Cytochrome P450 Architecture and Cysteine Nucleophile Placement Impact Raloxifene-Mediated Mechanism-Based Inactivation

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ABSTRACT

The propensity for cytochrome P450 (P450) enzymes to bioactivate xenobiotics is governed by the inherent chemistry of the xenobiotic itself and the active site architecture of the P450 enzyme(s). Accessible nucleophiles in the active site or egress channels of the P450 enzyme have the potential of sequestering reactive metabolites through covalent modification, thereby limiting their exposure to other proteins. Raloxifene, a drug known to undergo CYP3A-mediated reactive metabolite formation and time-dependent inhibition in vitro, was used to explore the potential for bioactivation and enzyme inactivation of additional P450 enzymes (CYP1A2, CYP2C8, CYP2C9, CYP2C19, CYP2D6, CYP2E1, and CYP3A5). Every P450 tested except CYP2E1 was capable of raloxifene bioactivation, based on glutathione adduct formation. However, raloxifene-mediated time-dependent inhibition only occurred in CYP2C8 and CYP3A4. Comparable inactivation kinetics were achieved with \( K_i \) and \( k_{\text{inact}} \) values of 0.26 \( \mu M \) and 0.10 \( \text{min}^{-1} \) and 0.81 \( \mu M \) and 0.20 \( \text{min}^{-1} \) for CYP2C8 and CYP3A4, respectively. Proteolytic digests of CYP2C8 and CYP3A4 Supersomes revealed adducts to Cys225 and Cys239 for CYP2C8 and CYP3A4, respectively. For each P450 enzyme, proposed substrate/metabolite access channels were mapped and active site cysteines were identified, which revealed that only CYP2C8 and CYP3A4 possess accessible cysteine residues near the active site cavities, a result consistent with the observed kinetics. The combined data suggest that the extent of bioactivation across P450 enzymes does not correlate with P450 inactivation. In addition, multiple factors contribute to the ability of reactive metabolites to form apo-adducts with P450 enzymes.

Introduction

The cytochrome P450 (P450) enzyme family is largely responsible for the metabolism of xenobiotics (Guengerich, 2001; Wienkers and Heath, 2005). Although P450-mediated metabolism usually leads to detoxification, bioactivation and reactive metabolites formation can occur, increasing the risk of drug-induced liver injuries (Mitchell et al., 1973; Dahlin et al., 1984; Leung et al., 2012). A key differentiating factor for a xenobiotic to form a benign, oxidative metabolite or a reactive metabolite is the innate chemistry of the xenobiotic (Nelson, 1982). The term structural alert classifies a set of chemical functional groups prone to P450-mediated reactive metabolite formation (Kalgutkar and Didiuk, 2009; Stepan et al., 2011). Exposure of these functional groups to P450 enzymes is anticipated to increase the probability of reactive metabolite formation; therefore, strategies to identify and limit reactive metabolite formation are of importance.

Substrate interactions with the active site architecture of P450 enzymes also govern the regioselectivity of metabolite formation, potentially impacting bioactivation and exposure to reactive metabolites. For example, furafylline possesses a structural alert in the furan moiety. However, bioactivation has been shown to occur exclusively at the 8-methyl position of the xanthine moiety (Kunze and Trager, 1993). Docking studies with furafylline and CYP1A2 position the 8-methyl position toward the heme iron based upon a prominent interaction between the furan ring and Phe125 of the enzyme active site, highlighting the potential for the enzyme to direct metabolite regioselectivity (Lewis and Lake, 1996). P450 active site architecture not only orients the substrate, potentially limiting the formation of reactive metabolites, but also can sequester bioactivated metabolites through covalent trapping by accessible nucleophiles. Reactive metabolites

ABBREVIATIONS: P450, cytochrome P450; GSH, glutathione; MS, mass spectrometry; TDI, time-dependent inhibition.
trapped in the active site by nucleophilic amino acids directly limit tissue exposure, thereby decreasing the release of reactive metabolites and, indirectly, the potential toxicity of bioactivation via feedback inhibition (Pearson et al., 2007).

