Lipid Metabolites of Carbon Tetrachloride

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Abstract 5 min after intravenous injection into rats of 14C- or 35Cl-carbon tetrachloride, liver lipids were found labeled. Most of the radioactivity was found in the phospholipid fraction. The metabolites were shown to comprise a heterogeneous group of branched long-chain chlorinated fatty acids, probably containing the trichloromethyl side chain. Surviving liver slices also formed these metabolites. In a simple chemical system which generates trichloromethyl free radicals, carbon tetrachloride added to methyl oleate to form esters which behaved like the metabolites during counter-current distribution and urea adduction. The evidence strongly suggests the formation of these metabolites by free radical attack on unsaturated lipids. The relation of these observations to current theories of carbon tetrachloride intoxication is discussed.

Introduction
Poisoning by carbon tetrachloride has often been thought to result from its ability as a lipid solvent to destroy lipid-rich membranes of cells and subcellular particles. This view is probably incorrect. It fails to explain the variability in toxic dose, the alteration in susceptibility to poisoning produced by antioxidants (1, 2), and the greater toxicity of carbon tetrachloride than chloroform (3). Most striking is Slater's demonstration (4) that in equimolar doses, fluorotrichloromethane, a liquid similar to CCl4 in physical properties and solvent power, is not hepatotoxic at all. Other theories of CCl4 poisoning which attribute its effects to the unchanged CCl4 molecule (e.g., stimulation of catechol release (5), interference with blood flow to the centers of liver lobules (6)) are not supported by recent evidence (7, 8). Much of this evidence is discussed in Recknagel's comprehensive review (9).

Recent efforts have therefore been concentrated on the relation between the metabolism of CCl4 and its toxicity. In 1951, McCollister, Beamer, Atchison, and Spencer (10) showed that while most 14C-CCl4 breathed by monkeys was exhaled unchanged, a small fraction was converted to radioactive carbon dioxide, chloroform, urea, and one or more unidentified urinary organic anions. This group of products did not in itself suggest a toxic mechanism. In 1961, Butler (11) showed that the in vitro reaction of CCl4 with sulfhydryl compounds led to replacement of sulfhydryl hydrogen. He inferred that the reaction mechanism involved the homolytic cleavage of CCl4 into CCl3 radicals and free radicals with radical attack on the susceptible functional groups. In Butler's experiments, sulfhydryl groups were oxidized to disulfide with formation of chloroform. He suggested that in other circumstances, the trichloromethyl radical itself might replace the sulfhydryl hydrogen. Free radical attack on sulfhydryl groups might take place in vivo, but no such products have been identified yet in intoxicated animals. Carbon tetrachloride may poison by means of two other well characterized reactions of free radicals: the addition of halomethanes to olefins (12), and the chain peroxidation of dienes and polyenes (13). In vitro these reactions are best initiated by heat, ultraviolet light, or catalytic amounts of organic peroxides. In the case of addition to olefins, trichloromethyl radicals add to one carbon of the double bond, and depending on the reaction conditions, either Cl or H adds to the other. In the case of chain peroxidation of dienes and polyenes, an initiator, which can be the CCl3 radical, catalyzes the formation of peroxide radicals, but this reaction is self-perpetuating and proceeds without further catalyst. In both cases polymerization can result from one or more couplings of product radicals. Unsaturated lipids are obvious substrates for both free radical reactions.

Reynolds (14) reported that rat liver lipids were labeled after feeding either 14C- or 35Cl-carbon tetrachloride. Microsomal lipids were most heavily labeled. The studies described below demonstrate the labeling of liver lipids within 5 min after the intravenous injection of 14C- and 35Cl-carbon tetrachloride, confirming the observations of Reynolds. The metabolites comprise a heterogeneous group of branched chlorinated long-chain fatty acids. Surviving liver slices also formed these metabolites. These products share the properties of lipids synthesized by benzoyl peroxide-catalyzed addi-
tion of carbon tetrachloride to unsaturated esters. The structure of the metabolites strongly suggests their formation through attack by trichloromethyl radicals on unsaturated lipids.

