Suppression of cytochrome P450 reductase expression in hepatoma cells replicates the hepatic lipidosis observed in hepatic POR-null mice

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Running Title: Lipidosis in POR-suppressed hepatoma cells

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Text pages: 27
Tables: 1
Figures: 5
References: 23

Words

Abstract: 250
Introduction: 455
Discussion: 1,403

Abbreviations: POR, cytochrome P450 reductase; RT-PCR, reverse-transcriptase polymerase chain reaction; FBS, fetal bovine serum; PBS, phosphate-buffered saline; BSA, bovine serum albumin; CYP51, CYP51A1; LXR, liver X receptor; GC/MS, gas chromatography/mass spectrometry; CDCA, chenodeoxycholic acid, SREBP, sterol regulatory element binding protein.
ABSTRACT

Cytochrome P450 reductase (POR) is a microsomal electron transport protein essential to P450-mediated drug metabolism and sterol and bile acid synthesis. The conditional deletion of hepatic POR gene expression in mice results in a marked decrease in plasma cholesterol levels counterbalanced by the accumulation of triglycerides in lipid droplets in hepatocytes. To evaluate the role of cholesterol and bile acid synthesis in this hepatic lipidosis, as well as the possible role of lipid transport from peripheral tissues, we developed a stable, siRNA-mediated cell culture model for the suppression of POR. POR mRNA and protein expression were decreased by greater than 50% in McArdle-RH7777 rat hepatoma cells 10 days after transfection with a POR-siRNA expression plasmid, and POR expression was nearly completely extinguished by day 20. Immunofluorescent analysis revealed a marked accumulation of lipid droplets in cells by day 15, accompanied by a nearly two-fold increase in cellular triglyceride content, replicating the lipidosis seen in hepatic POR-null mouse liver. In contrast, suppression of CYP51A1 (lanosterol demethylase) did not result in lipid accumulation, indicating that loss of cholesterol synthesis is not the basis for this lipidosis. Indeed, addition of cholesterol to the medium appeared to augment the lipidosis in POR-suppressed cells, whereas removal of lipids from the medium reversed the lipidosis. Oxysterols did not accumulate in POR-suppressed cells, discounting a role for LXR in stimulating triglyceride synthesis, but addition of chenodeoxycholate significantly repressed lipid accumulation, suggesting that the absence of bile acids and loss of FXR stimulation lead to excessive triglyceride synthesis.
Cytochrome P450 reductase (POR) is the obligate electron donor to the cytochromes P450 found on the endoplasmic reticulum, which includes most P450s involved in drug metabolism, but also P450s essential to physiological processes such as cholesterol, steroid, and bile acid synthesis. Indeed, the genomic inactivation of POR in mice results in embryonic lethality by day 14 (Shen et al., 2002), and developmental defects associated with POR deletion appear to be due in large part to the loss of cholesterol synthesis (Otto et al., 2003; Schmidt et al., 2009). To explore the role of microsomal P450s in drug metabolism and circumvent the embryonic lethality associated genomic deletion of POR, the conditional, liver-specific deletion of P450 reductase was reported by two groups in 2003 (Gu et al., 2003; Henderson et al., 2003). Unexpectedly, these hepatic POR-null mice developed a severe hepatic lipidosis characterized by the accumulation of di- and triglycerides and monounsaturated fatty acids (Weng et al., 2005; Mutch et al., 2007), even while serum triglyceride and cholesterol levels were decreased. As POR was not known to be involved in triglyceride synthesis or secretion, the basis for the lipidosis was not readily apparent.

