Competitive Inhibition of Bile Acid Synthesis by Endogenous Cholestanol and Sitosterol in Sitosterolemia with Xanthomatosis

Department of Medicine and the Sammy Davis, Jr. National Liver Institute, University of Medicine and Dentistry of New Jersey-New Jersey Medical School, Newark, New Jersey 07103; and Veterans Administration Medical Center, East Orange, New Jersey 07019

Abstract

The 7α-hydroxylation of two cholesterol analogues, sitosterol and cholestanol, and their effect on the 7α-hydroxylation of cholesterol were measured in rat and human hepatic microsomes. In untreated rat liver microsomes, the 7α-hydroxylation of cholesterol was higher than that of cholestanol (1.4-fold) and sitosterol (30-fold). After removal of endogenous sterols from the microsomes by acetone treatment, the 7α-hydroxylation of cholesterol was similar to that of cholestanol and only fourfold higher than that of sitosterol. Cholestanol and sitosterol competitively inhibited cholesterol 7α-hydroxylase in both rat and human liver microsomes, with cholestanol the more potent inhibitor. Patients with sitosterolemia with xanthomatosis, who have elevated microsomal cholestanol and sitosterol, showed reduced cholesterol 7α-hydroxylase activity relative to the activity in control subjects (13.9 and 14.7 vs. 20.3±0.9 pmol/nmol P<450 pmol per min, P<0.01). Enzyme activity in these patients was 40% higher when measured in microsomes from which competing sterols had been removed. Ileal bypass surgery in one sitosterolemic patient decreased plasma cholestanol and sitosterol concentrations and resulted in a 30% increase in hepatic microsomal cholesterol 7α-hydroxylase activity. Cholesterol 7α-hydroxylase appears to have a specific apolar binding site for the side chain of cholesterol and is affected by the presence of cholestanol and sitosterol in the microsomal substrate pool. Reduced bile acid synthesis in sitosterolemia with xanthomatosis may be related to the inhibition of cholesterol 7α-hydroxylase activity by endogenous cholesterol analogues.

Introduction

According to current information, the main pathway for the degradation of cholesterol1 in mammals is its conversion into bile acids. The major rate-controlling step in bile acid synthesis is the 7α-hydroxylation of cholesterol catalyzed by the microsomal enzyme, cholesterol 7α-hydroxylase (EC 1.14.13.17). Although newly synthesized cholesterol is the preferred substrate for this enzyme, other hepatic microsomal cholesterol pools can be used for the transformation to bile acids. About 300–500 mg of bile acids are formed daily in humans under normal conditions but, when stimulated, bile acid synthesis can be increased two to threefold. Up to 1 g of cholesterol thus can be 7α-hydroxylated and transformed into bile acids daily via this pathway.

For the past few years, bile acid synthesis has been measured by assessing the activity of hepatic microsomal cholesterol 7α-hydroxylase. The most commonly used assay is the isotope incorporation method, in which radioactive cholesterol is incorporated into the microsomal cholesterol pool and the formation of radioactive 7α-hydroxycholesterol is determined (1). Results obtained by this method may be confounded by the endogenous pool of microsomal sterols, especially when the size and composition of this pool vary. Various cholesterol analogues accompany cholesterol in virtually every mammalian tissue and may affect 7α-hydroxycholesterol formation (2, 3). Significant amounts of cholestanol are present in the liver of patients with the rare, inherited lipid storage disease cerebrotendinous xanthomatosis (4), and large amounts of plant sterols, in addition to cholestanol, are deposited in the liver of patients with sitosterolemia with xanthomatosis (5).

This study describes the 7α-hydroxylation of sitosterol and cholestanol and their effect on the 7α-hydroxylation of cholesterol in human and rat liver microsomal preparations in vitro. Enzyme activities were measured in the absence of endogenous microsomal sterols by a previously validated method that involves the extraction of endogenous sterols by acetone (1, 6), and compared with those obtained in untreated microsomes. The effect of the endogenous sterol substrate pool on cholesterol 7α-hydroxylase activity in patients with sitosterolemia with xanthomatosis was also evaluated.

