CUMENE HYDROPEROXIDE-SUPPORTED DENITRIFICATION OF 2-NITROPROPANE IN UNINDUCED MOUSE LIVER MICROSOMES

ELIZABETH K. MARKER* and ARUN P. KULKARNI†
Toxicology Program, Department of Environmental and Industrial Health, School of Public Health, The University of Michigan, Ann Arbor, MI 48109, U.S.A.

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Abstract—1. Cumene hydroperoxide supported oxidative denitrification of 2-nitropropane was investigated in uninduced mouse liver microsomes.
2. The cytochrome P-450 peroxygenase catalyzed reaction resulted in the production of nitrite and acetone.
3. Several lines of evidence suggested the involvement of multiple forms of cytochrome P-450.
4. Acetone production was at least two times greater than nitrite release possibly due to sequestration of nitrite in the reaction mixtures.

INTRODUCTION
The metabolism of nitro-containing compounds by mammalian systems has been an area of study for many years. Earlier, the focus was on the reduction of the nitro group classically illustrated by p-nitrobenzoic acid reduction to p-aminobenzoic acid (Fouts and Brodie, 1957). Oxidative metabolism of 2-nitropropane (2NP) by rat liver microsomes was demonstrated by Ullrich et al. (1978). This novel reaction involved cytochrome P-450 (P450)-mediated denitrification of 2NP to yield nitrite and acetone as products. The involvement of P450 was substantiated by an increase in nitrite release after phenobarbital (PB) and 3-methylcholanthrene (3MC) induction, in vitro inhibition by metyrapone, α-naphthoflavone and carbon monoxide (CO), and reversal of CO inhibition by monochromatic light at 450 nm. The importance of this reaction was amplified in a subsequent study (Sakurai et al., 1979) which demonstrated the applicability of this reaction mechanism to other aliphatic nitro compounds. In contrast to the essential absence of denitrification activity reported by Ullrich et al., (1978) for hepatic microsomes from untreated rats, our work with 2NP in an NADPH-dependent system (Marker and Kulkarni, 1985, 1986) has shown significant nitrite releasing ability in uninduced CD~ mouse liver microsomes. That this activity was observed without induction of P450 by chemical pretreatment enhances the potential toxicological significance of hepatic 2NP metabolism. A similar reaction has also been reported in rabbit liver microsomes for 2-nitro-1-phenylpropane (Jonsson et al., 1977).

A few studies have reported reductive microsomal denitrosation of nitrosamines (Appel et al., 1980; Appel and Graf, 1982). Appel and Graf (1982) demonstrated nitrite release from nitrosamines in a reconstituted microsomal P450 system from PB induced pig liver. Spectral binding studies reported by these authors suggested a mechanism involving reductive cleavage of the nitroso group with a secondary amine as the leaving group. The nitroso group was thought to remain bound to the reduced P450 as nitric oxide which was non-enzymatically displaced from its binding site by molecular oxygen. The liberated nitric oxide was then partially converted to nitrite.

The report of Ullrich et al. (1978) retains some ambiguity as to the reaction mechanism of 2NP metabolism. The authors noted that lowering the oxygen concentration from 21 to 4% had no effect on the rate of 2NP denitrification which led them to believe that reduction of the nitro group to an amine was unlikely to occur. In theory, however, if oxidation is the only pathway involved in the metabolism of 2NP then a low oxygen concentration should have caused a proportional decrease in P450-dependent oxidative nitrite release. Thus, it was not clear whether the reaction was exclusively oxidative. The failure to observe the oxygen dependence of the reaction suggests the possibility that at least part of the nitrite release from 2NP in the NADPH-coupled reaction was through an oxygen-independent mechanism(s). This has also been suggested by investigations on 2NP denitrification in an NADPH-dependent mouse microsomal system (Marker and Kulkarni, 1986).

