Comparative Mammalian Metabolism of Vinyl Chloride and Vinylidene Chloride in Relation to Oncogenic Potential

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Elucidation of the role of vinyl chloride metabolites in the various reaction sequences which comprise the metabolic pathway, including the interaction of reactive metabolites with some purine and pyrimidine residues of target-organ DNA, provides some explanation for the (oncogenic) properties associated with the original substance. Comparative investigation of the biological fate of vinylidene chloride reveals an agent of low oncogenic potential which is likely to be damaging only under special circumstances, and species differences which suggest that the mouse is more susceptible than the rat towards vinylidene chloride oncogenicity.

The research work with which this communication is concerned is based on the idea that knowledge of the biology of the reactive metabolites of chemical carcinogens in the mammal, including the precise nature of the chemical changes to the DNA of the nucleus, ought to give an insight into the (oncogenic) properties of the parent compounds.

In tracer studies, N-acetyl-S-(2-hydroxymethyl)cysteine was shown to be a major vinyl chloride metabolite in rats, but according to the method of protective esterification that was used so a derivative either of N-acetyl-S-(2-chloroethyl)cysteine or of N-acetyl-S-(2-hydroxyethyl)cysteine was isolated from body fluids (1, 2). Thus, by Fischer-Speier methylation, N-acetyl-S-(2-chloroethyl)cysteine was obtained, and with diazomethane, N-acetyl-S-(2-hydroxyethyl)cysteine. It might be stated in passing that throughout the investigations described, mass spectrometry, involving electron impact (EI) and chemical ionization sources and multiple-ion detection and all combinations of these facilities, was used extensively both for product identification and analysis and for the purposes of detection. Treatment of the O-methyl ester of N-acetyl-S-(2-hydroxyethyl)cysteine (a) with the methanol–HCl reagent gave a mixture of N-acetyl-S-(2-chloroethyl)cysteine (b), and S-(2-chloroethyl)cysteine, and conversely, the O-methyl ester of N-acetyl-S-(2-chloroethyl)cysteine (b) was hydrolyzed rapidly by water to that of N-acetyl-S-(2-hydroxyethyl)cysteine (a) (2). Hence, the reversible reaction processes connecting the two substances would seem to be modulated through the intermediacy of episulfonium ion (c) and formation of this ion would in fact be rate-limiting in respect of the hydrolysis of...
N-acetyl-S-(2-chloroethyl)cysteine. Nucleophilic attack of OH⁻ on the episulfonium ion would be expected to give olefin (3), and in fact, N-acetyl-S-vinylcysteine (d) (2) was recovered from the urine of vinyl chloride-treated animals whenever diazomethane esterification was used to protect S-containing metabolites.

Surprisingly, N-acetyl-S-(2-hydroxyethyl)cysteine O-methyl ester was methylated with neutral methanol, and the O-methyl esters of N-acetyl-S-(2-methoxy[¹⁴C]ethyl)cysteine plus N-acetyl-S-[¹⁴C]vinyl-cysteine degrade to give the volatile [¹⁴C]S-(2-methoxyethyl) (prop-1 or 2-enyl) sulfide. Although the mechanism of formation was not investigated, we felt that acetaldehyde, a known dissociation product of S-vinylcysteine-derived S-vinylcysteine-S-oxide (4) might undergo concerted condensation with N-acetyl-S-(2-methoxyethyl)cysteine leading to elimination of thermodynamically stable glyoxylate. [There is an analogy for such a concerted condensation reaction in the work of Dabritz and Virtanen (4) on the tear-producing volatile components of the onion.]

The half-mustard S-containing metabolites of vinyl chloride did not behave as mutagens in the Ames test (2).

Thiodiglycollic acid is another major vinyl chloride metabolite (1).

In order to determine whether vinyl chloride yielded chloroacethylene oxide in vivo, the biogenesis of several vinyl chloride metabolites and related compounds were investigated in rats (2). S-(2-Hydroxyethyl)cysteine gave 0.5% of the authentic thiodiglycollic acid, and this result was seen to be highly significant, because of the instability (v. supra) of the starting material under exceedingly mild conditions of reaction. The metabolic pathway concerned [Eq. (1)] appears to include endgroup oxidation (I), amino-acid transamination (II), and oxidative decarboxylation (III), and the results of the animal feeding experiments suggest that chloroacetaldehyde (g) chloroacetic acid (h), and S-(2-carboxymethyl)cysteine (i) might lie on a common pathway connecting vinyl chloride (e) with thiodiglycollic acid (j). However, other evidence implies that chloroacetic acid (h) does not belong to this metabolic pathway (e-j). Thus, < 0.1% has even been detected in the body fluids of any of our vinyl chloride-treated animals. Either there is a high rate of turn-over or this compound is not a major vinyl chloride metabolite. The latter possibility seems more likely, since relatively large amounts are produced in vinylidene chloride metabolism, and in those animals, thiodiglycollic acid accounts for an even greater proportion of the dose than in parallel experiments with vinyl chloride. A feasible metabolic pathway for thiodiglycollic acid from chloroacetic acid and involving cysteine desulphhydrase is unacceptable. Experiments with unlabeled vinyl chloride in rats in which the cysteine-cystine pools had been labeled adequately with [¹⁴C] gave [¹⁴C]thiodiglycollic acid, showing that a part of the C-skeleton must be derived in fact from cysteine. In rats treated with chloroacetaldehyde, the presence of thiodiglycollic acid and N-acetyl-S-(2-hydroxyethyl)cysteine, but not of chloroacetic acid, among the urinary metabolites was established by mass fragmentometry.