Methods

Human subjects: Six control subjects (aged 16–54) and two sisters with sitosterolemia with xanthomatosis (aged 24 and 26) were included in the study. Of the six control specimens, four were obtained from the “Liver Tissue Procurement and Distribution System” (National Insti-
tutes of Health contract NO1-DK-62274, University of Minnesota Hospital, Minneapolis, MN) and two were liver biopsies obtained for diagnostic histologic evaluation from normolipemic subjects with normal liver morphology and function tests. Complete clinical descriptions of the sitosterolemic subjects have been published previously (7). Liver biopsies were obtained from two untreated sitosterolemic patients and repeated in one patient 9 mo after ileal bypass surgery. The experimental protocol was approved by the Human Study Committees of the Veterans Administration Medical Center, East Orange, NJ and the University of Medicine and Dentistry of New Jersey Medical School. Informed consent was obtained from each subject.

Animals. Male Sprague-Dawley rats (Taconic Farms, Germantown, NY) weighing 100–150 g were fed ad lib. ground rat chow (Ralston-Purina Co., St. Louis, MO) containing 2.5% cholestanol for 7 d. The rats were decapitated and their livers were removed immediately and chilled on ice. The animal protocol was approved by the Animal Studies Committee at the University of Medicine and Dentistry of New Jersey.

Purification and labeling of substrates. [4-14C]Cholesterol (New England Nuclear, Boston, MA) was diluted with unlabeled cholesterol (Supelco, Inc., Bellefonte, PA) to a specific activity of 5 × 10^4 dpm/μmol and purified by column chromatography on an AgNO₃-silicic acid column (2). The purified radioactive cholesterol contained < 0.06% 7α-hydroxycholesterol, 0.01% 7β-hydroxycholesterol, and 0.04% cholestanol as determined by thin-layer chromatography (TLC) on an AgNO₃-impregnated silica gel G plate (8) and by scintillation counting (Intertechnique SL 4000, Fairfield, NJ). [3α-3H]-Cholesterol (specific activity 1.4 × 10^7 dpm/μmol) was prepared by reduction of 3 mg 5α-cholest-3-one with 1 mg NaBH₄, in 0.5 ml methanol for 2 h. The product was isolated by preparative TLC on silica gel G plates (Brinkmann Instruments, Westbury, NY) using chloroform/acetone (99:1 vol/vol). The major spot, R₀/0.5, was eluted with chloroform/methanol (90:10 vol/vol) and was found to be > 99% pure by TLC and zonal radioactivity scanning. It was further purified by chromatography on a AgNO₃-silicic acid column (2). The purified product contained < 0.04% cholesterol, 0.04% 7α-hydroxycholest-4-en-3-one and 0.01% 7β-hydroxycholesterol as seen by TLC-scintillation counting. [4-14C]Sitosterol (Amersham Corp., Arlington Heights, IL) was diluted with unlabeled sitosterol (Supelco, Inc.) to a specific activity of 3.6 × 10^4 dpm/μmol and was purified from its oxidation products by column chromatography as described for cholesterol (2). It was further purified by HPLC on a liquid chromatograph (ALC 201; Waters Associates, Milford, MA) with a refractive index detector (model 401) and a radial-pack μBondapak C₁₈ reversed-phase column (10 μm particle size). The mobile phase consisted of methanol/chloroform/water (90:2:10 vol/vol/vol), at a flow rate of 2 ml/min (operating pressure 2,000 psi). The pure product was eluted at 24.8 ml (Table I) and contained < 0.05% 7α-hydroxyisotestoster, 0.02% 7β-hydroxyisotestoster (as determined by TLC-scintillation counting), 0.03% campesterol, and 0.02% stigmasterol (as determined by HPLC-scintillation counting).

