The hydroxyl radical (·OH) has been considered to be one of the most reactive oxygen species produced in biological systems. It has been shown that ·OH can cause DNA, protein, and lipid oxidation. One of the most widely accepted mechanisms for ·OH production is through the transition metal-catalyzed Fenton reaction. Pentachlorophenol (PCP) was one of the most widely used biocides, primarily for wood preservation. PCP is now ubiquitously present in our environment and even found in people who are not occupationally exposed to it. PCP has been listed as a priority pollutant by the U.S. Environmental Protection Agency (EPA) and classified as a group 2B environmental carcinogen by the International Association for Research on Cancer (IARC). The genotoxicity of PCP has been attributed to its two major quinoid metabolites: tetrachlorohydroquinone and tetrachloro-1,4-benzoquinone (TCBQ). Although the redox cycling of PCP quinoid metabolites to generate reactive oxygen species is believed to play an important role, the exact molecular mechanism underlying PCP genotoxicity is not clear. Using the salicylate hydroxylation assay and electron spin resonance (ESR) secondary spin-trapping methods, we found that ·OH can be produced by TCBQ and H₂O₂ independent of transition metal ions. Further studies showed that TCBQ, but not its corresponding semiquinone radical, the tetrachlorosemiquinone radical (TCSQ), is essential for ·OH production. The major reaction product between TCBQ and H₂O₂ was identified to be trichloro-hydroxy-1,4-benzoquinone (TrCBQ-OH), and H₂O₂ was found to be the source and origin of the oxygen atom inserted into this reaction product. On the basis of these data, we propose that ·OH production by TCBQ and H₂O₂ is not through a semiquinone-dependent organic Fenton reaction but rather through the following novel mechanism: a nucleophilic attack of H₂O₂ to TCBQ, leading to the formation of an unstable trichloro-hydroperoxyl-1,4-benzoquinone (TrCBQ-OOH) intermediate, which decomposes homolytically to produce ·OH. These findings represent a novel mechanism of ·OH formation not requiring the involvement of redox-active transition metal ions and may partly explain the potential carcinogenicity of the widely used biocides such as PCP and other polyhalogenated aromatic compounds.
PCP is a potent carcinogen. Following chronic exposure of B6C3F1 mice to PCP, hepatocellular carcinomas or adenomas, hemangiosarcomas, and phaeochromocytomas were observed in F334/N rats (5). In contrast, in the blood of occupationally exposed workers, the median level of PCP was found to be as high as 19580 ppb (range, 6000–45200 ppb) (3). Postmortem analysis of serum, tissue, and urine samples from individuals who died from PCP intoxication showed tissue PCP concentrations of 20–140 ppm (ppm) and urine concentrations of 28–96 ppm. The most likely source of exposure is PCP-treated wood products by way of the food chain. In groups of individuals who are not specifically exposed to PCP, net daily intake estimated in eight countries varied from 5 to 37 µg. Net intake was between 51 and 157 µg per day in residents of homes made of PCP-treated logs (3). While PCP itself does not show any reactivity toward DNA, its thiol and glucuronide conjugates were also found in animals (11, 15).

Redox cycling of compounds with a quinoid structure is a well-known phenomenon (12, 13). The cyclic (auto)oxidation and reduction reactions with the intermediary formation of semiquinone radicals can produce large amounts of reactive oxygen species (ROS) by reducing molecular oxygen to superoxide (O$_2^-$), which in turn can induce oxidative stress (12, 13). However, it is generally accepted that O$_2^-$ itself does not directly attack DNA but only after its dismutation to hydrogen peroxide (H$_2$O$_2$) and subsequent metal-mediated cleavage to hydroxyl radical (·OH). This reaction sequence is called the Haber–Weiss reaction or superoxide-driven Fenton reaction (M represents transition metals, especially iron and copper) (14):

$$2O_2^- + 2H^+ \rightarrow H_2O_2 + O_2$$

$$O_2^- + M^{n+} \rightarrow O_2 + M^{(n-1)+}$$

$$H_2O_2 + M^{(n-1)+} \rightarrow \cdot OH + OH^- + M^{n+}$$

While PCP itself does not show any reactivity toward DNA, TCHQ was found to induce single strand breaks in isolated DNA (10), a variety of cell lines (11, 15–17), and liver of mice (18, 19). TCHQ also induced micronuclei and mutations at the
lating DNA and forming DNA adducts (24, 28, 29). Recently, Lin et al. showed (28) that low degree of oxidative and direct DNA damage was produced by high concentrations of TCBQ (1–5 mM). However, the molecular mechanisms underlying such damage are still unknown.