In order to establish unequivocally that denitrification of 2NP can proceed by an oxidative mechanism, the reaction was evaluated using mouse hepatic microsomes supplemented with cumene hydroperoxide (CHP). Organic hydroperoxides, through the peroxygenase activity of P450, can replace NADPH, NADPH cytochrome P450 reductase, and molecular oxygen in numerous types of P450-mediated reactions (O'Brien, 1978). When P450 acts as a peroxygenase the ferric ion in the active site remains in an oxidized state throughout the reaction cycle and, in the case of 2NP, any nitrite...
release or acetone production would be due to a totally oxidative reaction. In this communication we report CHP-supported denitrification of 2NP in uninduced CD mice liver microsomes.

**MATERIALS AND METHODS**

**Chemicals**

2NP (94%), metyrapone (2-methyl-1,2-di-3-pyridyl-1-propanone) and 1-naphthoflavone were obtained from Aldrich. HEPES (N-2-hydroxyethyl piperazine-N'-2-ethane sulfonic acid, sodium salt) was supplied by U.S. Biochemical, Cleveland, Ohio while imidazole was purchased from Eastman, Rochester, N.Y. Sigma Chemical Co., St Louis, Mo. supplied Tris (hydroxymethyl)ammonomethane, HCl (Tris), glycine, and mixed isomers of butylated hydroxyanisole (BHA), SKF-525A (2-diethylaminoethyl-2,2-diphenylvalerate, HCl) was the generous gift of Smith, Kline and French, Philadelphia, Pa. Metyrapone (2-methyl-1,2-di-3-pyridylpropane) and 1-naphthoflavone were obtained from Aldrich. CHP was purchased from Matheson, Coleman and Bell, Cincinnati, Ohio. The hydrochloride salt of N-oxycoumarine was synthesized in the laboratory. All other chemicals were reagent grade and available commercially.

**Animals**

Male CD mice were obtained from Charles River Breeding Laboratories, Portage, Mich. Mice were maintained on a 12 hr light/dark cycle and allowed to acclimate for one week after receipt. Food and water were available ad libitum. All mice were between 8 and 14 weeks of age at sacrifice.

**Isolation of microsomes**

Mice were sacrificed by cervical dislocation and livers quickly removed, weighed, placed in 15 ml ice cold homogenization buffer consisting of 50 mM Tris, 0.1 mM EDTA and 0.25 M sucrose, pH 7.4, and homogenized by 6-8 strokes of a motor driven Teflon pestle in a precooled glass mortar. The homogenate (15-20%) was centrifuged at 4°C for 20 min at 9000 g to remove cell debris and mitochondria. The post-mitochondrial supernatant was centrifuged at 4°C for 60 min at 106,000 g. The cytosol was discarded and the microsomal pellet washed by resuspension in a fresh 15 ml aliquot of ice cold homogenization buffer, followed by recentrifugation at 106,000 g for 60 min. The washed microsomes were resuspended in 2.0 ml sucrose (0.25 M).

**CHP-supported enzyme assay**

After protein determination by the Biuret method (Gornall et al., 1949) microsomes were diluted to a concentration of 2.0 mg of protein/ml in the specified buffer. Preliminary experiments indicated that, when CHP was included, the presence or absence of microsomes inactivated by boiling (10 min at 100°C) had no effect on non-enzymatic nitrite release from 2NP. Therefore, experiments were performed using control incubations containing 0.9 ml of the specified buffer to which all other reaction components except microsomes were added. Total activity was assayed in a flask containing 1.0 mg/ml active microsomal protein, the indicated concentrations of 2NP and CHP, and the specified buffer. Final volume of the complete reaction system was 1.0 ml for both the control (non-enzymatic) and total activity systems. Concentrations reported for reaction components are final concentrations.