**METHODS**

Unlabeled carbon tetrachloride, certified ACS grade, was obtained from Fisher Scientific Company, Pittsburgh, Pa., *14C*carbon tetrachloride, 7.2 mc/mm, was obtained from Nuclear-Chicago Corporation, Des Plains, III., and *36Cl*-carbon tetrachloride, 0.35 mc/mm, from the Radiochemical Center, Amersham, England. Methyl oleate, 99% pure, and ampoules of BF₃-methanol, were obtained from Applied Science Laboratories, Inc., State College, Pa., and Lipomul fat emulsion for intravenous injection from the Upjohn Co., Kalamazoo, Mich. Silicagel G was purchased from Merck, Darmstadt, Germany, and silicic acid, 100 mesh, A.R., from Mallinkrodt. Liquiflor, a concentrated solution of 2,5-diphenyloxazole (POPOP) and 2-p-phenylenelbis(5-phenyloxazole) (POPOP) in toluene, were obtained from New England Nuclear Corp., Boston, Mass. A 25-fold dilution in toluene provides a solution of POPOP 4 g/liter, and POPOP 50 mg/liter.

Solvents were redistilled before use, unless otherwise noted.

Radioactivities of samples containing *14C* or *36Cl* were assayed in the double channel Nuclear-Chicago model 720 scintillation counter. Discriminator settings were chosen so that the isotopes could be assayed simultaneously.

Radioactive carbon tetrachloride was administered intravenously to rats without additional unlabeled carbon tetrachloride. The tracer was mixed with Lipomul, yielding a stable aqueous radioactive preparation, since the tracer dissolved in the fat particles of the emulsion. *14C*-carbon tetrachloride was supplied in a vacuum ampoule. The top of the ampoule was broken under a layer of Lipomul, permitting the radioactive vapor to mix rapidly with the emulsion. *36Cl*-carbon tetrachloride, of lower specific activity, was supplied as a liquid, and was added to the Lipomul with a Hamilton microsyringe. To assay the radioactivity of the Lipomul-CCl₄ mixture, we distributed 5 μl in the two phase system, heptane (4 ml)-isopropanol (4 ml)-water (3 ml), and counted an aliquot of the upper (heptane) phase.

None of the fat in Lipomul was labeled by mixing with carbon tetrachloride. An alternative intravenous preparation, CC₄-propylene glycol-water, yielded identical liver metabolites, but this system incorporated less tracer and introduced a glycol, so that it was abandoned in favor of Lipomul.

To obtain liver lipids, we homogenized livers or liver slices in chloroform-methanol 2:1 (v/v). Homogenates were filtered, the filtrates washed with 0.2 volume of water, and the lipids recovered by evaporating the chloroform and resolvling the residue. "Phospholipids" refers to lipids precipitated in the cold by adding 9 volumes of acetone to a chloroform solution of total lipid extract.

Glycerol esters were hydrolyzed by refluxing in 4 N HCl for 3 hr. The reaction mixture was cooled and distributed in about 30 volumes of the two phase system petroleum ether-isopropanol-water (4:4:3 v/v). Fatty acids were recovered from the upper phase by evaporating the solvent in a rotary evaporator. Methyl esters were prepared from the free fatty acids by heating the acids 1 min at 100°C in BF₃-methanol. Lipids were not saponified in alkaline alcohols, since these reagents dehalogenate.

Phospholipids were fractionated by chromatography on silicic acid according to the procedure of Hanahan, Dittmer, and Warashina (15). The eluting solvents were evaporated in a rotary evaporator and the residues dissolved in petroleum ether before assay of lipid phosphorus (16) and *36Cl* radioactivity.

Countercurrent distributions of fatty acids and of fatty acid methyl esters were performed using the solvent systems of Ahrens and Craig (17). For fatty acids, the solvent phases were prepared by mixing heptane, 97.5% acetic acid, methanol and acetonitrile (4:1:1:1 v/v) and allowing phase separation. For methyl esters, the phases were obtained after equilibration of equal volumes of heptane and acetonitrile. Radioactivities were assayed after countercurrent distribution by evaporating an aliquot of each upper phase in a counting vial and resolvling the residue in the scintillation solution.

Branched-chain compounds were distinguished from straight-chain compounds by the formation of urea adducts (18, 19). When urea and a straight-chain hydrocarbon are dissolved together in methanol, inclusion complexes precipitate spontaneously. These complexes consist of hollow cylindrical crystals of urea in which the hydrocarbon chain is included. Branching of the chain tends to prevent its inclusion in the urea complex, and branched compounds remain mostly in the supernatant. Fatty acid methyl esters were dissolved in hot 15% methanolic urea, and the adducts precipitated spontaneously overnight at room temperature. The complexed esters and the excluded esters were recovered separately by distributing the precipitate and the supernatant in the system petroleum ether-methanol-water.

