Progressive liver fibrosis in rats develops when they are fed a diet deficient in choline. This diet also results in a pronounced and selective decrease in the liver microsomal content of a phase I drug-metabolizing enzyme belonging to the cytochrome P-450III gene family. Because P-450III cytochromes characteristically catalyze the N-demethylation of erythromycin, we believed that the production of breath CO₂ from erythromycin would be dramatically reduced in choline-deficient rats. However, when 12 choline-deficient rats were compared with 9 control rats, the reduction in CO₂ production from erythromycin (mean decrease 71%) was essentially identical to that from aminopyrine (mean decrease 69%), a substrate believed to be metabolized normally by the hepatocyte in fibrotic liver disease. Furthermore, we found that the relative erythromycin and aminopyrine demethylase activities were comparable when measured in vitro in liver microsomes prepared from the choline-deficient rats. To determine the molecular basis for the erythromycin demethylase activity in the choline-deficient rats, the liver microsomes were subjected to immunoblot analysis using a variety of polyclonal and monoclonal antibodies capable of distinguishing individual P-450III-related proteins. Our studies confirm that a major erythromycin demethylase belonging to the P-450III family, termed P-450p, was greatly reduced in the choline-deficient rat liver. However, the specific concentration of a second P-450p-related protein was essentially normal and that of a third P-450p-related protein was actually increased in the choline-deficient rat liver. These changes occurred over weeks and months on the choline-deficient diet and are not consistent with "feminization" of the liver. The P-450p-related proteins appeared to catalyze erythromycin demethylase activity because antibodies recognizing them inhibited the majority (62%) of this activity in choline-deficient rat microsomes. Finally, RNA extracted from choline-deficient rat livers was hybridized on Northern blots with synthetic oligonucleotide probes that identify the only two known rat P-450III cDNAs, PCN-1 and PCN-2. Prolonged choline deficiency appeared to have no consistent effects on expression of these RNA species. We conclude that despite the down-regulation of a major erythromycin demethylase, this catalytic activity is relatively preserved in the choline-deficient rat, at least in part because of differential regulation of P-450III cytochromes. Erythromycin therefore appears to offer little advantage over aminopyrine as a substrate for assessing liver function in this model of chronic liver disease. (Hepatology 1990;12:1371-1378.)

The ability to metabolize many drugs is impaired in patients with cirrhosis. Based on this observation, it has been proposed that the severity of chronic liver disease and cirrhosis can be quantitated in at least some patients by measuring their ability to metabolize a test dose of aminopyrine (reviewed in reference 1). Aminopyrine is demethylated in the liver by the mixed function oxidase system, which is made up of multiple families of genetically related enzymes collectively termed cytochromes P-450. Because a large proportion of the carbon in the cleaved methyl group promptly appears in the breath as CO₂, the ability to demethylate aminopyrine can be estimated in a patient simply as the rate of production of [14C N-methyl] aminopyrine (2). Enthusiasm over the potential usefulness of the aminopyrine breath test has been tempered, however, by the observation that the result is influenced by many other concomitantly administered medications (reviewed in reference 1). Furthermore, ethanol consumption appears to increase the rate of production of CO₂ from aminopyrine (3), presumably by inducing liver cytochrome(s) P-450 capable of demethylating aminopyrine. The breath test may therefore be difficult to interpret in the large population of patients with alcoholic liver disease.