All flasks were pre-incubated at 37 ± 1°C in a shaking water bath for 5 min. Following the addition of 2NP the reaction was initiated with CHP. The incubation time was 5 min after the addition of CHP unless otherwise specified. Nitrite was determined by a slight modification of the method described by Ullrich et al. (1978). The reaction was terminated by the removal of 0.5 ml of the reaction mixture to 0.5 ml ice cold zinc acetate (0.5 M in 50% ethanol) followed by the addition, while vortexing, of 0.5 ml ice cold 0.5 M sodium carbonate. The precipitate formed was removed by centrifugation at 2750 g for 5 min. A 0.5 ml aliquot of the clear supernatant was mixed with 1.0 ml sulfilamine (2% in 6N HCl) followed by 0.25 ml N-(1-naphthyl)ethylenediamine (0.16% in 6N HCl). After 15 min, absorbance of the colored product was measured at 540 nm. Enzymatic nitrite release was quantified from the difference between the total activity flask and the non-enzymatic activity flask. Standard curves were prepared using sodium nitrite under identical conditions.

**Identification and quantification of acetone**

Acetone was identified as the second oxidation product of 2NP by gas liquid chromatography (GC). All reaction components with the exception of 2NP and CHP were placed in 10 ml serum bottles kept on ice. After capping and sealing, the vials were preincubated at 37 ± 1°C for 5 min in a shaking water bath. The reaction was started by introducing 2NP (105 mM) through the rubber stopper with a Hamilton syringe followed by CHP (3.0 mM). (Optimal concentrations of 2NP were not used due to its interference with elution of the internal standard at the chromatography step.) After 5 min further incubation the reaction was stopped by addition through the stopper of 1.0 ml ice-cold 0.5 M aqueous zinc acetate. Vials were then placed on ice and the internal standard (IS) methylethylketone (MEK) (112 nmol/vial) was added. Vials remained on ice until head space analysis was performed.

GC was performed on a Varian 3700 chromatograph equipped with a flame ionization detector. Separation of the volatile components in the head space was achieved using a stainless steel column (6' × 1/8" o.d.) packed with 0.1% SP-1000 on 80/100 Carbopack C. The carrier gas was nitrogen with a flow rate of 24 ml/min. GC conditions were as follows: injector temperature, 150°C; detector temperature, 170°C; initial column temperature, 40°C; attenuation, 4 × 10^{-12}. The GC was programmed to hold the initial temperature for 5 min and then increase temperature 10°C/min to 150°C which was held for 10 min. For head space analysis, each vial was individually incubated for 30 min at 37 ± 1°C and 1 ml sample of the head space gas containing acetone was injected on column. A Finnegan mass spectrometer was used to confirm the presence of acetone as second product of the CHP-supported reaction. A purge and trap method was used to introduce volatiles dissolved in the reaction mixture onto the GC column interfaced with the mass spectrometer. The trap was a glass column 10" × 1/4" o.d. containing (in order from the inlet) silica gel, Tenax GC, and activated charcoal. The reaction mixture was purged into the cold trap by bubbling helium through the reaction mixture at 40 ml/min for 12 min. The trap was then desorbed for 2 min at 200°C onto the GC column. This column was a 6' × 2 mm i.d. packed column containing 0.1% SP-1000 on 80/100 Carbopack C. The column was maintained at room temperature during trap desorption. For this chromatography the flow rate of helium was 20 ml/min. Chromatography was begun at 50°C held for 2 min, and the temperature then increased at 8°C/min until 2NP appeared at which time the run was terminated.

Acetone standards were made for each assay in buffer and microsomes and were run through the assay procedure as if they were test samples except that CHP was excluded. Acetone from these standards eluted at a retention time of 3.6 min while MEK had a retention time of 9.7 min. Vials in which metabolism of 2NP occurred also contained a peak eluting at the same retention time as acetone. The peak height ratio of acetone to MEK in the samples increased with reaction time suggesting that acetone is an enzymatic reaction product. Quantification of acetone was by com-
CHP-supported 2NP oxidation

RESULTS

Similar to the NADPH and oxygen dependent reaction reported for hepatic microsomes from PB or 3MC induced rats (Ullrich et al., 1978) and uninduced mice (Marker and Kulkarni, 1985, 1986) nitrite release from 2NP was observed when uninduced male CD1 mouse hepatic microsomes were supplemented with CHP. Through a series of experiments, the CHP-supported reaction was characterized and the results are presented in the accompanying figures and tables.