1.3. Pro-oxidant vs Antioxidant Equilibrium. The pro-oxidant potential of biological systems and their antioxidant capacity usually are in an approximate equilibrium (14). In normal cells, a primary defense against oxidative damage is provided by small molecule antioxidants such as glutathione (GSH) and ascorbate, which are present in millimolar concentrations. However, these defense mechanisms can be overwhelmed by xenobiotics such as PCP and its metabolites that induce the production of excessive ROS, which can result in damage to biological macromolecules such as DNA (12–14).

It should be noted that the concentration of ROS, including H2O2, in cells under normal physiological conditions is low, but ROS concentration may be significantly increased in cells that are subjected to oxidative stress conditions such as exposure to PCP metabolites, as we showed recently in NIH 3T3 cells (30). TCHQ treatment was shown to cause more than 60% GSH depletion in liver tissues of mice, possibly by forming GSH conjugates (10, 12, 13, and 19). It was suggested that depletion of GSH and other antioxidants by PCP metabolites could abolish the protective ability of the cell against ROS and lead to DNA damage (19). It is thus reasonable to hypothesize that PCP metabolite-induced DNA damage could be prevented if the levels of intracellular antioxidants were raised through supplementation of dietary antioxidants. Indeed, it has been demonstrated recently (31) that oral administration of antioxidant vitamin E and diallyl sulfdide 3 h before each PCP challenge significantly protected against elevation of hepatic 8-OH-dG levels in male B6C3F1 mice, while vitamin C, epigallocatechin gallate, and ellagic acid showed partial protection. These findings indicate that PCP-induced oxidative DNA damage in the target organ liver can be blocked by a number of dietary antioxidants.

2. Mechanism of Protection by Desferrioxamine (DFO) against TCHQ-Induced DNA Damage

2.1. Protection by DFO against TCHQ-Induced DNA Damage Was Not Due to Its Binding of Iron but Rather to Its Scavenging of the Reactive TCSQ'. As discussed above, TCHQ has been identified as one of the main toxic metabolites of PCP. TCHQ can induce DNA single strand breaks and has also been implicated in PCP-associated genotoxicity. The ability of TCHQ to induce DNA damage has been previously attributed to its ability to form 'OH through the classic metal-dependent Fenton reaction (see above). This notion was based on the fact that TCHQ-induced DNA damage was completely prevented by DFO (also called desferal and deferoxamine). DFO has been used as an iron chelating agent for the treatment of iron overload. This includes clinical cases of individuals who have ingested toxic oral doses of iron salts or require multiple blood transfusions, such as in the treatment of β-thalassaemia. DFO is a linear trihydroxamic acid siderophore that forms a kinetically and thermodynamically stable complex with ferric iron, ferrirxoamine (Figure 2). Its high binding constant (log β = 31) and its redox properties (E° = −0.45 V) render the bound iron unreactive for the catalysis of oxygen radical production, as has been implicated in a variety of biological processes. It has been classically assessed that the prevention of damage by DFO was a sufficient proof for the role of loosely bound iron in the injurious processes. Although DFO has been repeatedly used to probe metal-catalyzed hydroxyl radical formation in biological processes.
systems, recent studies demonstrated the ability of this trihydroxamate compound to act as a radical scavenger, in addition to and independent of its iron binding properties (for a summary, see ref 32). Diethylenetriaminepentaacetic acid (DTPA) is an analogue of the widely used chelating agent ethylenediamine-tetraacetic acid (EDTA). DTPA could also form a kinetically and thermodynamically stable complex with ferric iron ($\log K = 28; E^0 = + 0.03$ V). Both DFO and DTPA have been widely used to study the role of iron in various chemical and biological systems; therefore, we employed these two structurally different but relatively specific iron chelating agents to probe whether iron played any role in TCHQ-induced DNA damage. We found (32) that DFO protected against TCHQ-induced DNA single strand breaks in isolated DNA, while other iron chelators such as DTPA did not (Figure 3). To better understand its underlying molecular mechanism, the autoxidation process of TCHQ yielding TCSQ intermediate was studied in the presence and absence of these two iron chelating agents. We found that DFO led to a marked reduction in both the concentration and the half-life of TCSQ. Interestingly, the decay of TCSQ was accompanied by the concurrent formation of DFO-nitroxide radicals (DFO$^\cdot$), which contains the structural component $\text{CH}_2-\text{NO}^\cdot-\text{CO}^-$ and gives a characteristic nine-line spectrum as a result of splitting of the nitroxide nitrogen coupling ($\alpha^N = 7.9$ G) by two protons [$\alpha(2)^{1H} = 6.3$ G] from the neighboring CH$_2$ group. These effects have been demonstrated by both UV-visible and electron spin resonance (ESR) spectral methods (Figure 4–7). In contrast, DTPA had no detectable effect on TCHQ autoxidation. These results suggest that the protection by DFO against TCHQ-induced DNA damage was not due to its binding of iron but rather due to its scavenging of the reactive TCSQ$^\cdot$ (32).