We have recently described a breath test that uses erythromycin as a substrate (4). We believed that erythromycin might be preferable to aminopyrine as a substrate for assessing liver function for two reasons. First, the major erythromycin demethylases in liver
have been identified and belong to the P-450III gene family (5). Because P-450III cytochromes have been reasonably well characterized in rat and man (6-13), medications likely to influence erythromycin demethylation are known (4, 6-12) and ethanol should have no effect (13, 14). Second, the available evidence suggested that in chronic liver disease and cirrhosis, the ability to demethylate erythromycin might be reduced to a greater extent than the ability to demethylate aminopyrine. Chronic liver disease appears to have differential effects on the regulation of individual cytochromes P-450 and their corresponding catalytic activities (15-18). Moreover, Murray et al. (18) showed that the concentration of at least one major P-450III cytochrome was less than 5% of normal in microsomes prepared from rats given fibrotic liver disease by prolonged choline deficiency. In the same microsomes, however, the concentrations of some other cytochrome P-450 proteins were essentially normal (18). This indicated that P-450III cytochromes may be selectively down-regulated in the fibrotic liver, which, in turn, would be expected to result in loss of the hepatocyte's intrinsic ability to demethylate erythromycin. On the other hand, Reichen et al. (19) have recently presented data suggesting that the hepatocyte's intrinsic ability to demethylate aminopyrine is not significantly altered in rats with fibrotic liver disease. These authors proposed that the reduced ability to demethylate a test dose of aminopyrine in chronic liver disease reflects only the reduced hepatocyte mass and the alterations in liver blood flow that characteristically accompany cirrhosis. Taken together, these observations suggested that the cytochromes P-450 involved in aminopyrine demethylation, which have not yet been identified, are probably present in near normal concentration in cirrhotic liver hepatocytes, whereas the concentration of forms primarily involved in erythromycin demethylation may be significantly decreased. Since reduction in hepatocyte mass and liver circulatory changes should have comparable effects on the metabolism of both substrates, we reasoned that there might be greater impairment in production of CO2 from erythromycin than from aminopyrine in rats with liver fibrosis. If so, the erythromycin breath test might provide a very sensitive parameter of liver function in chronic disease and cirrhosis.

To test this idea, we administered the erythromycin and aminopyrine breath tests to groups of rats given fibrotic liver disease by prolonged nutritional deficiency. Contrary to our expectation, we found that when these rats were compared with healthy rats, the fall in CO2 production from each substrate was essentially identical. To investigate the molecular basis for the relative preservation of P-450III catalytic activity, the diseased liver microsomes were analyzed using antibodies that are capable of distinguishing members of the P-450III gene family. We confirmed that a major liver P-450III protein is almost completely repressed in the choline-deficient (CD) animals. However, the microsomal concentration of a second P-450III-related protein was unchanged and that of a third related protein was actually increased.

MATERIALS AND METHODS

Animals and Treatments. Male, Sprague-Dawley rats (80 to 100 gm) were obtained from Charles River, Inc., (Wilmington, MA) and housed two to five per cage with free access to water. After acclimatization, rats had free access to either a custom high-fat, CD chow (ICN Nutritional Biochemicals Diet #901387, ICN, Cleveland, OH) that was free of methionine, folic acid, and vitamin B12 (referred to as the "choline-deficient" or "CD" diet) or an identical "control" chow supplemented with choline chloride (165 mg/100 gm), methionine (230 mg/100 gm), folic acid (198 µg/100 gm) and vitamin B12 (3 µg/100 gm) (referred to as the "control" diet). Rats in both groups gained weight, but at 7 mo rats on the CD diet weighed 45% less than rats on the control diet (mean 398 gm vs. 719 gm). After 2 wk to 7 mo on the diets, rats underwent breath tests and/or were killed by decapitation, and liver tissue assays were performed.

After the animals were decapitated, the liver was perfused through the hepatic vein with PBS, frozen in liquid nitrogen, and stored at -70°C. Hepatic hydroxyproline content was assayed as previously described (20) from 4 gm of liver tissue taken from the right lobe. Hepatic hydroxyproline was elevated an average of 2.2-fold in livers of 13 rats maintained for 7 mo on the CD diet relative to the livers of 9 rats maintained on the supplemented diet (4.02 vs. 1.81 µmol hydroxyproline/gm liver; p < 0.001 according to an unpaired t test). Histological examination revealed normal histology in a randomly selected supplemented control rat, whereas fatty infiltration and fibrosis were present in a randomly selected CD diet animal (not shown). These studies confirmed that significant liver fibrosis had occurred as a result of the CD diet. All studies were approved by the University of Michigan Committee on Animal Use.

Breath Tests. The erythromycin breath test (ERMBT) was performed essentially as described (4). Rats received tail vein injections of 0.5 to 1.0 µCi/100 gm body weight [14C N-methyl] erythromycin (54.3 mCi/mmol, New England Nuclear Corp., Boston, MA) and did not concomitantly receive unlabeled erythromycin. Trace doses of erythromycin were used because the pharmacological doses given in prior studies (4) quickly killed the animals who had received the CD diet. The aminopyrine breath test (APBT) was performed by injecting 0.5 µCi/100 gm body weight of [14C dimethyl] aminopyrine (101.8 mCi/mCi, New England Nuclear Corp.) into the tail vein. After the tail vein injections, the rats were placed in a water-sealed polyurethane breath chamber and CO2 was collected at intervals and assayed for 14C (4). The results of each breath test are expressed as the percent of injected 14C recovered during the first hour after injection.