The effect of pH on denitrification activity is illustrated in Fig. 1. The major activity peak was seen at pH 8.6 with a shoulder at pH 7.6. These results correlate well with those found in our investigations of the NADPH-dependent reaction in uninduced mice (Marker and Kulkarni, 1986) and in induced rats (Ullrich et al., 1978). Consequently, all further experiments were conducted at both pH 7.6 and 8.6.

The effect of different buffers at two pHs on relative nitrite releasing activity was studied. Considering both pHs, relative activity was from 9 to 58% greater in 0.1 M Tris than in HEPES buffer, sodium or potassium phosphate buffers, potassium chloride/boric acid/NaOH buffer, or glycine/NaOH buffer. Therefore, 0.1 M Tris was used in all subsequent experiments.

To determine the optimal concentration of CHP for denitrification, a range from 0.1 to 10.0 mM were tested. At pH 7.6 nitrite release was marginally greater at 2.0 mM than at 3.0 mM, while at pH 8.6, 3.0 mM was slightly greater than 2.0 mM. Since no statistical difference among the values was observed from 1.0 to 5.0 mM CHP at pH 7.6 or pH 8.6, 3.0 mM CHP was used for all subsequent experiments. Higher concentrations (>5 mM) were less effective.

Substrate (2NP) concentration optima were found to be 262 mM at pH 7.6 and 157 mM at pH 8.6 (Fig. 2). The apparent $K_m$ values determined from...
these data were 83 mM at pH 7.6 and 65 mM at pH 8.6. In addition, the CHP-supported reaction was found to be linear with respect to protein concentration up to 3.0 mg/ml at both pHs (Fig. 3).

Time dependence of the reaction was also assessed and linearity was observed up to 15 min at pH 7.6 but only up to 7.5 min at pH 8.6 (Fig. 4). Non-linearity after 7.5 min at pH 8.6 may be due to inactivation of P450 by a combination of high pH and CHP-induced lipid peroxidation over time with accompanying membrane disruption (Kulkarni and Hodgson, 1981). The routinely used 5 min reaction time is within the linear portion of the response curve in all the cases.

Under optimal conditions the specific activities observed for nitrite release were: 17.1 at pH 7.6 and 16.3 at pH 8.6. Acetone was identified by GC and confirmed by MS as second product of the oxidative denitrification of 2NP (Fig. 5).

The results of the balance study are shown in Table 1. A difference of over 200% between acetone production and nitrite release was observed. The possible reasons for this finding may involve the chemistry of nitrite and CHP and are discussed below.

The effects of chemicals known to modify the peroxygenase activity of P450 were tested and the results are presented in Table 2. The most effective inhibitor at both pH 7.6 and pH 8.6 was n-octylamine which inhibited the reaction by 30 and 41% respectively. Differential effects were seen for BHA and SKF-525A. BHA decreased activity 34% at pH 8.6 but only 13% at pH 7.6. SKF-525A, on the

<table>
<thead>
<tr>
<th>pH</th>
<th>Acetone</th>
<th>Nitrite</th>
<th>acetone/nitrite</th>
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</thead>
<tbody>
<tr>
<td>7.6</td>
<td>16.50 ± 0.70</td>
<td>7.28 ± 1.18</td>
<td>2.27</td>
</tr>
<tr>
<td>8.6</td>
<td>20.84 ± 0.49</td>
<td>8.65 ± 0.70</td>
<td>2.41</td>
</tr>
</tbody>
</table>

Specific activity is expressed as nanomoles of acetone produced or nitrite released/min/mg protein. The concentration of 2NP used was 105 mM (optimal concentrations: pH 7.6, 262 mM; pH 8.6, 157 mM). At pH 7.6, n = 4; at pH 8.6, n = 3. See "Materials and Methods" for further details.
other hand, produced 23% inhibition at pH 7.6 but caused a non-significant decrease in activity at pH 8.6. No significant decrease in activity was observed with imidazole, metyrapone or α-naphthoflavone.