Interestingly, we found that DFO could also dramatically enhance the hydrolysis (dechlorination) of TCHQ (and TCBQ) to form chloranilic acid (2,5-dichloro-3,6-dihydroxy-1,4-benzoquinone, DDBQ) (32). The exact underlying molecular mechanism is currently under investigation in our laboratory, and we suspect that a novel Lossen rearrangement reaction might...
be involved. As compared to TCBQ, the DDBQ molecule was considered to be more stable, less reactive, and much less toxic; therefore, the enhanced formation of DDBQ from TCBQ catalyzed by DFO also contributed to its reduction of the toxicity of TCHQ. Furthermore, the enhanced conversion of TCHQ to DDBQ reduced the possibility of redox cycling between TCBQ and TCHQ in the cell, whereby repeatedly producing TCSQ.

2.2. Protection by DFO against TCHQ-Induced Cyto- and Genotoxicity in Human Fibroblasts. Then, we extended our study from isolated DNA to human fibroblasts (cell line GM05757 from the Human Genetic Mutant Repository (Camden, NJ) were used) and from DFO to other hydroxamic acids. Cyto- and genotoxic effects of PCP metabolites were evaluated, respectively, by the MTT and “comet” assay (also called single cell gel electrophoresis). We found (33) that coincubation of DFO provided marked protection against both the cyto- and the genotoxicity induced by TCHQ. Pretreatment of the cells with DFO followed by washing also provided protection, although less efficiently as compared to the simultaneous treatment (Figure 8). Similar patterns of protection were observed for three other hydroxamic acids: aceto-, benzo-, and salicyl-hydroxamic acid (AHA, BHA, and SHA). Spectral studies showed that the three hydroxamic acids tested other than DFO also effectively scavenged the reactive TCSQ and enhanced the formation of the less reactive and less toxic chloranilic acid (Figure 9). Dimethyl sulfoxide (DMSO), an efficient ‘OH scavenger, provided partial protection only at high concentrations. The results of this study demonstrated that the protection provided by DFO and other hydroxamic acids against TCHQ-induced cyto- and genotoxicity in human fibroblasts is mainly through scavenging of the observed reactive TCSQ and not through prevention of the Fenton reaction by the binding of iron in a redox-inactive form (33).