When both breath tests were performed on the same rats, the APBT did not influence the results of the ERMBT when the tests were separated by 24 hr.

Microsomal Assays. The liver tissue remaining after hydroxyproline determination was used for preparation of microsomes by differential centrifugation (21). Up to 100 µg of microsomal protein was then subjected to 10% SDS-PAGE, and the separated proteins were transferred to nitrocellulose sheets. The nitrocellulose sheets were then reacted with antibodies as previously described (22). The monoclonal antibodies 1G8 and 13-7-10 have been previously characterized (9, 23) and were gifts from Dr. Philip Guzelian. The antibody used to recognize the ethanol-inducible cytochrome P-450 (P-450IE1) was a gift from Dr. Minor J. Coon. Erythromycin demethylase and aminopyrine demethylase activities were determined in liver microsomes by the Nash
**RESULTS**

**Aminopyrine and Erythromycin Demethylase Activity.** ERMBT and APBT were administered to CD and control rats. The production of CO₂ from aminopyrine was diminished by approximately 70% in the CD rats compared with that observed in the controls (Table 1). This decrease is comparable to that previously reported in rats with established cirrhosis (19). When the ERMBT was administered, the production of CO₂ from erythromycin was also reduced approximately 70% in the deficient rats (Table 1). Furthermore, in the rats that had received both breath tests, there was excellent correlation between the production of CO₂ from each substrate (r = 0.99, data not shown). The comparable falls in the production of CO₂ from aminopyrine and erythromycin did not support a selective loss of *in vivo* P-450III catalytic activity in the CD rats. However, it was possible that factors other than liver content of the responsible cytochromes P-450 became rate limiting in the production of CO₂ from these substrates in the CD rats.

We therefore directly examined whether erythromycin demethylase activity was selectively reduced in microsomes prepared from the liver of each CD rat. As shown in Table 2, the ability to demethylate aminopyrine in the CD rat liver microsomes was approximately 50% of that seen in control microsomes. Comparable falls in microsomal aminopyrine demethylase activity have been reported previously in cirrhotic rats (19). Since the intrinsic ability of the hepatocyte to demethylate aminopyrine appears to be normal in the
cirrhotic liver (19), the 50% decrease in specific activity may reflect dilution of the hepatocyte microsomes with those derived from other cell types (such as macrophages and bile ductular epithelia) that are abundant in the diseased liver.

The erythromycin demethylase activity was also significantly lower in the hepatic microsomes prepared from CD rats compared with control microsomes (Table 2). However, the fall in erythromycin demethylase activity was not greater than that in aminopyrine demethylase activity. This was also true when the respective demethylase activities were estimated for the entire liver (Table 2). Hence, the catalytic activity characteristic of P-450IIIA appeared to be relatively preserved in the CD liver microsomes, an observation that seemed at odds with the report that at least one major P-450IIIA protein should be virtually absent (18).

**Immunological Analysis of Liver Microsomes.** The liver microsomes prepared from each rat were analyzed on immunoblots to determine whether expression of P-450III cytochromes had in fact been repressed in the CD rats. ImmunobLOTS were first developed with a polyclonal antibody that selectively recognizes a major rat liver P-450III cytochrome termed P-450p (8, 11). In uninduced rats, P-450p is generally detectable only in male liver (9, 11). The P-450p specific antibody we used was prepared byabsorbing anti-P-450p IgG (goat) to liver microsomes prepared from a female rat (11). The absorption step removes antibodies that recognize epitopes common to P-450p and to other P450III proteins present in the female rat liver (9). As is shown in Figure 1A, P-450p was detected in the control liver microsomes but was barely detected, if at all, in the CD liver microsomes, essentially confirming the observations of Murray et al. (18). The repression of P-450p appeared to be a selective effect because there was no detectable difference between control and CD microsomes in the content of immunoreactive P-450 (P-450IIIE1) (data not shown). However, P-450p could still be induced in the CD rat by pretreatment with dexamethasone (Fig. 1A), indicating that the hepatocyte remained capable of expressing this enzyme. It therefore appeared that erythromycin demethylation was being catalyzed in the CD rat liver by enzymes other than P-450p.

