB contaminated sediments, and creosote contaminated aquifer sediments. The PCDD/Fs concentrations in active microcosms were shown to decrease at higher rates than in chemical and biological controls. Peri-dechlorinated was found under anaerobic methanogenic conditions^[63]. Disse compared PCDD/Fs concentrations after aerobic and anaerobic digestion of sewage sludge. The aerobic digestion occurred in ventilated batch reactors leading to a significant degradation of $PCDD/Fs^{[64]}$.
(iii) Other methods for the destruction of $PCDD/Fs$. Hilarides *et al.* showed that gamma irradia-

tion is both a technically and economically feasible method to degrade PCDD/Fs. Approximately 99% degradation of 2, 3, 7, 8-TCDD on a model soil was achieved using gamma radiation at a dose of 800 k~~[~~~ . Oku used alkali-metal hydroxide in **1,3-dimethyl-2-imidazolidinone** to destroy PCDD/Fs. Highly efficient destruction ($> 99.99\%$) of PCDD/Fs was achieved^[66].

Acknowledgement This work was supported by the Chinese Academy of Sciences and National Natural Science Foundation of China (Grant No. 29707003) .

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(Received September **11, 1998)**