Enzymes, cofactors, and reagents. NADPH and DTT were purchased from Calbiochem-Behring Corporation (La Jolla, CA); EDTA, Triton X-100, NaF, deoxycholate (DCA), and diiserylphosphatidylcholine (DLPC) from Sigma Chemical Co. (St. Louis, MO); [3H]7α-hydroxycholesterol (50 mCi/mmol) from New England Nuclear. Cholestyramine was a gift from Bristol-Myers Co. (Evansville, IN). 7-Ketocholesterol, 7α-, and 7β-hydroxycholesterol were synthesized as described by Fieser et al. (9). 7-Ketocholestanol, 7α-, and 7β-hydroxycholestanol were synthesized as described by Wintersteiner and Moore (10). 7-Ketosisterol, 7α- and 7β-hydroxyisotestoster were synthesized by oxidation of isomer acetate with tertiary butyl perbenzoate as described by Starka (11). NADPH-cytochrome P-450 reductase was purified from rat liver microsomes according to the method of Yasukochi and Masters (12).

Sterol determination in hepatic microsomes. Known aliquots of untreated or acetone-treated microsomal suspensions were refluxed for 3 h with 25% KOH in 95% ethanol (wt/vol). Sterols were extracted with hexane (6) and analyzed by capillary GLC (13) as trimethylsilyl (TMSi) ethers derivatives on a gas chromatograph (model 5890; Hewlett-Packard Co., Palo Alto, CA) equipped with a flame ionization detector and fitted with an open tubular fused silica column (0.32 mm × 26 m) that was internally coated with a 0.21-μm film of CP Wax 52CB (Chrompack, Inc., Bridgewater, NJ). The following operating conditions were used: injection temperature 275°C; column temperature (isothermal) 210°C; flame ionization detector temperature 295°C; helium carrier gas flow, 1.0 ml/min, and split ratio, 10:1.

Preparation and assay of cholesterol 7α-hydroxylase. Rat livers were excised at ~ 10 a.m. to minimize diurnal variations in bile acid synthesis, and all subsequent operations were carried out at 4°C. Microsomes were isolated and washed as described previously (1). The final pellet was suspended in a buffer (0.1 M potassium phosphate buffer, 20% glycerol, 1 mM EDTA, 5 mM DTT, and 50 mM NaF; pH 7.4) at a protein concentration, as determined by Lowry et al. (14), of ~30 mg protein/ml and stored at ~80°C until assayed. To remove endogenous sterols, 1 ml of the microsomal suspension was added dropwise, with stirring, to 40 ml acetone at ~10°C. When the suspension had settled, the precipitate was collected by centrifugation at 1,000 g for 5 min, washed, dried, ground, and stored as described previously (6). Before use, an aliquot of a known weight of the acetone powder (also referred to as acetone-treated microsomes) was homogenized in the above buffer. The cytochrome P-450 content of the untreated and acetone-treated microsomes was determined from the CO difference spectrum between 450 and 490 nm of the reduced sample using the extinction coefficient of 91 mM⁻¹·cm⁻¹ (15). These acetone-treated microsomal preparations contained 0.2–0.7 nmol P-450 and < 0.4 nmol sterols/mg protein. Acetone treatment of the microsomes removed 90% of the phospholipids as assayed by the method of Chen et al. (16) and 50% of the NADPH-cytochrome P-450 reductase activity as assayed by the method of Yasukochi and Masters (12).