It has recently been appreciated that rat liver contains multiple P-450III cytochromes (9, 25, 27). We reasoned that P-450III cytochromes other than P-450p (and those not recognized by the absorbed polyclonal antibody to P-450p) might account for the relative preservation of erythromycin demethylase activity in the CD rat liver. Immunoblots of the rat liver microsomes were therefore developed with the unabsorbed anti-P-450p IgG, which would be expected to recognize multiple members of the P-450III gene family. As shown in Figure 1B, blots developed with the unabsorbed IgG confirmed that the concentration of P-450p had been greatly reduced in the CD microsomes. However, a second protein band was recognized by the unabsorbed antibody in the CD microsomes, reflecting a protein that migrated more rapidly than did P-450p on the gels (Fig. 1B). To estimate the proportion of erythromycin demethylase activity that was being catalyzed by proteins recognized by this antibody, microsomes prepared from representative control and CD rats were incubated with the anti-P-450p IgG or with nonimmune IgG (10 mg/nmol spectrally determined P-450) before measuring catalytic activity toward this substrate. Erythromycin demethylase activity was inhibited to a comparable extent in liver microsomes prepared from each animal (54% and 62% inhibition in control and deficient microsomes, respectively, compared with the activity seen in the microsomes containing nonimmune IgG). We concluded that the proteins recognized by the nonabsorbed anti-P-450p IgG catalyzed most erythromycin demethylase activity in the livers of both CD and control animals.

We next used monoclonal antibodies in an attempt to determine the identity of the P-450p–related protein that appeared to be induced in the CD liver microsomes. As shown in Figure 1C, the induced protein appeared to be recognized by a monoclonal antibody termed 1G8. 1G8 has been previously shown to recognize P-450p (9), and the absence of P-450p in the CD microsomes was also confirmed using this antibody (Fig. 1C). In addition, 1G8 recognizes a second protein that is male specific in uninduced rats, and this protein migrates slightly faster than P-450p on SDS-PAGE (9, 28). As shown in Figure 1C, 1G8 appeared to recognize the P-450p–related protein present in the CD microsomes, and this protein exactly comigrated with the faster migrating protein present in control liver microsomes (hereafter referred to as “p2”).

**Table 2. Results of erythromycin demethylase activity and aminopyrine demethylase activity in liver microsomes from rats on a control or CD diet**

<table>
<thead>
<tr>
<th>Diet</th>
<th>ERMD (nmol/min/mg protein)</th>
<th>APD (nmol/min/mg protein)</th>
<th>ERMD* (nmol/min/liver)</th>
<th>APD* (nmol/min/liver)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (n = 9)</td>
<td>1.19 (.13)*</td>
<td>2.92 (.38)</td>
<td>363 (64)</td>
<td>888 (144)</td>
</tr>
<tr>
<td>CD (n = 13)</td>
<td>0.89 (.29)</td>
<td>1.48 (.23)</td>
<td>256 (85)</td>
<td>425 (101)</td>
</tr>
<tr>
<td>Percent change</td>
<td>-25%</td>
<td>-50%</td>
<td>-30%</td>
<td>-50%</td>
</tr>
</tbody>
</table>

*Determined as (specific activity) × (microsomal protein from entire liver).

*p values in parentheses are standard deviations.

*p < 0.01 determined by an unpaired t test.

ERMD = erythromycin demethylase; APD = aminopyrine demethylase; CD = choline deficient.
Finally, the immunoblots were developed with another monoclonal antibody, termed 13-7-10, that identifies additional P-450p-related proteins present in liver microsomes prepared from untreated male and female rats (9, 28). As shown in Figure 2, the reactive protein present in untreated male liver migrates more rapidly on PAGE than does the female reactive protein, as has been previously reported (9, 28). It has been shown (9) that the 13-7-10 reactive protein present in control male liver microsomes exactly comigrates with p2. Liver microsomes prepared from our CD (male) rats contained a reactive protein indistinguishable from the male-specific 13-7-10 reactive protein on immunoblots (Fig. 2). In addition, the specific content of this immunoreactive protein appeared to be identical in microsomes prepared from either the CD or control rats (Fig. 2 and data not shown).