Disruption of POR activity would be expected to prevent cholesterol synthesis as well as its catabolism to bile acids. The net result in hepatic POR-null mice is a dramatic decrease in circulating cholesterol and bile acids. Because the liver serves as a depot for lipids from the peripheral tissues, it is unclear to what extent extrahepatic lipids contribute to the accumulation of triglycerides in hepatic POR-null mice. Weng et al.
(2005) found an increase in the expression of a gene responsible for fatty acid uptake (
$Cd36$) and a decrease in the expression of genes involved in fatty acid oxidation ($Cpt1a,
Ascl1$). Mutch et al. (2007) made similar observations, concluding that, although hepatic
fatty acid synthesis was neither increased nor decreased in POR-null liver, lipid secretion
was impaired and that this, coupled with continual uptake from the circulation, lead to
the lipidosis. Indeed, Finn et al. (2009) were able to show that placing these mice on a
fat-deficient diet completely prevented the hepatic lipidosis and the fall in circulating
cholesterol and triglycerides. But why the disruption of POR activity, manifested as a
block to cholesterol and bile acid synthesis, would result in this impaired secretion of
triglycerides was not elucidated. As a first step toward answering this question, we have
developed a cellular model of the hepatic POR-null mouse by using cultured hepatoma
cells and siRNA to suppress POR expression. This model appears to faithfully replicate
the lipidosis seen in whole animals and has allowed us to explore the role of
extracellular lipids, cholesterol, and bile acids in the accumulation of triglycerides in
these cells.
MATERIALS AND METHODS

Construction of a POR siRNA expression plasmid. siRNA sequences for the rat POR mRNA were identified using a commercial program (Integrated DNA Technologies, San Diego, CA). Three sequences were identified and each was synthesized by IDT, along with its complementary sequence, to generate “hairpin” formation oligonucleotides. BamHI and HindIII restriction sites were included at the ends to allow cloning into the pSilencer4.1-CMVneo siRNA expression vector (Applied Biosystems, Foster City, CA) after annealing the complementary pairs. Each plasmid was prepared and transfected into McA-RH7777 rat hepatoma cells and POR mRNA expression was tested after 2, 4, 6, 8, and 10 days by reverse-transcriptase polymerase chain amplification (RT-PCR). The plasmid yielding the best suppression was selected for further use; the siRNA sequence corresponds to nucleotides 2372-2390 in the 3’-nontranslated region of the rat POR mRNA. A plasmid containing a scrambled siRNA sequence served as the wild-type control.

Transfection and culture of hepatoma cells. McArdle-RH7777 (McA-RH7777, CRL-1601, ATCC) rat hepatoma cells (American Type Culture Collection, Manassas, VA) between passages 12-22 were cultured in six-well plates in Dulbecco’s Modified Eagle Medium supplemented with 1x penicillin-streptomycin mixture and 10% fetal bovine serum (FBS, Invitrogen, Carlsbad, CA) at 37°C under a humidified atmosphere of 5% CO₂. For plasmid transfection, best results were obtained by mixing the plasmid DNA with Fugene 6 transfection reagent (Roche Applied Science, Indianapolis, IN) according to the
manufacturer's instructions at a ratio of 1 µg DNA:5 µl reagent, allowing complexes to form for a minimum of 30 min at room temperature. The cell culture medium then was replaced with antibiotic-free medium containing 10% FBS and the transfection mixture was added drop-wise to the cells with gentle swirling. The medium was replaced after 24 h with standard medium; cells were split as necessary and assayed for expression beginning after 5 days. For cholesterol-supplemented medium, cholesterol was added dropwise to FBS in ethanol (5 mg/ml) with vortexing to give a final medium concentration of 50 µg/ml (Meyer et al., 1990). For the preparation of delipidated medium, FBS was mixed with an equal volume of diethyl ether with vigorous mixing, after which the ether was removed and the delipidated serum was sterile-filtered for use (Slater and Robertson, 1979).

siRNA suppression of CYP51. Four plasmids for the suppression of rat CYP51A1 (CYP51, lanosterol demethylase) were purchased (SureSilencing™ shRNA, SABiosciences, Frederick, MD) and two were tested for their ability to suppress CYP51 expression in McA-RH7777 cells, following the transfection protocol described above. A plasmid containing a scrambled siRNA sequence served as the negative control. Cells transfected with the 'yellow' plasmid were selected and maintained in the presence of 1500 µg/ml geneticin (Invitrogen).