3. Molecular Mechanism of PCP Metabolite-Induced Genotoxicity

3.1. Metal-Independent Production of ‘OH by PCP Metabolite and $H_2O_2$. The above findings suggest that iron was not involved in TCHQ-induced DNA damage. In another word, TCHQ-induced DNA damage may not be due to the iron-mediated ‘OH production through the classic Fenton reaction! Then, the question became what was the underlying molecular mechanism for PCP metabolites-mediated ‘OH production? To test whether ‘OH can be produced by PCP metabolites, we first employed the well-known salicylate hydroxylation method. HPLC with electrochemical detection was used to measure the levels of 2,3- and 2,5-dihydroxybenzoic acid (DHBA) formed when ‘OH reacts with salicylate. We found (34) that TCHQ...
and H$_2$O$_2$ could produce both 2,3- and 2,5-DHBA when incubated with salicylate. Their production was markedly inhibited by the 'OH scavenging agents DMSO and ethanol. In contrast, their production was not affected by the nonhydroxamate iron chelators and the copper-specific chelator. Similar effects were also observed with TCBQ and H$_2$O$_2$. On the basis of these results, we suggested that 'OH was produced by TCHQ and H$_2$O$_2$, possibly through a metal-independent Fenton-like reaction (34).

Because the salicylate hydroxylation method cannot provide direct evidence for 'OH formation, a more specific method, such as secondary radical ESR spin trapping with 5,5-dimethyl-1-pyrroline N-oxide (DMPO), is needed to further substantiate and extend our previous observations. A typical DMPO/'OH signal, and DMPO/CH$_3$ signal derived from 'OH attack on DMSO, will be more conclusive evidence for 'OH production from H$_2$O$_2$ and TCHQ or TCBQ (14) (Figure 10). We found (35) that when incubated with DMPO, TCBQ and H$_2$O$_2$ produced the DMPO/'OH adduct. In contrast, incubation of either compound alone did not cause 'OH formation (Figure 11). The formation of DMPO/'OH was markedly inhibited by the 'OH scavenging agents DMSO and formate, with the concomitant formation of the characteristic DMPO adducts with 'CH$_3$ and 'COO', respectively (Figure 12). These secondary radical ESR spin-trapping results provided definitive evidence that 'OH could indeed be produced by TCBQ and H$_2$O$_2$.

Then, a critical question arose: Was the production of 'OH by TCBQ and H$_2$O$_2$ metal-dependent or -independent? To answer this question, the potential role of catalytic transition metals contaminating the DMPO/TCBQ/H$_2$O$_2$ reaction system was carefully examined by using several structurally different and relatively specific metal chelating agents for iron and copper (36–38). Neither the DMPO/'OH signal nor the DMPO/CH$_3$ signal produced by the DMPO/TCBQ/H$_2$O$_2$ system in the absence and presence, respectively, of DMSO was affected by the addition of various nonhydroxamate iron chelating agents, viz., bathophenanthroline disulfonate (BPS), ferrozine, and ferene, as well as the copper-specific chelating agent bathocuproine disulfonate (BCS) (Figure 13). In addition, no significant decrease in the DMPO/'OH and DMPO/CH$_3$ signal was observed by low concentrations (≤10 µM) of the trihydroxamate iron chelating agent DFO. These DFO concentrations should be sufficient to chelate any trace amounts of iron contaminating the chelated-pretreated buffer. However, the formation of DMPO/'OH and DMPO/CH$_3$ was abolished by high concentrations of DFO (≥1 mM), with the concurrent formation of the DFO·. As discussed before, the inhibition of 'OH production by DFO was not due to its iron-binding capacity but rather due to its ability to scavenge TCSQ·. Similar marked inhibition of DMPO/'OH and DMPO·/CH$_3$ formation was observed with another TCSQ· scavenger, benzohydroxamic acid. In addition, even when trace amounts of iron [Fe(II), 0.5 µM] were added to the DMPO/TCBQ/H$_2$O$_2$ system, no increase in 'OH production was observed.

In contrast, the formation of both DMPO/'OH and DMPO·/CH$_3$ by the DMPO/Fe(II)/H$_2$O$_2$ system in the absence and presence, respectively, of DMSO was almost completely inhibited by the nonhydroxamate iron chelating agents BPS, ferrozine, and ferene, as well as the hydroxamate iron chelating agent DFO. No concurrent formation of the DFO· was detected, indicating that DFO acted by chelating iron in this classic Fenton system. These results clearly demonstrated that the production of 'OH by TCBQ and H$_2$O$_2$ is independent of transition metal ions.