Over the last decade, the X-ray crystal structures of P450s offered structural insights to the active site topography and defined key residue interactions with cocrystallized ligands (Johnson and Stout, 2005; Dong and Wu, 2012). This structural information facilitates the location of nucleophilic residues within the active sites of P450 enzymes. A substrate prone to reactive metabolite formation could be used to probe the location of relevant active site nucleophiles. P450-mediated reactive metabolite formation and subsequent enzyme inactivation typically have one of three fates: the reactive metabolite adds the heme, forms a metabolic intermediate complex with the heme, or covalently adds the apo-protein (Kalugtka et al., 2007). Alkynes and alkyl amine-containing compounds are classic examples where the reactive metabolites react with the heme and heme iron, respectively (Chan et al., 1993; Hanson et al., 2010). In this instance, the reactive metabolite formed has relatively the same potential for reactivity across P450 enzymes given the conserved nature of the heme. However, some reactive metabolites are quenched preferentially by nucleophilic amino acid residues in the active site, which leads to P450 inactivation (Baer et al., 2007; Henne et al., 2012; Lin et al., 2012). The structure of reactive metabolites is commonly inferred by the identification of glutathione (GSH) adducts (Baillie et al., 1989; Dieckhaus et al., 2005). The cysteine thiol in GSH represents a delocalized (soft) electrophile with vast reactivity and is physiologically relevant given the high cellular concentrations. As such, GSH-adducted xenobiotics are commonly inferred to be the chemical entities responsible for P450 inactivation through apo-adduct formation.

Raloxifene undergoes bioactivation and the formation of GSH adducts in vitro by both CYP3A4 and CYP3A5. However, raloxifene is a time-dependent inhibitor of only CYP3A4 (Pearson et al., 2007). The difference in inhibition was attributed to a single cysteine residue (Cys239) that is present in CYP3A4 but not CYP3A5. Cys239 sits at the interface between the active site and bulk solvent. Blocking this residue with iodoacetamide and/or mutating the cysteine to alanine led to an increase in the formation of reactive metabolites from raloxifene as inferred from the increase in GSH adducts. The increase in GSH-conjugated raloxifene. The increase in GSH-adduct raloxifene was even more dramatic in CYP3A5. Although the raloxifene adduct findings are not physiologically relevant due to the limited role of P450 metabolism in the in vivo clearance (Kemp et al., 2002), other xenobiotics including (R)-N-(1-(3-(4-ethoxyphenyl)-4-oxo-3,4-dihydropyridine-2,3-dipyrinylmethyl)-11-(pyridin-3-ylmethyl)-2-(4-tribromoethoxy)phenyl)acetamide (AMG 478) (Henne et al., 2012) and carbamazepine (Kang et al., 2008) have been shown to or proposed to form adducts with Cys239.

The work presented herein investigates the potential role of active site cysteine residues across a panel of P450 enzymes for their ability to trap reactive metabolites. Raloxifene was used as a probe to explore the potential for bioactivation and inactivation across eight P450s (CYP1A2, CYP2C8, CYP2C9, CYP2C19, CYP2D6, CYP2E1, CYP3A4, and CYP3A5). In addition, the amino acid architectures of the P450 active sites were examined and discussed in terms of the observed kinetic data.

**Materials and Methods**

**Materials.** Superoxides coexpressed with cytochrome b_{5} and P450 reductase were purchased from BD Biosciences (San Jose, CA). Trypsin and protease K were purchased from Roche Diagnostics (Indianapolis, IN). Raloxifene, tolbutamide, diclofenac, dextromethorphan, dextrophan, midazolam, 1-hydroxymidazolam, 6β-hydroxytestosterone, paclitaxel, reduced GSH, and chlorzoxazone were purchased from Sigma-Aldrich. (St. Louis, MO). (S)-Mephénytoin was purchased from Enzo Life Sciences, Inc. (Farmingdale, NY). 6-Hydroxy-docetaxel and reduced β-NADPH were purchased from Calbiochem (San Diego, CA). 4’-Hydroxy-(S)-mephénytoin, 4’-hydroxyclofenac, and 6-hydroxychlorzoxazone were purchased from BD Biosciences. Phenacetin and acetaminophen were purchased from MP Biomedicals (Solon, OH). Testosterone was purchased from Steraloids (Newport, RI). All of the other chemicals and liquid chromatography solvents were acquired from commercial sources and were of the highest grade available.