Thin-layer chromatography on Silicagel G impregnated with silver nitrate was used to distinguish saturated from unsaturated esters. Plates 700 μ thick were prepared by a modification of the method of Kaufmann and Wessels (20).

**RESULTS**

**In vivo studies**

As early as 5 min after intravenous injection of radioactive carbon tetrachloride, rat liver lipids were labeled. The analyses described below were done on lipids obtained 15 min after injection. Rats were given either *36Cl*-carbon tetrachloride or "doubly labeled" carbon tetrachloride, a mixture of *14C* and *36Cl*-CCl₄. The use of carbon tetrachloride labeled only with *36Cl* permitted simultaneous comparison of chlorinated metabolites with standard *14C*-palmitic acid and its methyl ester during countercurrent distribution and urea adduction. Use of doubly labeled CCl₄ permitted calculation of the number of chloride atoms per molecule of CCl₄ incorporated into lipid.

**Liver Lipids after Injection of *36Cl*-Carbon Tetrachloride**

Male rats of the Sprague-Dawley strain weighing 125-150 gm were fasted overnight. They were each injected intravenously with 1 ml Lipomul containing 8 × 10⁶ cpm *36Cl*-carbon tetrachloride. 15 min later the rats were decapitated, and the liver lipids were extracted. The lipid extracts contained about 0.5% of the injected radioactivity. The extracts were pooled, and the phospholipids
precipitated with acetone. 65% of the radioactivity was found in the acetone-precipitable lipids. None of the activity in the acetone precipitate was due to Cl-chloride ion, since shaking a wet solution of the phospholipids with powdered silver carbonate did not reduce the lipid radioactivity.

**Distribution of radioactivity in phospholipid fractions.** To determine if any class of phospholipid was preferentially labeled, we fractionated a portion of the acetone-precipitable lipid on a silicic acid column (15). Table I presents the elution sequence, and the lipid phosphorus and Cl activity of each fraction. From the finding that all of the phospholipid fractions were labeled, as well as the labeling of acetone-soluble lipids, it appeared probable that the fatty acids were labeled. The remainder of the phospholipid was hydrolyzed, and the fatty acids recovered. To this fatty acid mixture, C-palmitic acid was added in an amount chosen to make the C and Cl radioactivities similar. 2/3 of this fatty acid mixture was then esterified in BF3-methanol.

**Countercurrent distribution of labeled fatty acids and methyl esters.** Phospholipid fatty acids containing Cl-acids and added C-palmitic acid were distributed through twenty transfers in the system heptane-acetonitrile-acetic acid-methanol, lower phase 10 ml, upper phase 6.8 ml. 5-ml aliquots of each upper phase were evaporated in counting vials and the residues dissolved in scintillation fluid for assay of radioactivity. Fig. 1 shows the distribution of the free acids. Fig. 2 shows the distribution of the methyl esters (heptane-acetonitrile, 10 ml upper phase, 10 ml lower phase, 8 ml of upper phases assayed).

The labeled metabolites thus behave as a mixture of long-chain acids. To show that the chlorinated lipid is not a single substance, consider the simultaneous countercurrent distribution of two pure substances, A and B, with partition coefficients K and K. Call the concentration of A in the rth tube Ar, and of B in the rth tube Br. It can be readily shown that if any two adjacent tubes are compared, say the rth and the (r + 1)th,

\[
\frac{A_{r+1}/A_r}{B_{r+1}/B_r} = K/K.
\]

Since the right side is constant, so must be the left. Calculation from the data used to plot the curves in Figs. 1 and 2 shows that this ratio varies with successive pairs of adjacent tubes. The chlorinated lipid is therefore heterogeneous.

**Urea adduction.** A portion of the methyl ester mixture containing Cl-esters and C-methyl palmitate was retained from the last 10 tubes of the countercurrent distribution. Solvents were evaporated and the esters dissolved in hot 15% urea in methanol. Urea adducts were allowed to precipitate overnight at 25°C. Adducted esters in the precipitate and excluded esters in the supernatant were recovered. Table II shows the isotope ratio in the original mixture, in the adduct, and in the supernatant. The chlorinated metabolites are clearly mainly branched-chain lipids.