To see whether repression of P-450p and induction of the related protein were due to an immediate effect of the diet, rats were placed on the CD and control diets. Members of each treatment group were killed 2 wk later and at the end of each consecutive month on the diets. Microsomes prepared from each rat were analyzed on immunoblots developed with the monoclonal antibody 1G8. As shown in Figure 3, the loss of immunoreactive P-450p and the appearance of the related protein occurred gradually over time on the CD diet.

**mRNA Studies.** To date only two complete cDNAs corresponding to rat P-450III genes have been cloned and sequenced, and these have been termed PCN-1 and PCN-2 (25), which correspond to the genes P450III A1 and P450III A2, respectively (5). We sought to determine whether regulation of these mRNAs was altered in the CD rat. In addition, by comparing PCN-1 and PCN-2 mRNA content in control and CD rat livers, we hoped it would be possible to tentatively identify the P-450III-related proteins recognized by our antibodies. This would, however, require that the loss of P-450p and the appearance of the related protein (Fig. 3) had resulted in the CD rat from alterations in regulation at steps before mRNA translation. Accordingly, total RNA extracted from the livers of control and CD rat livers was analyzed on Northern blots hybridized with 30-mer synthetic oligonucleotide probes corresponding to hypervariable regions of PCN-1 and PCN-2 cDNA (25). PCN-1 RNA has been reported to be inducible by dexamethasone but not to be readily detected in liver RNA prepared from control male or female rats (25). Our results confirmed these observations (Fig. 4). We also found that PCN-1 mRNA was undetectable in the livers from each of the CD rats (Fig. 4).

We found that PCN-2 mRNA was readily detected in male but not female liver and was not inducible by dexamethasone (Fig. 4); this confirms prior reports (25).
Among individual male liver RNA samples, the intensity of hybridization with the PCN-2 probe varied remarkably (Fig. 4). However, there were no consistent differences in PCN-2 RNA concentrations between the CD and control liver RNA (Fig. 4). Moreover, when liver microsomes prepared from each of these rats were subjected to immunoblot analysis, there were no obvious correlations between the concentrations of any of the P-450p-related proteins and liver content of PCN-2 mRNA (data not shown). To assure that comparable amounts of RNA were being analyzed from each rat, the blot was also hybridized with a synthetic oligonucleotide specific for P-450IIE1.

We observed generally higher amounts of P-450IIE1 mRNA in female than in male rats (Fig. 4 and data not shown), and this has been observed by others (14). No consistent differences in hybridization intensity were observed between the CD and control liver RNA samples (Fig. 4).

**DISCUSSION**

We have confirmed that the liver content of a major erythromycin demethylase (P-450p) is markedly reduced in a commonly used rat model for chronic liver disease (Figs. 1 and 3). Contrary to our initial hypothesis, however, the production of CO₂ from erythromycin in these rats was not more significantly impaired than that from aminopyrine (Table 1). At least in part, this appears to result from relative preservation of erythromycin demethylase activity in the diseased liver (Table 2). Our observations are supported by those of Murray et al. (18), who reported that another catalytic activity characteristic of P-450III cytochromes (testosterone 6β-hydroxylase activity) also appeared to be relatively preserved in CD rat liver microsomes.

We believe that our immunochemical analysis of the CD liver microsomes provides an explanation for these observations. Although the concentration of P-450p was dramatically reduced in the CD rat liver microsomes, the concentration of one P-450III–related protein appeared to be similar in control and CD rat liver microsomes (Fig. 2). Moreover, the microsomal concentration of another P-450III–related protein was markedly increased in the CD liver microsomes (Figs. 1 and 3). Because the non-P-450p proteins identified by the monoclonal antibodies have not yet been purified, it has not been possible to directly demonstrate that these proteins are capable of erythromycin demethylation. Nonetheless, most erythromycin demethylase activity in CD liver microsomes (essentially lacking P-450p) could be inhibited by preincubation with anti-P-450p IgG, indicating that most of this activity was due to P-450p–related proteins. Our observations suggest that the loss of erythromycin demethylase activity attributable to P-450p is compensated for by the induction of another P-450III cytochrome capable of catalyzing this reaction.