**Measurement of Raloxifene IC_{50} Values, IC_{50} Shift Assays and Time-Dependent Inhibition.** IC_{50} values of raloxifene were determined against both testosterone and midazolam. Before conducting IC_{50} measurements, K_{m} and k_{cat} values for both testosterone and midazolam were determined for each enzyme studied (R. S. Foti, unpublished observations) and the results were consistent with published data (Dickmann et al., 2012). Varying concentrations of raloxifene (0–100 μM) were preincubated with enzyme (1 pmol per 100 μl final incubation volume) and substrate (at the predetermined K_{m} value) for 5 min before the initiation of the reaction with NADPH. The reactions were quenched after 5 min for midazolam or 20 min for testosterone with an equal volume (100 μl) of 0.5 μM tolbutamide in acetonitrile. Incubations were run in triplicate, and IC_{50} values were determined in Prism 5.0 (GraphPad Software, Inc., San Diego, CA). The IC_{50} values were used to demonstrate the potential for raloxifene to reversibly bind to the eight P450s under investigation and provide an appropriate initial estimate for raloxifene concentrations in the subsequent time-dependent inhibition (TDI) experiments. IC_{50} shift experiments were performed under similar conditions and as reported previously (Henne et al., 2012). For raloxifene IC_{50} and TDI studies, a 4000 QTRAP from Applied Biosystems (Foster City, CA) was used and interfaced to a high-performance liquid chromatograph (Shimadzu, Kyoto, Japan). Mass spectrometry (MS) parameters for each metabolite measured are listed in Supplemental Table 1. Common to all of the compounds were the dwell time, 500 μs; curtain gas, 10 psi; IonSpray voltage, 4500; and source temperature, 400°C. IonSpray gases 1 and 2 were set at 40.

**Raloxifene Metabolism and GSH Adduct Formation.** Individul P450 enzymes (10 pmol) were preincubated for 5 min at 37°C before the addition of 10 mM NADPH to a final concentration of 1 mM (1 ml total volume) in the presence of 10 μM raloxifene and 1 mM GSH. Aliquots were removed from the incubation at 0, 3, 6, 9, 12, and 15 min and placed into 100 μl of cold acetonitrile spiked with 0.5 μM tolbutamide for an internal standard. All of the incubations were run in triplicate. Remaining concentrations of raloxifene were measured using the 4000 QTRAP using a multiple reaction monitoring scan with the quadrant 1 (Q1) m/z 474.3 and the Q3 m/z 111.9. The declustering potential was 66, and the collision energy was 47. Tolbutamide was monitored with Q1 m/z 271.2 and Q3 m/z 91.1. Common to both compounds were the dwell time, 500 μs; curtain gas, 10; IonSpray voltage, 4500; and source temperature, 400°C. IonSpray gases 1 and 2 were set at 40. TDI was characterized for CYP3A4 and CYP2C8 based on the IC_{50} shift results. The assay was performed with a two-step procedure in a 96-well format. Each time point was sampled in triplicate. Inactivation in preincubation mixtures was conducted in potassium phosphate buffer (100 mM, pH 7.4) containing CYP3A4 or CYP2C8 Supersomes and raloxifene.
(0–10 μM). After equilibration for 3 min at 37°C, reactions were initiated by the addition of NADPH (1 mM) in a final incubation volume of 0.2 ml. Reactions proceeded for 0, 2, 4, 6, or 10 min, at which time aliquots (10 μl) were transferred to an activity assay for the assessment of remaining P450 activity. Incubations (prewarmed to 37°C) contained NADPH (1 mM) and either midazolam for CYP3A4 (10 μM) or paclitaxel for CYP2C8 (200 μM) in potassium phosphate buffer (100 mM, pH 7.4). The final volume was 0.2 ml. Reactions were allowed to proceed for 5 min and were terminated by the addition of an equal volume of acetonitrile containing tolbutamide (0.5 μM) as an internal standard. Quenched mixtures were vortex-mixed and centrifuged, and the resulting supernatants were analyzed via liquid chromatography-tandem mass spectrometry as described above. Kinetic parameters were determined using Prism. The natural log of the percentage of remaining CYP3A4 activity versus preincubation time was plotted for each inhibitor concentration, and initial inactivation rate plots were fitted to the following relationship:

\[ k_{\text{obs}} = (k_{\text{inact}} \times I) + k_{\text{inc}} \]