Thus, it is probable that in vivo chloroethylene...
oxide (f) was formed (5) from vinyl chloride (e) and transformed spontaneously (6) into chloroacetaldehyde (g); there is supporting evidence (7–10) for vinyl chloride epoxidation in vitro. This supposition is supported by the facts that chloroacetaldehyde affords both N-acetyl-S-(2-hydroxyethyl)cysteine and thiodiglycolic acid in vivo and that S-(2-carboxymethyl) cysteine has been identified by mass fragmentometry amongst the hydrolytic products of an hepatic extract prepared from vinyl chloride-treated animals. Since chloroacetaldehyde and chloroethylene oxide are mutagenic in the Ames test (11–13) and in Chinese hamster V79 cells (14), they may be relevant to vinyl chloride carcinogenicity.

Respective formation of 9β-D-2′-deoxy ribofuranosylimidazo-[2,1-i]purine or 3β-D-2′-deoxy ribofuranosyl-2-oxo-2,3-dihydroimidazo-[1,2-c]pyrimidine from deoxy adenosine or deoxy cytidine by reaction with chloroacetaldehyde (15) or chloroethylene oxide was readily confirmed. Recognition of the nucleoside units of DNA that were modified by reaction with active vinyl chloride metabolites in vivo provides opportunity for the construction from appropriate animal data of the corresponding dose-response, time-response relationships, in comparison with the ones for tumor incidence/occurrence in those animals. The presence of these two imidazoneucleoside derivatives has now been established by mass fragmentometry (16) in the enzymic hydrolysate of modified rat-liver DNA, prepared from rats, which had been exposed chronically to vinyl chloride (250 ppm in their drinking water) for 1 year (Fig. 1). A smaller proportion of the 9β-D-2′-deoxy ribofuranosylimidazo-[2,1-i]purine, than would have been expected to have been formed, was found both in the animal experiments with vinyl chloride and in model reactions between chloroacetaldehyde and calf thymus DNA (16). This observation is consistent with some degree of DNA depurination brought about by the reaction of vinyl chloride, and in our model experiments, we have found evidence for the presence of the detached purine, viz., imidazo-[2,1-i] purine. Hence, the alkylation that produces imidazo-derivative formation (with DNA) labilizes the N9-purine β-glycoside linkage, which leads to depurination. The gap so produced might then be filled by various bases, resulting in “mispairing” during DNA replication. These results are very important, because in general, there is excellent agreement between the severe damaging effect of depurination to DNA and mutagenicity (17–19).

Thus, in retrospect, one would suspect vinyl chloride of being mutagenic/carcinogenic.

On the other hand, vinylidene chloride (k) metabolism in rats gave thiodiglycolic acid (r) and

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Scheme suggesting the model reaction of chloroacetaldehyde with (calf-thymus) DNA and the biotransformation of hepatocyte DNA by vinyl chloride in vivo. Both reaction processes afford 3β-D-2′-deoxy ribofuranosyl-3β-oxo-2,3-dihydroimidazo-[1,2-c]pyrimidine (left-hand side) and 9β-D-2′-deoxy ribofuranosylimidazo-[2,1-i]purine (right-hand side).

an N-acetyl-S-cysteiny1-acetyl derivative (p) as major urinary metabolites, plus substantial amounts of chloroacetic acid (l), dithioglycolic acid (t) and thiodiglycolic acid (s) (20). It is probable that chloroacetic acid (l), which is a vinylidene chloride metabolite per se, lies on a major metabolic pathway for vinylidene chloride, since it affords several metabolites in common with vinylidene chloride (20).