Expression analysis by RT-PCR. Total RNA was isolated from cells grown to confluence in 6-well plates, with the use of TRIzol reagent (Invitrogen) and the concentration of RNA
was determined spectrophotometrically with the use of a Nanodrop instrument (ThermoFisher Scientific, Waltham, MA). RT-PCR was carried out with Superscript II Reverse Transcriptase (Invitrogen) and quantified on an Applied Biosystems 7000 instrument using gene-specific primer sets, as follows: POR, tctcacgaagacaccagtgc, cgtgttttgatcttgctga; CYP51, gtgccaaatgcagttttcct, tcagacagcgctcacaac. Dissociation curves were established for each PCR product. The detection and quantitation of nucleic acid levels was done by using the comparative Ct method (Livak and Schmittgen, 2001) with β-actin as the reference gene.

**Immunoblot analysis of protein expression.** Thirty micrograms of protein from cell lysates was fractionated by SDS-polyacrylamide gel electrophoresis on 8% gels and electroblotted to nitrocellulose (Bio-Rad, Hercules, CA). The membrane was blocked with 0.05% Tween-20 and 5% defatted milk for 1 h at room temperature and then incubated in this same buffer with rabbit polyclonal antibody to rat POR (Stressgen/Enzo Life Sciences, Plymouth Meeting, PA) or human CYP51 (Proteintech Group, Inc., Chicago, IL) overnight at 4°C with gentle shaking. The immunoblot was developed with a secondary antibody conjugated to horseradish peroxidase for 1 h at room temperature and the chemiluminescent image (Supersignal West Pico Chemiluminescent Substrate, Pierce/Thermo Scientific, Rockford, IL) captured and quantified by autoradiography on a Kodak Image Station. β-Actin served as a loading control, and was detected with a mouse monoclonal antibody (Sigma-Aldrich, St. Louis, MO).
**Immunofluorescence analysis.** Hepatoma cells transfected with an siRNA expression plasmid were plated on cover slips coated with collagen in 35 mm dishes at 0.5 x 10^6 cells per 1 ml in DMEM containing 5% FBS and 1% penicillin-streptomycin. At 5, 10, 15 and 20 days after transfection the cells were immunostained as follows: cover slip cultures were rinsed twice with ice-cold phosphate-buffered saline (PBS) and fixed by incubation with 4% paraformaldehyde for 30 min at room temperature. The cells were washed three times for 5 min with ice-cold PBS, treated with 0.1% Triton X 100 for 8 min at room temperature, washed with PBS, and quenched with 100 mM ammonium chloride for 20 min. The cells were washed with PBS and blocked in 1% bovine serum albumin (BSA) in PBS for 30 min, and incubated for 10 min with the Nile Red stain (Sigma-Aldrich) for lipids (1:5000). Stained cells were then washed and mounted on glass slides with use of ProLong antifade reagent (Invitrogen) containing DAPI nuclear stain for visualization on a Zeiss Axiovert 200M fluorescence microscope. For detection of POR expression, coverslips were incubated with the antibody to POR (1:100) described above in 1% BSA in PBS overnight, washed with PBS containing 0.1% BSA, and incubated in the dark with a secondary antibody conjugated to a fluorochrome (Alexa488 Fluor, Molecular Probes/Invitrogen) for 1 h. Images were captured at 400x unless otherwise indicated.