where \( k_{\text{obs}} \) is the observed inhibitor concentration, \( k_{\text{inact}} \) is the initial inactivation rate constant, and \( I \) is the inhibitor constant. The rate of enzyme inactivation is half the maximal rate, \( k_{\text{inact}} \), which time aliquots (10 μl) were transferred to an activity assay for the assessment of remaining P450 activity. Incubations were placed subsequently on ice and concentrated to a volume of 0.2 ml using a Savant SpeedVac (Thermo Fisher Scientific, Waltham, MA). Aliquots (400 μl) of ammonium bicarbonate buffer (50 mM, pH 8.1) and methanol (50 μl) were added to each incubation sample. Mixtures then were incubated with trypsin (Roche Diagnostics) in a 1:25 (w/w) ratio overnight at room temperature. After proteolysis, the pH was adjusted to 5 by the addition of 0.1% trifluoroacetic acid followed by the addition of acetonitrile (5% final volume). Digested sample mixtures were analyzed immediately by liquid chromatography-tandem mass spectrometry. An Accela 1250 high-performance liquid chromatography system coupled to an HTS PAL autosampler (LEAP Technologies, Carrboro, NC) interfaced with an LTQ-Orbitrap Velos mass spectrometer (Thermo Fisher Scientific) was used for peptide analysis. Peptide samples were injected onto a Jupiter C18 column (3 μm, 2.1 × 150 mm; Phenomenex, Torrance, CA). A flow rate of 0.2 ml/min was used, with a portion of the column elute (20%) diverted to the mass spectrometer. Mobile phase consisted of 0.05% formic acid in H2O (A) and 0.05% formic acid in acetonitrile (B). Initial conditions were 98% A, with a linear gradient: 2% B for 2 min, 2 to 95% B over 35 min, and 95% B for 5 min. Ions were detected in positive mode; precursor ions were selected for collision-induced dissociation were produced using 35% collision energy and a 1.0-Da isolation window. The resulting data were searched via SEQUEST embedded in Proteome Discoverer 1.2 (Thermo Fisher Scientific). A protein preparation script embedded in Maestro (Schrodinger, Portland, OR) was used to prepare the P450 structures before use in CAVER. In brief, each protein Data Bank was imported and automatically corrected for missing hydrogen atoms. All of the cocrystallized water molecules were removed, and the iron formal charge was defined at +3. Protonation states for the histidine residues were defined and automatically corrected for missing hydrogen atoms. All of the cocrystallized water molecules were removed, and the iron formal charge was defined at +3. Protonation states for the histidine residues and heavy atoms in arginine, glutamine, and histidine side chains also are assigned automatically. The protein hydrogen bond network also is optimized before a final restrained minimization, which allows the hydrogen atoms to freely minimize within the protein. Prepared P450 structures then were explored using CAVER as described previously (Petrek et al., 2006). In brief, the active site was defined by an active site water axial to the heme ligand. The default number of channels was explored with 4 Å of nearest neighbor. Protein cavities and tunnels were identified by analyses of a series of molecular dynamics trajectory files with CAVER and visualized with PyMOL (Schrodinger).

**Results**

**Raloxifene Metabolism and GSH Conjugate Formation by P450 Enzymes.** Fig. 1 depicts the metabolic scheme for the P450-mediated metabolism of raloxifene in vitro. All eight P450 isoforms tested exhibited different amounts of raloxifene oxidation. CYP2C19 showed the largest extent of raloxifene hydroxylation (Fig. 2). In contrast, no evidence of CYP2E1 metabolite formation from raloxifene was observed. Metabolism experiments conducted in the presence of GSH determined the relative percentage of reactive metabolite formation. CYP3A5 yielded the most GSH-related adducts when compared with the other P450 enzymes (Fig. 2; Supplemental Fig. 1). Differences in the regioselectivity of GSH adduct formation also were observed. For example, only CYP1A2, CYP3A4, and CYP3A5 demonstrated the potential to form adducts designated GSH1 and GSH2. GSH1 and GSH2 adducts are derived from the β-o-quinone (Pearson et al., 2007; Fig. 1). In contrast, all of the P450 enzymes examined formed GSH adducts from the precursor, diquinone methide, except for CYP2E1, where no GSH adducts were observed, which is consistent with a lack of raloxifene metabolism.