There is a strong supposition that detoxification of chloroacetic acid (l) is effected through a glutathione S-acetyl transferase-catalyzed reaction process and ensuing degradative sequence for the resulting carboxymethylglutathione (n), and that this represents the principal metabolic pathway for chloroacetic acid and a major one for vinylidene chloride. Thiodiglycolic acid is the ultimate detoxification product, and previous work (2) established the biotransformation of S-(2-carboxymethyl) cysteine (g) into that substance. A feasible metabolic pathway to thiodiglycolic acid from chloroacetic acid and involving cysteine desulphydrase is unacceptable. In experiments (rats) with unlabeled vinylidene chloride in which the cysteine-cystine pools had been labeled with 14C, labeled thiodiglycolic acid resulted, and a part of the C-skeleton of
that substance must be derived in fact from cysteine (20). Formation of a small amount of [14C]thiodiglycollic acid (t) and hence of the intermediate [14C]thiodiglycollic acid) (s) is reconcilable with the action of Michaelis's (21) unspecific β-thionase, which would lyse a small proportion of the preponderating [14C]thiodiglycollic acid.

Moreover, Kolbe electrolysis (22) of one molecular proportion of the [14C]thiodiglycollic acid metabolite from [1-14C]1,1-dichloroethylene or [1-14C]chloroacetic acid gave one equivalent of 14CO2 (23), and this evidence is consistent with the transformation of vinylidene chloride into chloroacetic acid by a mechanism involving migration of one Cl atom and the loss of the other one (20, 23). Hence, the metabolic pathway which was tentatively proposed for the biotransformation of vinylidene chloride into thiodiglycolic acid does in fact operate in rats.

It is equivocal whether the very small amounts of CO2 and urea are produced by the action of epoxide hydratase on 1,1-dichloroethylene oxide or by a minor oxidative pathway for chloroacetic acid.

There is a strong supposition that the N-acetyl-S-cysteinylacetetyl derivative (p), which is a metabolite of vinylidene chloride, but not of chloroacetic acid, may be formed in fact from 1,1-dichloroethylene oxide through the agency of glutathione S-epoxide transferase to afford S-glutathione acetetyl chloride (m) and its subsequent reactions (20). This supposition is important, since the reactivity displayed by 1,1-dichloroethylene oxide appears to be relevant to the possible interaction of reactive vinylidene chloride metabolites with mouse kidney DNA (Fig. 2), which is a prerequisite of tumor initiation (24). Such interaction would be analogous to that of vinyl chloride with rat-liver DNA in vivo, which forms imidazo derivatives with some nucleoside residues (16). Further work is in progress to investigate this hypothesis.

Comparative studies (25) provide clues of differences between rats and mice in the processing of vinylidene chloride (Table 1). Thus, in mice, the production of thiodiglycolic acid is considerably reduced and the formation of the N-acetyl-S-cysteinylacetetyl metabolite is increased. The higher β-thionase activity in mice than in rats accounts for the greater conversion of thiodiglycolic acid into dithioglycolic acid via thioylglycolic acid in the former species of animal. Yllner's (26) mice excreted a proportion of a dose of chloroacetic acid as unchanged starting acid. Thus, in mice, the metabolic pathway from chloroacetic acid to thiodiglycolic acid seems to be readily saturable, possibly on account of an inadequacy in the reaction catalysed by glutathione S-acyl transferase. Under these circumstances, detoxification of

![Figure 2. Scheme suggesting the feasible interaction of reactive vinylidene chloride metabolites, 1,1-dichloroethylene oxide and chloroacetyl chloride, with adenosine and cytidine respectively.](image)

Table 1. Relative proportion of products from metabolism of chloroacetic acid and vinylidene chloride in rats and mice.

| Substrate          | Metabolite                          | Yield of metabolites, % |
|--------------------|-------------------------------------|-------------------------|
| Chloroacetic acid  | Chloroacetic acid                    | Rats  | Yllner | BKJ-DEH |
|                    | Thiodiglycolic acid                  | 90    | 37     | 30-40   |
|                    | N-Acetyl-S-(2-carboxymethyl) cysteine| 2     | 40     | 40      |
| Vinylidene chloride| Chloroacetic acid                    | 3     | –      | –       |
|                    | Thiodiglycolic acid                  | 37    | 3      | –       |
|                    | Thioglycolic acid                    | 3     | 5      | –       |
|                    | Dithioglycolic acid                  | 5     | 20     | –       |
|                    | N-Acetyl-S-cysteinylacetetyl derivative| 48    | 70     | –       |

1,1-dichloroethylene oxide by glutathione S-epoxide transferase and the modification of DNA by 1,1-dichloroethylene oxide or chloroacetyl chloride would be expected to be more significant in mice than in rats. This diagnosis of species susceptibility seems to accord with Maltoni's (24) discovery of vinylidene chloride oncogenicity in (the kidneys of) mice.
Vinylidene chloride emerges as an agent of low, perhaps very low, oncogenic potential, which can be damaging only in a special set of biological circumstances, which we have partially defined and on which work is continuing.

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