In standard incubations with untreated microsomes, aliquots (0.1–0.5 nmol P-450) were preincubated for 2 min at 37°C in a final volume of 0.5 ml buffer (100 mM potassium phosphate, 0.1 mM EDTA, and 1 mM DTT, pH 7.4) containing 100 nmol [4-14C]-cholesterol solubilized with 0.15% Triton X-100. The reaction was stopped...
initiated by the addition of 0.6 μmol NADPH. Incubations were carried out with shaking for 10 min at 37°C, and were terminated by the addition of 15–20 vol of methylene chloride/ethanol (5:1 vol/vol). Zero time controls were run with each experiment. The 7α-hydroxycholesterol formed was extracted, acetylated with [3H]acetic anhydride, and separated by TLC on silica gel G (Analtech, Inc., Newark, DE) as previously described (17). The acetylation step is needed to eliminate the underestimation of product formation that is due to the dilution of the labeled cholesterol by the endogenous cholesterol pool that also serves as substrate in the untreated microsomes. [3H]-Cholestanol and [4-14C]sitosterol were used as substrates only with rat liver microsomes and their respective 7α-hydroxylated products were separated by TLC on silica gel G (Analtech, Inc.) without prior acetylation. The acetylation step was not necessary because endogenous cholesterol and sitosterol were not present in the rat liver microsomes. The TLC plates were developed with diethyl ether and the pertinent spots were visualized by spraying with 3.5% phosphomolybdic acid in isopropanol (6). The observed Rf values were: cholesterol, cholestanol, or sitosterol, 0.88; 7α-hydroxycholesterol or 7α-hydroxysterol, 0.40; 7β-hydroxycholesterol or 7β-hydroxysterol, 0.54; 7α-hydroxycholesterol, 0.55; and 7β-hydroxycholesterol, 0.46. The radioactivity of the pertinent spots was determined in a liquid scintillation counter.

When acetone-treated microsomes were used, the same assay conditions as described above were used except that 0.025 mg DLPC, solubilized in 0.12 mM Na-DCA, and 2 units of NADPH-cytochrome P-450 reductase were added to the incubation mixture to replace membrane phospholipids and NADPH-cytochrome P-450 reductase lost in the preparation of acetone-treated microsomes. The 7α-hydroxylated products were separated on silica gel G plates without prior acetylation because of the absence of endogenous sterols. The cholesterol 7α-hydroxylase activity in acetone-treated microsomes, assayed in this reconstituted system, was comparable to that of the untreated microsomes and has previously been shown to be similarly affected by bile acid malabsorption (cholestyramine treated) and by cholic acid feeding (6). We have noted that cytochrome P-420, which is devoid of enzyme activity, is formed in various amounts during the preparation of the acetone powder and if included in the calculations (e.g., when enzyme activity is expressed per milligram protein) it could lead to an underestimation of cholesterol 7α-hydroxylase activity. All results are expressed in picomoles of product formed per nanomole microsomal cytochrome P-450 per minute.

**Competitive inhibition assay.** To determine the effect of cholestanol and sitosterol on cholesterol 7α-hydroxylase activity in both rat and human acetone-treated microsomes, increasing amounts of unlabeled cholestanol (5–300 μM) or sitosterol (25–300 μM) were added to the assay systems together with the labeled cholesterol substrate. The concentrations of endogenous cholestanol and sitosterol in hepatic microsomes from patients with sitosterolemia and xanthomatosis were at least 5 and 25 μM, respectively, under the assay conditions used. When the type of inhibition was determined, varying concentrations of [4-14C]cholesterol were used with rat acetone-treated microsomes and its conversion to 7α-hydroxycholesterol was measured in the presence of two different concentrations of unlabeled cholestanol or sitosterol (50 and 100 μM for cholestanol, 100 and 200 μM for sitosterol). Double reciprocal plots were used to determine the apparent Km values in the presence and absence of inhibitory sterols.

Statistical significance of the data was determined by the unpaired t test. In the comparison of sitosterolemic cholesterol 7α-hydroxylase activities with control values, a normal distribution for the control data was assumed and a 0.01 confidence limit was calculated (18).

**Results**

7α-Hydroxylation of cholesterol, cholestanol, and sitosterol. The formation of 7α-hydroxycholesterol (2, 19) and 7α-hydroxycholesterol (2) by untreated hepatic microsomes have been reported previously and the same identification procedures of reaction products were used with the acetone-treated microsomes. To obtain enough material to carry out the identification of 7α-hydroxycholesterol, the products from large-scale (300–500-fold) incubation mixtures were combined and purified by TLC as described in Methods. The fraction corresponding to 7α-hydroxycholesterol (Rf = 0.4) was subjected to GLC analysis after trimethylsilylation, and was found to have an identical retention time to synthetic 7α-hydroxycholesterol (3). Its specific radioactivity after dilution with authentic unlabeled 7α-hydroxycholesterol (1,100 dpm/μmol) remained constant after crystallization from acetone/water and methanol/water.