**GC-MS analysis of cellular sterols.** For oxysterol analysis, hepatoma cells cultured for 12 days in standard medium were homogenized at 4°C for 5 min in 1 ml of absolute ethanol.
containing 10 μg/ml butylated hydroxytoluene. The homogenates were subjected to centrifugation at 12,000g for 20 min, the supernatant set aside, and the pellets resuspended and re-extracted with sonication and centrifugation in a methylene chloride/methanol mixture (1:1, v/v, 1 ml) containing 10 μg/ml butylated hydroxytoluene. This supernatant fraction was combined with the ethanol extract and the solvents removed by evaporation under N2 gas. The extracted sterols were resuspended in 50 μl of methylene chloride/hexane (8:2, v:v) and derivatized with trimethylsilane. Mass spectra were acquired by the University of Kentucky Mass Spectrometry Facility on a Trace Gas Chromatograph with a DB5-ms (Agilent/J&W) 30 m x 0.25 mm (0.25 μm df) column coupled to a Thermo-Finnigan PolarisQ ion-trap mass spectrometer, using selective ion monitoring. The following sterols were run as standards and monitored in the cell extracts: 22β-hydroxycholesterol, 24,25-dihydrolanosterol, 25-hydroxycholesterol, and 27-hydroxycholesterol. For measurement of lanosterol in CYP51-suppressed cells, lipids were extracted twice with 2 ml of hexane, the solvent evaporated under N2 gas, and the samples derivatized by resuspension in 50 μl of bis (trimethylsilyl) trifluoroacetamide: pyridine (1:1).

Analysis of triglycerides. Cells were cultured and treated in triplicate in 96-well plates, and then were pooled and assayed in duplicate. Triglyceride content was determined with 30 μg cell lysate protein using the LabAssay Triglyceride Kit from Wako Pure Chemical Ind. (Richmond, VA).
RESULTS

Transfection of a plasmid that expresses siRNA complementary to rat cytochrome P450 reductase mRNA into McA-RH7777 rat hepatoma cells decreased POR mRNA in these cells by greater than 80% by day 5, and this suppression was maintained through 20 days of culture (Fig. 1A). POR protein expression, measured by immunoblot analysis at days 10 and 20, was decreased to 28 and 16% of control cell levels, respectively (Fig. 1B). Immunocytochemical analysis of these cells at 20 days post-transfection with an antibody to cytochrome P450 reductase confirmed the near complete loss of POR expression (Fig. 2). Staining of these cells with Nile Red stain for lipids demonstrated a diffuse increase in lipid content throughout the cell with numerous and prominent intracellular lipid vesicles of various sizes. Triglyceride content in POR-suppressed cells was increased at 5 and 10 days, and was statistically significant at 15 days (Table 1). This pattern of lipidosis closely resembles that observed in hepatocytes of hepatic POR-null mice (Gu et al., 2003; Henderson et al., 2003).

To determine if the increase in lipid content in POR-suppressed cells was due to the loss of cholesterol synthesis, CYP51 expression was suppressed in McA-RH7777 cells by siRNA. CYP51 uniquely catalyzes the demethylation of lanosterol in the cholesterol biosynthetic pathway and has no other known biochemical roles. Moreover, it requires cytochrome P450 reductase for activity, and thus its suppression blocks cholesterol synthesis at the same step as does POR suppression. CYP51 mRNA was decreased by 86% at 10 days (Fig 3A) and CYP51 protein was decreased by 79% at 15 days (Fig. 3B);
lanosterol content in these cells increased by up to 8-fold, similar to the 10-fold increase seen in POR-suppressed cells (data not shown). However, CYP51-suppressed cells had significantly lower triglyceride content than did control cells or POR-suppressed cells (Table 1) and displayed no evidence of lipidosis (Fig 3C), arguing that the lipidosis observed in POR-suppressed cells does not result solely from the loss of cholesterol synthesis.

The conclusion that loss of cholesterol synthesis alone is not responsible for the lipidosis is further supported by the observation that addition of cholesterol to the culture medium appeared to augment the accumulation of lipids in POR-suppressed cells (Fig 3D). As cholesterol addition had no effect on the lipid content of control hepatoma cells, this observation raised the possibility that the inability of POR-suppressed cells to eliminate cholesterol might lead to the generation of oxysterols in these cells. Oxysterols serve as potent ligands for the liver X receptor (LXR), which stimulates triglyceride synthesis, accumulation, and hepatic steatosis (Kalaany and Mangelsdorf, 2006; Moya et al., 2010). To test this hypothesis, the abundance of three oxysterols whose formation does not require cytochrome P450 reductase (22β-hydroxycholesterol, 25-hydroxycholesterol, and 27-hydroxycholesterol) was determined by gas chromatography/mass spectrometry (GC/MS) in control and POR-suppressed hepatoma cells. As shown in Fig. 4, these three oxysterols were less abundant in POR-suppressed cells than in control cells, indicating that the loss of cholesterol catabolism does not
result in the accumulation of cholesterol oxidation products that might otherwise lead
to an LXR-mediated stimulation of triglyceride accumulation.