**DISCUSSION**

Earlier, the involvement of P450 in the metabolism of 2NP was clearly shown (Ullrich et al., 1978). However, the contribution of an oxidative mechanism in the overall denitrification process is difficult to assess in an NADPH-dependent microsomal system when oxygen-independent pathways are concurrently operative.

Ullrich et al. (1978) working with an NADPH-dependent rat liver microsomal system, observed no change in nitrite release from 2NP when the oxygen concentration was reduced from 21 to 4%. It is possible that 4% oxygen is sufficient for the NADPH-dependent system to function at full capacity although a decrease in activity is more likely. Alternatively, a reductive mechanism similar to that postulated for nitrosamines by Appel, et al. (1982) may be operating. Of lesser importance is the contribution of hydrogen peroxide derived from the NADPH oxidase activity of P450 which could promote oxidative metabolism of 2NP through the peroxygenase activity of P450 (Estabrook et al., 1984). It has been postulated that 2NP may cause hepatic lipid peroxidation (Zitting et al., 1981). The lipid peroxides formed during this process, in concert with preformed lipid peroxides, may contribute to a certain extent to oxidative metabolism of 2NP as has been demonstrated for benzo(a)pyrene (Morgenstern et al., 1981; Dix and Marnett, 1983). Lastly, if P450 were to act as a true peroxidase then a chain reaction mechanism analogous to that reported for 2NP metabolism by horseradish peroxidase (HRP) is also possible (Porter and Bright, 1983).

Our earlier work involving NADPH-dependent P450-mediated 2NP metabolism in uninjured mouse liver microsomes (Marker and Kulkarni, 1986) indicated that anaerobic conditions produce a substantial but not complete elimination of nitrite releasing activity. Decline in nitrite release due to anaerobiosis at pH 7.6 was 83.7% and at pH 8.8 was 74.4% suggesting that other oxygen-independent mechanisms for the metabolism of 2NP are operating in the presence of NADPH. The oxygen sensitivity of these reactions is unknown at present.

The ability of organic hydroperoxides such as CHP to replace NADPH, NADPH cytochrome P450 reductase and molecular oxygen in microsomal P450-mediated metabolism has been recognized for some time (O’Brien, 1978). It has been suggested that P450, in the presence of hydroperoxides, can perform all reactions typical of NADPH-dependent P450 oxidative metabolism of xenobiotics with the possible exception of N-hydroxylation (O’Brien, 1978). The main advantage is that only oxidative reactions proceed in this system. Thus, a study of an oxidative pathway is possible without the interference of reductive mechanisms which apparently occur in an NADPH-supplemented microsomal system.

The investigations presented here describe for the first time denitrification of a nitroalkane (2NP) by hepatic microsomal P450 in the presence of the organic hydroperoxide, CHP. Specific activities of 17.1 and 16.3 at pH 7.6 and 8.6 respectively were observed in CHP-supplemented hepatic microsomes from uninjured CDl mice. In the NADPH-dependent reaction, mouse hepatic microsomes exhibited maximal specific activities of about 8.0 and 6.4 at pH 7.6 and pH 8.8 respectively (Marker and Kulkarni, 1985, 1986). This is in contrast to Ullrich et al. (1978) who reported specific activities of 10 in PB induced rats and 0.5 in 3MC induced rats. Values for microsomes from untreated rats were not reported but were stated to be very low or undetectable.

The enzymatic nature of the CHP-supported reaction is evident from the observation that boiled microsomes reduce nitrite release to the non-enzymatic level. Linearity of the reaction with protein concentration and time also suggests an enzyme catalyzed reaction. Acetone, the expected second product of the oxidative denitrification of 2NP, was identified by GC and confirmed by MS.