It should be pointed out that the reciprocal changes in the concentrations of P-450p and the second reactive protein (Fig. 3) might also be compatible with P-450p degradation to a slightly smaller protein in the CD liver. However, this seems unlikely for several reasons. First, both the unabsorbed and absorbed polyclonal antibody preparations appeared to have higher affinity for P-450p than for the second protein, whereas the reverse appeared to be the case with the monoclonal antibody 1G8 (Fig. 1). This suggests that P-450p contains multiple epitopes not present in the protein induced in the CD liver. Second, the CD rat hepatocytes were clearly capable of expressing P-450p when pretreated with dexamethasone (Fig. 1A). Finally, two attempts to purify the induced protein from our CD liver microsomes have failed using techniques similar to those used to purify P-450p (S. Wrighton, Personal communication, 1989).
These observations suggest that the protein induced in the CD liver is related to, but structurally distinct from, P-450p. We believe that this is the first evidence to date that some cytochromes P-450 may actually be induced by chronic liver disease or prolonged nutritional deficiency.

Several conclusions can be drawn from our data about the mechanisms involved in the differential expression of P-450III proteins we observed in the CD animals. First, because the reciprocal changes in the specific concentrations of P-450p and the related protein were only evident after weeks on the diet (Fig. 3), they were not acute effects of the nutritional deficiency. However, since the CD rats gained weight much more slowly than did the controls, it is possible that starvation alone may account for the differential regulation we observed. Second, it is unlikely that the changes we observed reflect “feminization of the liver” as has been previously postulated (18). The CD rats appeared to maintain expression of a male-specific P-450III-related protein, and a related protein that is female-specific was never detected (Fig. 2). In addition, the P-450III-related protein that was induced in the CD liver was indistinguishable from a protein that is male specific in control animals (Figs. 1C and 3). Finally, the male-specific mRNA PCN-2, was readily detected in each CD rat liver (Fig. 4).

Our observations also supply some additional information regarding the P-450III gene family. When uninduced male rat liver microsomes are subjected to SDS-PAGE, the 13-7-10 reactive protein (Fig. 2) exactly comigrates with the faster migrating 1G8 reactive protein (9). It had therefore not been previously possible to show that these were distinct proteins rather than a single protein sharing both epitopes (9). In the CD liver, however, the IG8-reactive protein was significantly induced, whereas the microsomal concentration of the 13-7-10 reactive protein was unchanged, indicating that the proteins are not identical. Unfortunately, we were unable to use our rats to indicate whether any of the three immunoreactive proteins reflected expression of either the PCN-1 or PCN-2 gene. The concentrations of PCN-1 or PCN-2 mRNA did not appear to correlate well with the specific content of any of the P-450p-related proteins in the livers analyzed (Fig. 4). However, regulation of these proteins in the CD rats may not be at the level of transcription.

It is tempting to speculate that the loss of P-450p may have physiological significance in these animals. Selvy (29) showed that rats pretreated with the prototypical inducer of P-450p, pregnenolone 16α-carbonitrile, were resistant to the liver toxicity of many drugs. This appeared to result from an increased ability to detoxify and/or eliminate potential toxins. Furthermore, others have shown that pregnenolone 16α-carbonitrile–treated rats are resistant to dimethylsulfamine-induced hepatomas (30). The selective loss of P-450p in the CD rat may therefore result in decreased detoxification ability, which may contribute to the progressive liver injury associated with the diet. On the other hand, some P-450III cytochromes appear to be involved in the bioactivation of various mycotoxins, including aflatoxin B (31, 32). Differential effects on regulation of P-450III cytochromes may therefore contribute to the higher incidence of spontaneous hepatomas observed in rats maintained on the deficient diet for more than 7 mo (33).

It has in fact been postulated that hepatomas observed in these rats result from enhanced susceptibility to aflatoxin present in trace amounts in the diet (34). Because the P-450III1 cytochrome has been widely implicated in carcinogenesis (14 and references therein), our incidental observations indicating that regulation of this enzyme did not appear to be altered in the CD rat liver may also be important.

Finally, it should not be concluded from our studies that the ERMBT will offer no advantage over the APBT in monitoring liver function in patients. Although P-450III cytochromes are highly conserved in mammals (7) and are major constitutive enzymes in human liver (4, 6), species differences in their regulation exist (7). Furthermore, the P-450III cytochromes have been characterized with regard to induction or inhibition by medications and are not induced by ethanol (6-14). The use of erythromycin therefore has theoretical advantages over aminopyrine where the responsible enzymes have yet to be identified. Clinical studies using the ERMBT are now underway.

REFERENCES
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