The catabolism of cholesterol to bile acids is dependent upon multiple P450 oxidations,
several of which, including the regulatory step, are dependent on cytochrome P450
reductase. As bile acids act through the farnesoid X receptor (FXR) to suppress the
synthesis of fatty acids, triglycerides, and bile acids (Zhang and Edwards, 2008), the loss
of bile acids in POR-suppressed cells might be expected to de-repress the synthesis of
fatty acids and lead to the accumulation of triglycerides, as has been found in FXR-null
mice (Sinal et al., 2000). To test this hypothesis, chenodeoxycholic acid (CDCA), a strong
FXR ligand, was added to POR-suppressed cells. As shown in Fig. 5A, the addition of 50
µM CDCA largely prevented the accumulation of lipids in these cells and decreased
triglyceride content below that of control and POR-suppressed cells (Table 1).

Although the results obtained with bile acid supplementation suggest that triglyceride
synthesis is up-regulated in POR-suppressed cells, studies with hepatic POR-null mice
indicated that genes involved in fatty acid and triglyceride synthesis are not up-
regulated in the livers of these mice (Wang et al., 2005; Weng et al., 2005). These
studies instead suggested that the accumulating triglycerides are derived largely from
the circulation. To test the role of lipid uptake in POR-suppressed hepatoma cells, lipids
were extracted from FBS and cells were cultured for up to 15 days in medium prepared
with this delipidated serum. POR-suppressed cells initially accumulated lipids to the
same extent as cells cultured in normal medium, and triglyceride levels were not statistically different from POR-suppressed or control hepatoma cells cultured in normal medium after up to 10 days of culture (Table 1). By 15 days, however, triglyceride levels were significantly elevated in POR-suppressed cells cultured in normal medium, but were greatly decreased in POR-suppressed cells maintained in delipidated serum. Lipid staining of these cells demonstrated marked lipidosis in cells cultured in normal medium for 15 days and a notable lack of lipids in cells cultured in the delipidated serum (Fig. 5B). Although the viability of POR-suppressed cells was limited to 15 days in delipidated serum, these results suggest that lipid uptake contributes to the lipidosis.
DISCUSSION

The present studies were undertaken to determine if suppression of cytochrome P450 reductase expression in an in vitro, hepatoma cell culture model would replicate the lipidosis observed in hepatic POR-null mice. The intent was to remove the whole animal physiological and compensatory components that might contribute to, or diminish, the effect of loss of cytochrome P450 reductase on lipid homeostasis in hepatocytes. Ultimately, the goal was, and is, to define the mechanism by which loss of this enzyme leads to triglyceride lipidosis; this goal is all the more interesting given that cytochrome P450 reductase is not known to play a role in triglyceride uptake, synthesis, catabolism, or secretion, and all the more challenging given the several pathways in which cytochrome P450 reductase participates in lipid metabolism in the cell.

siRNA-mediated suppression of cytochrome P450 reductase in hepatoma cells was effective in generating a lipidosis that closely resembles that seen in hepatic POR-null mice. In this cell culture model, POR mRNA was decreased to 20% of control levels at 5 days post-transfection, POR protein was similarly decreased by day 10, and lipid accumulation was clearly evident by day 15. These changes match the results obtained by Finn et al. (2009) with an inducible hepatic POR-null mouse in which hepatic triglyceride accumulation was evident 5 days after loss of POR expression. Notably, plasma cholesterol levels fell in concert with POR protein levels in these mice, whereas plasma and liver triglyceride levels took an additional 5 days to reach a new equilibrium.
Their results, and ours, suggest that changes in triglycerides are secondary to changes in cholesterol synthesis or catabolism.