Under the conditions employed, known modifiers of P450 mono-oxygenase/peroxygenase activity appear to be less effective inhibitors of 2NP denitrification than of other CHP-supported reactions. The inhibition patterns found in our experiments do not appear to be classifiable by the interaction of the inhibitor with P450 (Type I vs Type II ligands). The most effective inhibitor, under the conditions used, was n-octylamine, a Type II compound, which produced a statistically significant inhibition of nitrite

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**Table 2. Effects of modifiers of P450 peroxygenase activity on CHP-supported 2NP denitrification**

<table>
<thead>
<tr>
<th>Modifier</th>
<th>pH 7.6</th>
<th>pH 8.6</th>
</tr>
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<tbody>
<tr>
<td>None</td>
<td>100.0</td>
<td>100.0</td>
</tr>
<tr>
<td>Imidazole (1.0 mM)</td>
<td>102.5 ± 9.6</td>
<td>104.9 ± 13.2</td>
</tr>
<tr>
<td>n-Octylamine (1.0 mM)</td>
<td>70.2 ± 3.6*</td>
<td>58.3 ± 4.3*</td>
</tr>
<tr>
<td>Metyrapone (1.0 mM)</td>
<td>92.8 ± 4.1</td>
<td>92.5 ± 5.7</td>
</tr>
<tr>
<td>α-Naphthoflavone (0.2 mM)</td>
<td>90.1 ± 1.7</td>
<td>103.5 ± 8.2</td>
</tr>
<tr>
<td>Butylated hydroxyanisole (0.1 mM)</td>
<td>86.5 ± 2.8</td>
<td>66.0 ± 7.1*</td>
</tr>
<tr>
<td>SKF-525A (1.0 mM)</td>
<td>76.5 ± 3.5*</td>
<td>94.5 ± 1.8</td>
</tr>
</tbody>
</table>

*Statistically different from control: P < 0.01.
Each value represents the mean ± SE of determinations from four separate microsomal preparations. Mean specific activity for controls at pH 7.6 was 17.5 nmol of nitrite released/min/mg protein and at pH 8.6 was 16.2 nmol of nitrite released/min/mg protein.
release at both pHs tested. Imidazole, another Type II compound, however, was without effect.

Metyrapone and alpha-naphthoflavone also proved to be ineffective inhibitors. A widely range of values for inhibition by metyrapone of organic hydroperoxide-supported P450 reactions have been reported (Rahimtula and O’Brien, 1974, 1976; Ashley and Griffin, 1981; Cederbaum, 1983). These values vary from 80% inhibition of CHP-supported amino-pyrene \( N \)-demethylation by 0.1 mM metyrapone (Ashley and Griffin, 1981) to only 25% inhibition of \( t \)-butyl hydroperoxide and CHP-supported microsomal ethanol oxidation by 1.0 mM metyrapone (Cederbaum, 1983). Taken together these reports suggest that the degree of inhibition due to any particular agent may be dependent on the substrate and the animal species in question.

The effectiveness of these inhibitors may also depend in part on the P450 species involved in the reaction. The possible involvement of two or more P450 species in CHP-supported denitrification of 2NP is suggested by the presence of two peaks of activity at pH 7.6 and 8.6. This is further supported by different optimal substrate concentrations at each pH, and by pH dependent patterns of response to the inhibitors BHA and SKF-525A. SKF-525A showed a statistically significant inhibition of nitrite release at pH 7.6 but not at pH 8.6. BHA, which is metabolized by P450 in a CHP-supported system (Rahimtula, 1983) displayed a significant inhibition only at pH 8.6. Differing patterns of BHA inhibition due to the participation of multiple isozymes in oxidative reactions have also been postulated by Yang et al. (1981) working with aryl hydrocarbon hydroxylase activity in hepatic microsomes from control animals or animals induced with PB, 3MC or 2,3,7,8-tetrachlorodibenzo-p-dioxin. Varying affinities of different P450 species for metyrapone have also been postulated by Cinti (1978). Similar responses to changes in pH, substrate concentration and inhibitors observed in the NADPH-dependent reaction also implicate involvement of multiple forms of P450 in the denitrification of 2NP in uninduced mouse liver microsomes (Marker and Kulkarni, 1986). This contention is further supported by a report on the varying effectiveness of NADPH-dependent denitrification of 1-nitropropane and nitroethane by different reconstituted isozymes of P450 (Sato et al., 1982).