**Structural Characterization and In Silico Modeling of P450 Enzymes.** The Protein Data Bank files used for modeling were 2HI4 for CYP1A2, 1PQZ for CYP2C8, 1R9O for CYP2C9, 2F9Q for CYP2D6, and 1TQN for CYP3A4. For CYP2C19, a homology model was generated based upon the crystal structure of CYP2C9 (Protein Data Bank code 1R09). For CYP3A5, a homology model was generated based upon the crystal structure of CYP3A4 (Protein Data Bank code 1TQN).
greater than 2-fold. On the basis of the IC50 shift results, a CYP3A4 showed a shift in potency from control samples identify the potential for TDI (Table 1). CYP2C8 and

4.21 enzymes, except CYP2E1. The inhibition ranged from 0.39 to

onstrated the potential to inhibit each of the eight P450 pathway B, Fig. 1B, and oxidized 3’ hydroxy raloxifene, pathway A, Fig. 1C. Percentages of metabolites formed are represented in triplicate. +, Increased by 5x.

Raloxifene-Mediated P450 Inhibition. Raloxifene demonstrated the potential to inhibit each of the eight P450 enzymes, except CYP2E1. The inhibition ranged from 0.39 to 4.21 μM. For CYP1A2, the inhibition could not be determined readily based upon the inability for fits to converge on the data (Table 1). On the basis of metabolism and inhibition across P450 isoforms, the potential for raloxifene-mediated TDI was assessed. An IC50 shift experiment was used to identify the potential for TDI (Table 1). CYP2C8 and CYP3A4 showed a shift in potency from control samples greater than 2-fold. On the basis of the IC50 shift results, a full kinetic profile for TDI of CYP2C8 and CYP3A4 was performed. The K1 and k inactive values for CYP2C8 were 0.26 μM and 0.10 min⁻¹, respectively. For CYP3A4, the K1 and k inactive values were 0.81 μM and 0.20 min⁻¹, respectively (Table 1), and were similar to previous results (Chen et al., 2002).

Identification of the Mechanism for Raloxifene-Mediated TDI of CYP2C8. Raloxifene-mediated TDI in CYP2C8 was predicted to occur via apo-adduct formation based on quantitative recovery of heme and CO (D. A. Rock, unpublished observations). These results precluded heme adduct formation and a metabolic intermediate complex as mechanisms for CYP2C8 inactivation. Proteolysis of CYP2C8 after the incubation with raloxifene and NADPH (n = 4) yielded peptides covering 85 to 94% of the protein sequence, including a putative [M + 3H]³⁺ peptide adduct at m/z 724.0281 Da, corresponding to [M + H]⁺ = 2170.0699 Da. MS³ experiments with m/z 724.0281 serving as the precursor ion revealed doubly charged b and y ions (Fig. 3) highlighted in blue and red, respectively. The y ion series with 7 of the 14 ions identified provided sufficient sequence coverage to identify the modified peptide as LIDcFPGTHNKVNKLK²²², specifically, y₁₂ is crucial in the identification of Cys225 as the adducted amino acid by bioactivation of raloxifene to the diquinone methide species, with a change in m/z of 471.1504. In addition, the theoretical molecular weight of the adducted peptide was consistent with the experimentally derived molecular weight ([M + H]⁺ = 2170.0699), differing by 9.1 ppm. The b ion series exhibited similar results with six ions identified, including b₁, which is consistent with Cys225 reacting with the diquinone methide species of raloxifene. The reaction of an active cysteine with the diquinone methide species of raloxifene is similar to the previous report of the raloxifene adduct Cys239 of CYP3A4 (Bae et al., 2007). Subsequent MS³ analysis conducted using m/z 724.0281 to 914.92 (γ₁₂) as precursors generated product ions consistent with the adducted peptide (Supplemental Fig. 2). Search algorithms were employed using the raloxifene o-quinone mass as a potential contributing inactivating species, but no adducts were observed.

Structural Characterization of the Active Site Cysteine. The program CAVER has been used to find the most accessible path from the active site to the surface of a protein (Petrek et al., 2006). Mapping of each P450 enzyme with CAVER revealed a conserved access channel distal to the heme moving from the I helix toward the end of the G and G’ helices (Fig. 4). Despite this conserved channel, there were unique channels found for several of the P450s. For example, the primary access channel defined for CYP1A2 (Protein Data Bank code 2HI4) exits parallel to the I helix, between the G helix and the I helix such that it runs parallel to the I helix (green channel, Fig. 4A). A second egress channel begins to follow this trajectory but breaks toward the B’ helix (orange channel, Fig. 4A). In contrast, the cysteine residues with the closest proximity to the active site in CYP1A2 are located parallel to the heme adjacent to one another on the opposite side of the protein in the β2 sheet region, Cys405 and Cys406. For CYP2C8, there are two cysteine (Cys216 and Cys225) residues distal to the heme. Upon overlay of CYP2C8 and CYP3A4, the placement of Cys216 and Cys225 are in close proximity to that of Cys239 for the respective enzymes (Fig. 5). From the CAVER results, a unique access channel in CYP2C8 was identified between the B’ helix and the G’ helix (green channel, Fig. 4B). In contrast, the divergent channel in CYP3A4 was found to exit between the F’ helix and the G’ helix (blue channel, Fig. 4F). On the