To address the possibility that the lipidosis resulted from the loss of cholesterol synthesis in POR-suppressed cells (or from the fall in plasma cholesterol levels in hepatic POR-null mice), we supplemented the medium with cholesterol, but this appeared to augment, rather than attenuate the lipidosis. As hepatic cholesterol levels are normal in hepatic POR-null mice (Finn et al., 2009), it seems unlikely that a cholesterol deficit leads to the triglyceride accumulation; the cells and liver appear able to compensate for the loss of cholesterol synthesis by increasing cholesterol uptake from the medium or circulation. This conclusion is further supported by our observation that CYP51 suppression does not lead to lipidosis and, indeed, lowers triglyceride levels. Suppression of this enzyme is complementary and equivalent to POR suppression in the cholesterologenic pathway but, unlike POR suppression, does not affect other biochemical pathways. This provides strong evidence that loss of cholesterol synthesis per se is not sufficient to cause triglyceride accumulation.

Oxysterols are well-established agonists for the liver X receptor, a potent transcriptional inducer of genes involved in fatty acid synthesis. LXR agonists readily induce hepatic triglyceride accumulation (Kalaany and Mangelsdorf, 2006). However, studies with hepatic POR-null mice do not show a strong transcriptional response characteristic of LXR stimulation (Wang et al., 2005; Weng et al., 2005) indicating that oxysterols are
unlikely to be elevated in POR-null mice despite the inability of these liver cells to catabolize excess cholesterol to bile acids. Our cell culture studies support this conclusion, as three of the most common oxysterols, none of which are dependent on cytochrome P450 reductase for their synthesis, were less abundant in POR-suppressed cells than in control cells. There is, however, at least one additional sterol that could contribute to LXR stimulation: 24,25-dihydrolanosterol. Dihydrolanosterol was recently shown to be a moderate LXR agonist (Zhu et al., 2010); this sterol is formed from lanosterol when CYP51 is inhibited or absent. Interestingly, Zhu et al. (2010) showed that inhibition of CYP51 in 3T3L1 fibroblasts results in a triglyceride lipidosis, presumably by activation of LXR. Dihydrolanosterol is not abundant in POR-suppressed or CYP51-suppressed McA-RH7777 hepatoma cells (our data), but is abundant in the livers and hepatocytes of hepatic POR-null mice (Li and Porter, 2007), and may generate some level of LXR activation in these animals. But overall, the evidence from both POR-null mice and POR-suppressed cells does not support a significant role for oxysterols and other LXR agonists in the triglyceride lipidosis that follows loss of POR expression.

FXR is a bile acid receptor that down-regulates bile acid synthesis in the liver in the presence of excess bile acids. In the absence of FXR signaling triglyceride synthesis is greatly increased through de-repression of SREBP-1c signaling, which increases transcription of several lipogenic genes. Addition of a bile acid, CDCA, to POR-suppressed cells largely prevented the lipidosis and decreased triglycerides to below normal levels, arguing that FXR signaling is absent or greatly decreased in these cells;
this would be consistent with an inability of these cells to synthesize bile acids in the absence of cytochrome P450 reductase. FXR signaling also appears to be decreased in the livers of hepatic POR-null mice, consistent with a 90% reduction in the bile acid pool in these animals (Henderson et al., 2003). Expression of CYP7A1, a gene down-regulated by FXR, is increased in the livers of these mice by at least 3.5-fold (Wang et al., 2005; Weng et al., 2005), and similar increases in other genes negatively regulated by FXR are also seen. However, genes involved in fatty acid and triglyceride synthesis in the livers of these mice are not significantly up-regulated, leading the investigators in these studies to conclude that the lipidosis is not the result of increased triglyceride synthesis, but rather due to increased triglyceride uptake and decreased catabolism and secretion.