The balance study of acetone production to nitrite release in the CHP-supported reaction (Table 1) produced unexpected results. Since the presence and purity of acetone were confirmed by MS, the imbalance in the stoichiometry of acetone:nitrite production may be due to sequestration of nitrite in the reaction mixture. The results of a recent study (Pryor et al., 1985) investigating nitrosation of organic hydroperoxides provide, in part, a possible explanation for the presumed nitrite sequestration. These authors showed that, in hexane, CHP will react within seconds with an \( \text{NO}_2^-/\text{N}_2\text{O}_5^- \) mixture to form cumyl nitrate, cumyl nitrite, cumyl alcohol and acetophenone in ratios of 58:13:11:17 respectively. The mixture of \( \text{NO}_2^-/\text{N}_2\text{O}_5^- \) can be obtained from nitrite or nitrous acid through an \( \text{N}_2\text{O}_3^- \) intermediate (Anonymous, 1981). It is thus possible that a significant portion of the total nitrite generated during denitrification may be reacting with CHP and be undetectable by the methods employed. The possibility also exists that nitrite may be reacting in a similar fashion with lipid peroxides formed by the interaction of CHP with microsomal lipids and this may in part explain the difference in the acetone:nitrite ratio. Similarly lipid peroxide radicals produced by interaction of CHP with microsomal membranes could also interact with an \( \text{NO}_2^- \) radical (Pryor et al., 1982) contributing to the nitrite loss. It appears from this discussion that several secondary products may be generated and further studies are needed to identify and quantitate these products.

It is proposed that the first step in the mechanism for the CHP-supported oxidative denitrification of 2NP involves the peroxygenase activity of P450. However, the possibility of a subsequent or concurrent chain reaction mechanism similar to that reported for 2NP metabolism by HRP cannot be discounted at present (Porter et al., 1983). HRP Compound I was reported to initiate a chain reaction resulting in nitrite and acetone as products. The reaction, however, depended on the presence of propane-2-nitronate. This species is derived from 2NP but the equilibrium has a \( t_{1/2} \) of 150 min at pH 5.0 and thus is unlikely to contribute significantly to acetone production or nitrite release in the 5 min reaction time used in our study.

**CONCLUSION**

The evidence presented indicates that P450-mediated oxidative metabolism of 2NP occurs in CHP-supplemented hepatic microsomes from uninduced CD\(_1\) mice. The observed stoichiometry suggests that the specific activity based on acetone produced is the appropriate measure of 2NP denitrification in the CHP supported system since nitrite appears to be sequestered in the reaction mixture. The results suggest the generation of yet unidentified products resulting from secondary reactions of the unaccounted nitrite released. The possibility of a chain reaction mechanism in the CHP supported denitrification reaction exists but seems unlikely. It is suggested that, through the peroxygenase activity of P450, oxygen from CHP is introduced into 2NP possibly through a hydroxylated intermediate as suggested by Ullrich et al. (1978). Rearrangement of the unstable intermediate produces acetone with concomitant release of nitrite. This is supported by the identification of nitrite and acetone as the products in the CHP system. This does not imply, however, that the NADPH-dependent denitrification of 2NP in uninduced CD\(_1\) mouse liver microsomes is solely oxidative.

**SUMMARY**

Cytochrome P-450 (P450) mediated denitrification of 2-nitropropane (2NP) was studied in cumene hydroperoxide (CHP) supplemented hepatic microsomes from male CD\(_1\) mice. Under optimal conditions, the mean specific activities of 17.1 and 16.3 nmoles of nitrite released/min/mg microsomal protein were observed at pH 7.6 and 8.6, respectively. The oxidative metabolism of 2NP produced two
precisely monitored by acetone production rather than by quantification of nitrite released.


Sakurai H., Hermann G. and Ullrich V. (1979) "The interaction of aliphatic nitro compounds with the liver microsomal monoxygenase system." Biochem. Pharmac. 29, 341-345.