Optimal assay conditions for the 7α-hydroxylation of cholestanol and sitosterol were determined and compared with those of cholesterol, using acetone-treated microsomal preparations that contained < 0.4 nmol endogenous sterols per milligram protein (Fig. 1). The rates of formation of the 7α-hydroxylated products were linear up to 1 μM endogenous sterols (Fig. 1A). Reaction rates were proportional to the incubation time during the first 20 min (Fig. 1 B). The enzyme appeared to be saturated when the sterol concentration reached 200 μM (Fig. 1 C). The apparent Km value of 46 μM observed with acetone-treated microsomes was lower than the range of 80–225 μM reported for untreated microsomes (20–22) and closer to the value of 15 μM observed with butanol/acetone-treated microsomes (23).

**Substrate specificity.** Fig. 2 shows sterol 7α-hydroxylase activity with cholesterol, cholestanol, and sitosterol as substrates, determined in untreated (Fig. 2 A) and acetone-treated (Fig. 2 B) rat liver microsomal preparations. Cholesterol was the preferred substrate and the enzyme activity was 30-fold stronger in acetone-treated microsomes than in untreated microsomes.

![Figure 1](image_url) **Figure 1.** Effect of enzyme concentration, time, and substrate concentration on the 7α-hydroxylation of labeled cholesterol (○), cholestanol (●), and sitosterol (■). (A) Varying amounts of acetone-treated rat liver microsomes (0.4 nmol P-450/mg protein) were incubated at 37°C for 10 min in the presence of 200 μM labeled detergent-solubilized sterol, 0.05 mg/ml DLPC solubilized in 0.12 mM Na-DCA, 2 U NADPH-cytochrome P-450-reductase and 1.2 mM NADPH. The 7α-hydroxylated sterols formed were assayed by TLC and liquid scintillation counting. Each point is an average of two to three experiments, each performed in duplicate with correction for zero-time controls. The coefficient of variation of the assay was < 6%. (B) Assay conditions as described in A were used, except incubation time was varied and two different P-450 concentrations within the linear range were used for each sample. (C) Assay conditions as described in A were used, except the sterol concentration was varied and two different P-450 concentrations within the linear range were used for each sample.
greater for cholesterol than for sitosterol and 1.4 times greater than for cholestanol ($P < 0.0002$) when measured in the untreated microsomes (Fig. 2A). When the incubation was carried out with the acetone-treated microsomes from which endogenous cholesterol had been removed (Fig. 2B), cholesterol and cholestanol exhibited almost the same affinity toward the enzyme (71.8±7.3 and 63.0±4.8 pmol/nmol P-450/min, respectively; $P > 0.05$). The 7α-hydroxylation of sitosterol, in the acetone-treated microsomes, was significantly lower (18.4±0.8 pmol/nmol P-450/min) than that of cholesterol or cholestanol ($P < 0.001$), suggesting a lower affinity for the enzyme. However, it was increased eightfold relative to the value obtained with labeled sitosterol in the presence of endogenous cholesterol in untreated microsomes (18.4±2.4 vs. 2.4±0.6 pmol/nmol P-450/min; $P < 0.001$). Upon removal of endogenous cholesterol, the 7α-hydroxylation of cholestanol was also increased to 63.0±4.8 from 52.4±3.0 pmol/nmol P-450/min, though the difference was not statistically significant.

Figure 2. Substrate specificity of cholesterol 7α-hydroxylase in untreated (A) and (B) acetone-treated rat liver microsomes. The open, hatched, and solid bars represent sterol 7α-hydroxylase activity, assayed as described in Methods, with labeled cholesterol, cholestanol, and sitosterol as substrates, respectively. The data represent the means±SEM of six experiments, each done with duplicate samples. * significantly different from enzyme activity with cholesterol as substrate.