We tested the role of lipid uptake from the medium in our cell culture model by using delipidated medium. While in normal medium triglyceride content trended upward in POR-suppressed cells, reaching significance at 15 days, triglyceride content in cells grown in delipidated medium trended downward, also reaching significance at 15 days. At 15 days lipid vesicles were clearly absent in cells grown in delipidated medium, demonstrating that extracellular lipids are required for lipid content to increase in POR-suppressed cells. Because we did not measure fatty acid or triglyceride synthesis or uptake in these cells, we cannot determine to what extent synthesis and uptake contribute to the lipidosis, but it would appear that both play a role in this system. We do note that POR-suppressed cells grown in delipidated medium do not survive to 20 days, whereas control cells are still dividing at 20 days despite the absence of
extracellular lipids. Thus, an external source of lipid is essential to the survival of POR-suppressed cells; we suspect that this required lipid is cholesterol. Interestingly, POR-suppressed cells supplemented with CDCA also die after approximately 15 days, whereas control cells remain viable in this medium; because cholesterol and the normal complement of serum lipids are present in this medium the toxicity must not be due to the lack of a lipid but rather to presence of CDCA. Why POR-suppressed cells are sensitive to this concentration of a bile acid, while control cells are not, remains to be determined.

McA-RH7777 rat hepatoma cells retain a number of characteristics of liver parenchymal cells and are widely used in lipid metabolism studies, including cholesterol and triglyceride synthesis (Singh and Porter, 2006; Ko et al., 2009) and lipoprotein secretion (Rasouli et al., 2004). They more closely resemble the intact liver with regard to SREBP expression than do the also widely used HepG2 cell line (DeBose-Boyd et al., 2001), and thus are very attractive for studies on the regulation of hepatic lipid synthesis. They do not appear to be efficient at bile acid synthesis (Labonte et al., 2000), although we have not tested this. Further studies on the expression level of regulatory and catalytic genes in cholesterol, fatty acid, and bile acid synthesis in these cells would clearly help elucidate which pathways are being activated in the absence of POR expression in this defined in vitro system.
We have established a hepatoma cell culture system that replicates the hepatic lipidosis observed in hepatic POR-null mice. Studies with this model suggest that a loss of bile acid synthesis leading to enhanced triglyceride synthesis, coupled with uptake and perhaps altered metabolism of triglycerides and fatty acids, is responsible for the lipidosis observed in these cells. The extent to which these complementary pathways contribute to the lipidosis remains to be determined, and potential mechanisms by which triglyceride metabolism and secretion are altered in these cells have not been defined. Further studies with this model, as well as with hepatic POR-null mice, will be needed to delineate fully the interplay between cytochrome P450 reductase and triglyceride metabolism.
ACKNOWLEDGMENT

We thank Ms. Melissa Douglas for assistance with these studies. This work was supported by the National Institutes of Health National Center for Complementary and Alternative Medicine [Grants AT003488 and AT005235].

AUTHORSHIP CONTRIBUTION

Participated in research design: Porter, Banerjee, Stolarczyk, Zou

Conducted experiments: Banerjee, Stolarczyk, Zou

Performed data analysis: Porter, Banerjee, Stolarczyk, Zou

Wrote or contributed to the writing of the manuscript: Porter, Banerjee, Stolarczyk
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FIGURE LEGENDS

Fig. 1. Suppression of POR mRNA and protein in hepatoma cells. Hepatoma cells were transfected with the POR siRNA expression plasmid or a control vector (scrambled) and POR mRNA (A) and protein (B) expression were measured at various time-points by RT-PCR and immunoblot analysis, respectively. An immunoblot image is shown at the bottom corresponding to the time-points in B. Values represent the mean and standard deviation of two determinations done in duplicate; values that are statistically different from the corresponding control are indicated by asterisks, as determined by one-way ANOVA with Bonferroni’s Multiple Comparisons post-hoc test, $p < 0.05$. CoD5, Control cells, day 5; ExD5, Experimental (POR-suppressed) cells, day 5.

Fig. 2. Immunofluorescence analysis of POR siRNA-transfected hepatoma cells. Hepatoma cells were transfected with the POR siRNA expression plasmid or a control vector and cytochrome P450 reductase expression in these cells was visualized with a POR antibody followed by a fluorochrome-coupled secondary antibody. Lipid content was assessed by staining with Nile Red; nuclei are stained with DAPI nuclear stain (blue). Cells were stained 20 days after transfection; image magnification is 630x.