It should be noted that the metal-independent production of 'OH was not limited to TCBQ and H$_2$O$_2$ but was also observed in the presence of other halogenated quinones, that is, 2-chloro-, 2,5-dichloro-, 2,6-dichloro-, trichloro-, tetrachloro-, and tetrabromo-1,4-benzoquinone. In contrast, no 'OH production was detected from H$_2$O$_2$ and the nonhalogenated quinone, 1,4-benzoquinone, and the methyl-substituted quinones 2,6-dimethyl- and tetramethyl-1,4-benzoquinone (35).

3.2. Molecular Mechanism of Metal-Independent Production of 'OH by PCP Metabolites and H$_2$O$_2$. On the basis of the above experimental results, we first proposed (34, 35) that the production of 'OH by TCBQ and H$_2$O$_2$ might be through a metal-independent semiquinone-mediated organic Fenton reaction:

\[
\text{TCSQ}^+ + \text{H}_2\text{O}_2 \rightarrow \text{'OH} + \text{OH}^+ + \text{TCBQ}
\]

where TCSQ· substitutes for ferrous iron in the classic, metal-dependent Fenton reaction. This type of reaction between semiquinone radicals and H$_2$O$_2$ has been previously proposed by Koppenol and Butler (39), who suggested that if a quinone/semiquinone couple has a reduction potential of between −330 and +460 mV, it can theoretically bring about a metal-independent Fenton reaction. It was suggested that such reactions are thermodynamically feasible and do not require metal ions for catalysis (39, 40), which might be the case in this study, where the reduction potentials of the quinone/semiquinone couples for 2-chloro-, 2,5-dichloro-, tetrachloro-, tetrabromo-, and tetrachloro-1,4-benzoquinone are −100, +60, +200, +240, and +250 mV, respectively (41). These values are within the suggested range of −330 to +460 mV. In contrast, the reduction potentials for 2,6-dimethyl-1,4-benzoquinone and tetramethyl-1,4-benzoquinone of −430 and −600 mV, respectively (41), are outside this range, and indeed, no 'OH formation could be detected.

If the above mechanism was correct, then the production of 'OH from H$_2$O$_2$ and TCBQ should be dependent on the concentration of TCSQ·; that is, the higher the concentration of TCSQ·, the more 'OH should be produced. Furthermore, the main product of this reaction should be TCBQ. Using secondary radical ESR spin-trapping method, we found that DMPO/CH$_3$ and DMPO/'OH adducts can be produced by H$_2$O$_2$ and TCBQ in the presence of the spin-trapping agent DMPO and 'OH scavenger DMSO. However, no DMPO/CH$_3$ and DMPO/'OH adducts were detected from H$_2$O$_2$ and TCHQ (the reduced form of TCBQ), although high concentrations of TCSQ· could be produced during the autoxidation of TCHQ. Interestingly, if
TCHQ was quickly oxidized to TCBQ with myeloperoxidase (MPO), DMPO/CH₃ and DMPO/OH adducts could be detected again, similar to that produced by TCBQ (Figure 14). Furthermore, the formation of DMPO/CH₃ and DMPO/OH was found to be directly dependent on the concentrations of TCBQ and H₂O₂. These results strongly suggest that TCBQ, but not its corresponding semiquinone radical TCSQ, is essential for OH production. Therefore, the production of 'OH by TCBQ and H₂O₂ appears not to occur through a semiquinone-dependent organic Fenton reaction.