Enzyme inhibition and kinetics. The effect of cholestanol and sitosterol on the 7α-hydroxylation of cholesterol is demonstrated in Figs. 3 and 4. In these experiments, incubations were carried out with increasing amounts of unlabeled cholestanol or sitosterol added to acetone-treated microsomes. Radioactive cholesterol was the substrate and labeled 7α-hydroxycholesterol was the product measured. The results show that cholestanol is a far more potent inhibitor of cholesterol 7α-hydroxylation than sitosterol, probably because of its greater affinity for the enzyme as compared with sitosterol (Fig. 2). In the rat, sitosterol (100 μM) inhibited cholesterol 7α-hydroxylase activity 30%, whereas 100 μM cholestanol inhibited it 60% (Fig. 3). In acetone-treated microsomes from human liver, 100 μM sitosterol and 50 μM cholestanol inhibited cholesterol 7α-hydroxylase activity ~ 40% (Fig. 4). For the determination of the type of inhibition, enzyme activities were measured in acetone-treated rat liver microsomes with two different concentrations of cholestanol or sitosterol and increasing concentrations of labeled cholesterol substrate (Figs. 5 and 6). The Lineweaver-Burk double reciprocal plots showed straight lines that intersected at the same point on the ordinates, indicating competitive inhibition of cholesterol 7α-hydroxylase by cholestanol and sitosterol.

Figure 3. Inhibition of cholesterol 7α-hydroxylase activity by cholestanol and sitosterol in acetone-treated rat liver microsomes. The enzyme activity was assayed with [4,14C]cholesterol as described in Methods in the presence of detergent-solubilized unlabeled cholesterol or sitosterol. Each point represents the mean±SEM of two to three experiments run in duplicate. 100% cholesterol 7α-hydroxylase activity is 76.7±4.0 pmol/nmol P-450/min.
acetone-treated cholesterol 7α-hydroxylase activity was measured in untreated and acetone-treated microsomes, measured with increasing concentrations of labeled cholesterol in the presence of 0, 100, and 200 μM unlabeled sitosterol, S, cholesterol substrate concentration; V, cholesterol 7α-hydroxylase activity. Each point represents the average of two to three experiments run in duplicate.

Discussion

The results of this investigation emphasize the important effect of the microsomal sterol substrate pool on cholesterol 7α-hydroxylase activity. Optimal assay conditions were determined for measuring the 7α-hydroxylation of cholesterol and sitosterol in comparison to cholesterol in acetone-treated microsomes from which virtually all endogenous sterols had been removed. The enzyme was assayed at saturation levels of substrate (200 μM), with enzyme concentration of 0.2–1.0 μM P-450 and incubation time of 10 min that were in the linear range of enzyme activity. The membrane phospholipids lost during acetone treatment of the microsomes were replaced in the incubation mixture with DPLC, the major constituent of liver membrane phospholipids (24). The amount of Na-deoxycholate used to solubilize DPLC was shown previously not to inhibit microsomal cholesterol 7α-hydroxylase (2). Individual phospholipids were not identified since these lipids apparently do not act directly on cholesterol 7α-hydroxylase but facilitate interaction of the enzyme with NADPH-cytochrome P-450 reductase and can be replaced by an excess of the latter (25). Triton X-100 (0.15%) was used as a solubilizing agent for the substrates and was found to yield comparable enzyme activities to 0.3% Triton WR-1339 but higher activities than Cutsicum or Tween-80 used in earlier assays (6). The total sterol concentration that could be used with 0.15% Triton X-100 did not exceed 300 μM (Fig. 6, inset). Over this concentration, high variability of enzyme activity in replicates was observed, possibly due to incomplete solubilization of the sterols. Under the assay conditions used in these experiments, the variation coefficient of cholesterol 7α-hydroxylase activity of the acetone-treated microsomes was < 6%.