![Graph](image-url)

**TABLE I**

<table>
<thead>
<tr>
<th>Fraction No.</th>
<th>Eluting solvent</th>
<th>Total eluted lipid P</th>
<th>Total eluted lipid</th>
<th><strong>Cl</strong></th>
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<tbody>
<tr>
<td>I</td>
<td>CHCl3-CH3OH 7:1</td>
<td>1.2</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>II</td>
<td>&quot;</td>
<td>4:1</td>
<td>38.0</td>
<td>26</td>
</tr>
<tr>
<td>III</td>
<td>&quot;</td>
<td>3:2</td>
<td>55.0</td>
<td>48</td>
</tr>
<tr>
<td>IV</td>
<td>&quot;</td>
<td>1:4</td>
<td>5.7</td>
<td>17</td>
</tr>
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</table>

![Graph](image-url)

**Figure 1** Countercurrent distribution of Cl-fatty acid metabolites (in vivo labeling) together with added standard C-palmitate.

**Figure 2** Countercurrent distribution of methyl esters of Cl-fatty acid metabolites (in vivo labeling) together with added standard methyl C-palmitate.
Acetone-soluble liver lipids were likewise hydrolyzed, then analyzed by countercurrent distribution and urea adduction. Again, the radioactivity was found incorporated into branched, long-chain fatty acids.

**Liver Lipids after the Injection of Doubly Labeled CCl₄**

Fasted rats were injected with 0.5 or 1 ml of Lipomul containing a mixture of ¹⁴C- and ³⁵Cl-carbon tetrachloride. At 15 min, approximately 0.1% of the injected radioactivity was found incorporated in liver lipids. 68% of the lipid radioactivity was found in the acetone-precipitable lipids. Fatty acid methyl esters were prepared from the acetone-precipitable lipid for countercurrent distribution.

Fig. 3 shows the countercurrent distribution of doubly labeled methyl esters. Above the distribution curve is a plot of the ³⁵Cl/¹⁴C ratio in the lipid of each tube divided by the ratio in the injected dose, the quotient multiplied by four. It is seen that the predominant route to the labeled lipid involves the incorporation of three of the four chlorine atoms from each molecule of CCl₄, but some lipid is formed using all the CCl₄ chlorine atoms.

Formation of doubly labeled lipids by liver slices

Slices weighing 100–150 mg were taken from livers of starved rats. Two slices per flask were incubated at 37°C in 3.5 ml Krebs-Ringer bicarbonate buffer to which 0.5 ml Lipomul containing “doubly-labeled” CCl₄ was added. Slices and buffer in two control flasks were heated 1 min in a water bath at 100°C before the Lipomul was added. Incubations were stopped after 5, 15, 30, and 60 min. Control flasks were incubated 60 min. After incubation, slices were removed with forceps, rinsed copiously with saline, and the lipids were extracted. Table III shows the increasing incorporation of radioactivity into slice lipids with time in the surviving slices. The extracts were then pooled, fatty acid methyl esters again prepared from the acetone-precipitable lipid, and distributed in the countercurrent system heptane-acetonitrile. Fig. 4 shows that the labeled lipids behave similarly to those obtained in vivo.

**Free radical addition of carbon tetrachloride to methyl oleate**

Methyl oleate (250 mg), carbon tetrachloride (3.5 ml) containing ¹³C- and ³⁵Cl-tracer, and dibenzoyl peroxide (7 mg) were heated together under argon in a sealed ampoule 16 hr at 85°C. The ampoule was then heated 1 additional hr at 110°C to destroy the peroxide. After cooling, the reaction mixture was washed first with 10%

<table>
<thead>
<tr>
<th>Table II</th>
<th>Comparison of Urea Adduction of ³⁵Cl-Fatty Acid Methyl Esters with ¹³C-Methyl Palmitate</th>
</tr>
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<tbody>
<tr>
<td>Lipid mixture</td>
<td>³⁵Cl</td>
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<tr>
<td>Starting mixture</td>
<td>2730</td>
</tr>
<tr>
<td>Precipitate (urea adduct)</td>
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<td>Supernatant (excluded esters)</td>
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<table>
<thead>
<tr>
<th>Table III</th>
<th>Incorporation of ³⁵Cl Radioactivity into Lipid by Rat Liver Slices after Incubation in Radioactive CCl₄</th>
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<tbody>
<tr>
<td>Time (min)</td>
<td>cpm/mg wet tissue</td>
</tr>
<tr>
<td>5</td>
<td>158</td>
</tr>
<tr>
<td>15</td>
<td>326</td>
</tr>
<tr>
<td>30</td>
<td>480</td>
</tr>
<tr>
<td>60</td>
<td>1114</td>
</tr>
<tr>
<td>60 (control slices)</td>
<td>46</td>
</tr>
<tr>
<td>60 (control slices)</td>
<td>44</td>
</tr>
</tbody>
</table>