To get more information on the mechanism of 'OH production by TCBQ/H₂O₂, the time- and concentration-dependent production of DMPO/OH by TCBQ/H₂O₂ was studied. Two distinct phases were observed as follows: the first fast phase (about 30 s) and the second slower phase. This indicates that 'OH may be produced by two-step reactions between TCBQ and H₂O₂. When the TCBQ concentration was fixed at 0.1 mM, the rate of DMPO/OH production was dependent on the H₂O₂ concentration. It should be noted that DMPO/OH could be detected at H₂O₂ concentration as low as 10 µM. When the H₂O₂ concentration was fixed at 0.1 mM, DMPO/OH could be detected at a TCBQ concentration as low as 5 µM. Furthermore, UV-visible spectral studies showed that there was a direct interaction between TCBQ and H₂O₂, with the reaction mixture changing quickly from the original yellow color (λ_max = 292 nm) to a characteristic purple color (λ_max = 295 and 535 nm) in phosphate buffer (pH 7.4). The final reaction products between TCBQ and H₂O₂ were then identified by electrospray ionization quadrupole time-of-flight mass spectrometry (ESI-Q-TOF-MS). The mass spectrum of TCBQ is characterized by a four-chlorine isotope cluster at m/z 246 and traces of a three-chlorine isotope cluster at m/z 227. Tandem mass spectrometric analysis showed that the peak at m/z 227 could be fragmented to form the peak at m/z 197, which suggests that the peak at m/z 197 is solely derived from the peak at m/z 227 (Figure 15). These results indicate that the major reaction product between TCBQ and H₂O₂ was probably the ionic form of trichloro-hydroxy-1,4-benzoquinone (TrCBQ-OH). This was further confirmed by comparison with the authentic TrCBQ-OH synthesized according to a published method (42), which showed the same ESI-MS profile and the same retention time in HPLC.

To better understand the source and origin of the oxygen atom inserted into the reaction product TrCBQ-OH formed from the reaction between H₂O₂ and TCBQ, TCBQ was incubated with oxygen-18-enriched H₂O₂ ([18O]-H₂O₂). The mass spectra of the molecular ion region of deprotonated TrCBQ-OH, obtained with unlabeled and labeled H₂O₂, demonstrated the shift of the molecular ion isotope cluster peaks of the unlabeled compound with 2 mass units, as could be expected for the incorporation of ¹⁸O. These results indicate that H₂O₂ is the source and origin of the oxygen atom inserted into the reaction product TrCBQ-OH.

It has also been shown (43) that both TCBQ and H₂O₂ were consumed with a stoichiometric ratio of about 1:1, and H₂O₂ accelerated the rate of TCBQ decomposition by 2 orders of magnitude with the loss of chloride. Thus, the metal-independent production of 'OH by TCBQ and H₂O₂ may not be through a previously proposed semiquinone-mediated organic Fenton reaction. On the basis of the above experimental results and the fact that H₂O₂ is a better nucleophile than H₂O (44), we proposed a novel mechanism for 'OH production by H₂O₂ and TCBQ (45) (Figure 16): A nucleophilic reaction may take place between TCBQ and H₂O₂, forming an unstable trichloro-
Figure 16. Proposed novel mechanism for 'OH production by TCBQ and H₂O₂ (modified based on ref 45).

These findings represent a novel mechanism of 'OH and alkoxyl radical formation not requiring the involvement of redox-active transition metal ions and may partly explain the potential carcinogenicity of not only PCP but also other widely used polyhalogenated aromatic compounds such as 2,4,6- and 2,4,5-trichlorophenol, hexachlorobenzene, Agent Orange (the mixture of 2,4,5-trichlorophenoxyacetic acid (2,4,5-T) and 2,4-dichlorophenoxyacetic acid (2,4-D)), and the brominated flame retardant 3,3′,5,5′-tetrabromobisphenol A (TBBPA), since these compounds can be metabolized in vivo (47, 48) or dechlorinated chemically (49–52) to tetra-, di-, or monohalomethylated quinones. Our data suggest that TCBQ and other halogenated quinones may react with hydroperoxides and exert toxic effects through enhanced production of hydroxyl/alkoxyl radicals and hence increased DNA, protein, and lipid oxidation.

It should be noted, however, that many questions still need to be addressed, especially regarding the biological relevance of the reactions under study. For example: What is the rate constant for the reaction between TCBQ and H₂O₂ and could it be able to compete with the classic Fenton reaction? How does the reaction of the quinone with H₂O₂ compare kinetically with reactions with other good nucleophiles such as glutathione and other thiols, which are present at high concentrations in vivo? How rapid is the decomposition of the quinone-peroxide adduct? How efficient is the radical formation from a given amount of adduct; that is, how well do other pathways compete with this reaction? Could 'OH produced by this pathway be detected in cell culture or even in an animal model? Therefore, further investigations are needed to study whether these reactions occur and are relevant under physiological conditions or in vivo.

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