Table II. Effect of Endogenous Cholesterol Analougues on Hepatic Cholesterol 7α-Hydroxylase Activity in Humans

<table>
<thead>
<tr>
<th>Subject</th>
<th>Microsomes</th>
<th>Total nmol/mg protein</th>
<th>Cholesterol 7α-hydroxylase activity pmol/nmol P-450/min</th>
</tr>
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<tbody>
<tr>
<td>Control</td>
<td>Untreated</td>
<td>60.1</td>
<td>20.3±0.9</td>
</tr>
<tr>
<td></td>
<td>Acetone-treated</td>
<td>0.48</td>
<td>22.1±1.1</td>
</tr>
<tr>
<td>Sitosterolemia</td>
<td>Untreated</td>
<td>67.8</td>
<td>14.3†</td>
</tr>
<tr>
<td></td>
<td>Acetone-treated</td>
<td>0.54</td>
<td>20.0</td>
</tr>
</tbody>
</table>

*The composition of the plant sterols in microsomes from sitosterolemic patients was as follows: sitosterol, 8.6%; campesterol, 5.9%; 5α-sitosteranol, 0.9%; and 5α-campestanol, 0.7%. N.D. = none detected. † The mass of 7α-hydroxycholesterol formed by the untreated microsomes was determined by the isotope derivative method (17) to avoid underestimation by dilution with the endogenous cholesterol pool. When acetone-treated microsomes were used, the isotope incorporation method was used as described in Methods. The numbers represent means±SEM of six control subjects and the averages of two patients with sitosterolemia and xanthomatosis. ‡ Significantly lower than controls (P < 0.01). The concentrations of endogenous cholesterol and sitosterol present in the in vitro assay of the untreated microsomes from sitosterolemic patients were >5 and >25 μM, respectively, based on the use of 2–3 mg microsomal protein in a total volume of 0.5 ml incubation mixture.
In the study of substrate specificity, cholestanol was selected as the substrate because it contains the identical isooctyl side chain as cholesterol but does not retain its Δ^2-steroid nucleus. The plant sterol, sitosterol, was selected because it differs from cholesterol only by its additional ethyl substituent at C-24 of the side chain. Both compounds are present in abundant amounts in microsomes of patients with sitosterolemia with xanthomatosis (Table II). The affinity of the 7α-hydroxylating enzyme toward these substrates was evaluated in acetone-treated microsomes (Fig. 2 B). It was greatest for cholesterol and poorest for sitosterol. Cholestanol, which has the same isooctyl side chain as cholesterol, exhibited almost the same affinity toward the enzyme. The difference in the nuclear structure did not interfere with the enzymatic hydroxylation of cholestanol. These findings are consistent with the results of Bergstrom et al. (26) and Boyd et al. (27), who studied the 7α-hydroxylation of several cholesterol analogues and suggested that cholesterol 7α-hydroxylase has a specific binding site for the apolar side chain of cholesterol. This binding site may be required for the orientation of the substrate on the enzyme to allow the hydroxylation to occur stereospecifically at the 7α-position. When the 7α-hydroxylation of sitosterol was carried out with the untreated microsomes, it was barely detectable, as has been observed earlier by Aringer and Eneroth (3) and by Boyd et al. (27), but showed an eightfold increase upon removal of endogenous cholesterol (Fig. 2 B).

Similarly, the 7α-hydroxylation of cholestanol was lower in untreated relative to acetone-treated microsomes. This suggests that the large microsomal cholesterol pool competes with sitosterol and cholestanol and partially masks their 7α-hydroxylation. Only when the microsomal cholesterol pool is removed by acetone treatment is the 7α-hydroxylation of sitosterol shown to be significant.

In this study, cholestanol was shown to inhibit cholesterol 7α-hydroxylase activity up to 75%, whereas sitosterol inhibited it maximally 40% (Fig. 3). In earlier studies with the rat, low inhibition of cholesterol 7α-hydroxylase by cholestanol (2) and lack of inhibition by sitosterol (3) have been reported and were probably due to the confounding effect of the endogenous microsomal cholesterol pool in the untreated microsomes that were used. The double reciprocal plots demonstrated that both cholestanol and sitosterol were competitive inhibitors of cholesterol (Figs. 5 and 6). The apparent K_m of cholesterol 7α-hydroxylase measured in the presence of 50 μM cholestanol or 100 μM sitosterol was 87 μM compared with the K_m of 46 μM observed in the absence of any added inhibitor. With 100 μM cholestanol, the apparent K_m increased to 117 μM. Thus, cholestanol is a potent inhibitor of the conversion of cholesterol to its 7α-hydroxylated derivative and has similar affinity as cholesterol toward cholesterol 7α-hydroxylase. Sitosterol, on the other hand, is a less effective inhibitor, has much less affinity for the enzyme and can be 7α-hydroxylated up to 26% relative to the 7α-hydroxylation of cholesterol when competing endogenous cholesterol is removed.