**Figure 3** Countercurrent distribution of methyl esters of doubly labeled fatty acid metabolites (in vivo labeling).

**Figure 4** Countercurrent distribution of methyl esters of doubly labeled fatty acid metabolites (in vitro labeling).
A portion of the lipid residue was analyzed by countercurrent distribution. Fig. 5 shows the results. The radioactivity was incorporated into lipid.

Another portion of the lipid in the final reaction mixture was dissolved in hot 15% urea in methanol, and the adducts were allowed to form overnight at 25°C. Samples of the original methyl oleate, of the lipids in the ampoule at the end of the reaction, of the adducted esters, and of the excluded esters were analyzed by thin-layer chromatography on AgNO₃-silicic acid. The results (Fig. 6) demonstrated that a saturated product was produced, and that this product was branched. On a second AgNO₃-silicic acid plate, about 50 mg of the final reaction mixture was applied to the origin as a band, and the chromatography repeated. Bands corresponding to the saturated esters and the monoenes were scraped from the plate with a razor blade, and the esters eluted...
from the scrapings with ethyl ether. Radioactivities of both fractions were assayed. 97% of the radioactivity was found in the saturated esters.

**DISCUSSION**

These studies confirm Reynolds' observations that liver lipids are labeled both by \(^{14}C\) and \(^{35}Cl\)-carbon tetrachloride. More than half the labeling occurs in the phospholipid fraction. The studies reported here show that the new compounds are a heterogeneous group of branched long-chain fatty acids containing both carbon and chlorine. Surviving liver slices can also form these metabolites. In a simple chemical system which generates trichloromethyl free radicals, the addition of carbon tetrachloride to methyl oleate produced a saturated product which behaved like the metabolites during urea adduction and countercurrent distribution. The evidence suggests that the free radical mechanism is the route to the metabolites formed in vivo.

Any theory which relates the toxicity of carbon tetrachloride to its metabolism must satisfy at least two requirements: (a) it must explain the decreasing order of toxicity of CCI\(_4\), chloroform, and CCl\(_3\)F, and (b) the metabolic transformation should occur at least as soon as the morphologic change and should help explain it. Of the three halomethanes, carbon tetrachloride forms free radicals most easily, CCl\(_3\)F least (21). The rapidity and locus of labeling of liver lipid is compatible with the rate of onset and locus of the morphological changes after ingestion of a toxic dose of CCl\(_4\). The first effect on the liver demonstrable by electron microscopy is the disruption and vesiculation of the cisternae of the coarse endoplasmic reticulum and the dispersion of ribosomal particles (22, 23). At this stage these ribosomes are severely incapacitated in protein-synthesizing ability (24). Presumably the disruption of the ordered structure of the endoplasmic reticulum could result from damage to its lipid membranes.

Recknagel has proposed that the CCl\(_4\)· radicals initiate destructive chain peroxidation of unsaturated lipids (25). The active methylene bridges of dienes and polyene fatty acids are excellent targets for radical attack (13). Recknagel found that peroxides could not be directly identified in vivo, since their identification depends on determination of malonaldehyde, itself metabolized by liver. However, he was able to show increased diene conjugation after CCl\(_4\) administration, a reaction which accompanies lipid peroxidation. An alternative role for the radical reaction is the formation of chlorinated lipids, which may be unsuitable as membrane components. Cross-linking and polymerization of lipids containing unsaturated fatty acids might also result from free radical attack, along with peroxidation or chlorination. The protective effect of certain antioxidants against CCl\(_4\) poisoning (1, 2) is compatible with the radical mechanism, since these compounds can function as free radical inhibitors. Of course, evidence for a metabolic pathway is not proof that it is responsible for toxicity. The existence of the free radical mechanism seems well established. Whether any of its possible consequences account for the toxicity of CCl\(_4\) remains to be proven.

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**REFERENCES**