Fig. 3. Lack of cholesterol does not lead to lipid accumulation. Hepatoma cells were transfected with a CYP51 siRNA expression plasmid or a control vector (scrambled) and CYP51 mRNA (A) and protein (B) expression were measured at various time-points by RT-PCR and immunoblot analysis, respectively. An immunoblot image is shown at the
bottom corresponding to the time-points in B. Lipid accumulation in CYP51-suppressed cells is shown in (C) with Nile Red staining of cells after 10 and 15 days; the effect of cholesterol supplementation on POR-suppressed cells is shown in (D). Cells were cultured in standard medium supplemented from day 2 post-transfection with 50 µg/ml of cholesterol and stained with Nile Red lipid stain at days 10 and 15. Values in panels A and B represent the mean and standard error, n = 3, of duplicate determinations; values that are statistically different from the control group (CoD2, A) or the corresponding control (B) are indicated by asterisks, as determined by one-way ANOVA with Dunnett’s Multiple Comparisons post-hoc test (A) or Bonferroni’s Multiple Comparisons post-hoc test (B), p < 0.05. CoD5, Control cells, day 5; ExD5, Experimental (CYP51-suppressed) cells, day 5.

Fig. 4. Oxysterols do not accumulate in POR-suppressed cells. Control and POR-suppressed hepatoma cells were grown in normal medium for 12 days, after which lipids were extracted and GC/MS with selective ion monitoring was used to identify 22β-hydroxycholesterol (eluting at ~27.3 min) and 27-hydroxycholesterol (31.8 min). 25-Hydroxycholesterol could not be detected in either lipid sample (uppermost baseline tracing); lanosterol eluted at 28.25 min, and 24,25-dihydrolanosterol at 27.37 min.

Fig. 5. Bile acid supplementation or depletion of serum lipids prevents lipid accumulation. In (A) POR-suppressed hepatoma cells were cultured for 15 days in normal medium (left set of images) or medium supplemented with 50 µM CDCA (right
set of images); in (B) cells were cultured in normal medium (left set of images) or delipidated medium (right set of images) for 5, 10, or 15 days. All cells were stained for lipids with Nile Red.
# TABLE 1

## Triglyceride Content of Hepatoma Cells

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>POR siRNA</th>
<th>POR siRNA DLPS</th>
<th>POR siRNA CDCA</th>
<th>CYP51 siRNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 days</td>
<td>132.9 ± 19.8</td>
<td>184.8 ± 26.2</td>
<td>139.5 ± 23.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 days</td>
<td>143.8 ± 23.1</td>
<td>184.8 ± 30.8</td>
<td>125.3 ± 21.3</td>
<td>62.5 ± 11.6</td>
<td>*</td>
</tr>
<tr>
<td>15 days</td>
<td>154.8 ± 34.1</td>
<td>247.0 ± 39.6 *</td>
<td>72.9 ± 12.6 *</td>
<td>59.7 ± 9.3 *</td>
<td>34.1 ± 5.0 *</td>
</tr>
</tbody>
</table>

Means ± standard deviation, n = 2, in mg/mg cell lysate protein. DLPS, delipidated serum.

*, p < 0.05, Bonferroni’s Multiple Comparison Test with comparison to the control value for that time-point.
Figure 3

A

Fold difference in CYP51 mRNA relative to β-actin

B

CYP51 protein (relative values)

C

Control day 10  CYP51 siRNA day 10

Control day 15  CYP51 siRNA day 15

D

Control day 10  POR siRNA day 10

Control day 15  POR siRNA day 15

siRNA: (scrambled) — CYP51

siRNA: (scrambled) — CYP51

CYP51-suppressed

POR-suppressed
Figure 5

A  POR-suppressed cells

B  POR-suppressed cells

Normal medium, day 5

Delipidated medium, day 5

Normal medium, day 10

Delipidated medium, day 10

Normal medium, day 15

Delipidated medium, day 15

Normal medium

CDCA-supplemented