In this study, we have demonstrated for the first time that human liver microsomal cholesterol 7α-hydroxylase activity is preserved after the removal of endogenous sterols by acetone treatment. In human liver, as in the rat, 50 μM cholesterol or sitosterol inhibited cholesterol 7α-hydroxylase activity 35 or 16%, respectively. The importance of this observation concerns cholesterol 7α-hydroxylase activity in the rare inherited lipid storage disease, sitosterolemia with xanthomatosis, in which the overabsorption of plant sterols leads to the accumulation of sitosterol and 5α-saturated stanols in all tissues, including the liver (5). The reduced cholesterol 7α-hydroxylase activity in the patients’ untreated microsomes relative to controls was associated with increased amounts of endogenous cholestanol and plant sterols (Table II). Removal of endogenous sterols by acetone treatment of the microsomes resulted in a 40% increase in cholesterol 7α-hydroxylase activity in the sitosterelemic specimens but did not affect the enzyme activity in the control specimens. When cholestanol or sitosterol was added to human acetone-treated microsomes (Fig. 4) at concentrations (10 or 50 μM) that were similar to those present in assays of untreated microsomes from patients with sitostenolemia with xanthomatosis, cholesterol 7α-hydroxylase activity was reduced 9 and 16%, respectively. The greater inhibitory effect on enzyme activity of endogenous cholesterol analogues observed in microsomes from patients with sitosterolemia and xanthomatosis (40%, Table II) may be due to the presence of additional endogenous cholesterol analogues such as campesterol. Preliminary experiments in our laboratory showed that campesterol, the second most abundant plant sterol in the patients’ microsomal samples, inhibited cholesterol 7α-hydroxylase activity to a similar extent as sitosterol. In addition, the higher inhibitory effect of endogenous cholesterol analogues relative to that observed with exogenous inhibitors may be due to greater availability of endogenous substrate/inhibitors (28).

After ileal bypass surgery, endogenous sterols and 5α-stanols decreased 50% and hepatic cholesterol 7α-hydroxylase activity increased 30%. The rise in activity may be due to both a partial removal of competing cholesterol analogues and an induction of enzyme activity by the interruption of the enterohepatic circulation (2). Recently, Miettinen et al. (29) and Ilias et al. (30) reported that gallstones and bile from patients with cholelithiasis contained substantial amounts of cholestanol, plant sterols, and cholesterol precursors. Further, decreased bile acid pools (31) and low cholesterol 7α-hydroxylase activities (32) have been reported in patients with cholelithiasis. It is tempting to speculate that the lower bile acid synthesis in this condition relates to competitive inhibition of cholesterol 7α-hydroxylase by the associated abnormal microsomal sterol environment. It is important to emphasize that it is difficult to extrapolate an in vitro enzyme activity, expressed as picomole product formed per nanomole cytochrome P-450 per minute, to a daily bile acid synthetic rate expressed in milligrams bile acids formed per kilogram body weight per day. However, cholesterol 7α-hydroxylase activity reflects the rate of bile acid synthesis since increased or decreased enzyme activity, as measured in vitro, correlates with higher or lower bile acid production rates determined in vivo (2, 33). The conversion of sitosterol and cholestanol to primary bile acids has been demonstrated in humans (34) and rats (35, 36), which indicates that both cholesterol analogues are 7α-hydroxylated in vivo.
nous microsomal sterol substrate(s). Thus, analogues of cholesterol may reduce the transformation of cholesterol to bile acids, thereby hindering its removal from the body, and may contribute to premature atherosclerosis in sitosterolemia with xanthomatosis.

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References