TABLE 1
Raloxifene-mediated P450 inhibition kinetic constant

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>IC₅₀</th>
<th>IC₅₀ Shift</th>
<th>K₁</th>
<th>k inactive</th>
<th>Partition Ratio</th>
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<td></td>
<td>μM</td>
<td>μM</td>
<td>μM</td>
<td>min⁻¹</td>
<td></td>
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<td></td>
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</tr>
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NA, not available; NI, no inhibition.
basis of the CAVER results and known substrate recognition sites, each P450 enzyme was positioned with the I helix parallel to the z-axis and divided into four quadrants (Fig. 4). All of the cysteine residues were identified in each of the quadrants (Table 2).

Discussion

Mammalian P450 enzymes have a conserved globular tertiary structure (Johnson and Stout, 2005). At the core of P450 enzymes is a porphyrin heme, locked in place by electrostatic interactions with a number of conserved arginine residues and a ring of hydrophobic amino acids that sandwich the heme against the rigid, conserved I helix (Wester et al., 2004; Yano et al., 2004; Sansen et al., 2007; Halpert, 2011; Shah et al., 2011). Positioning the I helix of each P450 parallel to the z-axis, each of the isoforms was divided into four quadrants to explore the location of the cysteine residues (Fig. 4A). In this view, the potential substrate and metabolite recognition sites are almost exclusively positioned in Q1 across the different mammalian P450s. Therefore, the ability of a P450 enzyme to quench reactive metabolites is directed by the presence and accessibility of nucleophiles that can contact reactive metabolites in Q1. Reactive metabolites formed in P450 enzymes without matched nucleophiles could lead to uncontrolled reactive metabolite formation, resulting in an increased exposure to reactive metabolites and potential for toxicity. Given the defined oxidative metabolic pathway and the previous characterization of GSH adducts, raloxifene-mediated P450 bioactivation and inactivation were explored across eight P450 enzymes.

Raloxifene is an effective therapeutic in the treatment of breast cancer. In vivo, the exposure and elimination of raloxifene are governed by gut glucuronidation (Kemp et al., 2002). However, in vitro raloxifene is subject to CYP3A me-

Figure 3. Mass spectra of the CYP2C8 raloxifene-adducted peptide, 222-L1DcFPGTHNKVLKN236. Spectrum acquired at 24.1 min with parent [M + 3H]3⁺ = m/z 724.0281 corresponding to a mass error of 9.1 ppm. Blue ions (y ions) and red ions (b ions) masses are listed in the table to the right of the ions labeled on the tandem mass spectrometry spectra.

Figure 4. Active site access channels identified with CAVER in CYP1A2 (A), CYP2C8 (B), CYP2C9 (C), CYP2C19 (D), CYP2D6 (E), and CYP3A4 (F). The top three trajectories depicted as tubes showing the route followed by the center of mass of the substrate raloxifene. Each structure is divided into four quadrants using the I helix as the intersection point. Helices are labeled in corresponding red letters.
tabolism, resulting in the formation of multiple reactive metabolites, including a diquinone methide and o-quinone (Chen et al., 2002; Yu et al., 2004). The formation of the diquinone methide introduces an electrophilic structure capable of reacting at multiple sites over 12 Å, due to the extensive conjugation of the molecular structure (Fig. 1). The diquinone methide reacts with soft nucleophiles such as GSH and N-acetyl cysteine (Chen et al., 2002) but does not appear to react readily with more charge-localized nucleophilic residues such as lysine (D. A. Rock, unpublished observations). Additional reactivity of the diquinone methide has been shown toward acetic and propionic acid, suggesting the potential for the diquinone methide to react with aspartic or glutamic acid under nonphysiological conditions (Moore et al., 2010). However, the removal of Cys239 abolished TDI in CYP3A4, suggesting that ester formation with diquinone methide is too unstable to lead to TDI. In addition, no isolable aspartic or glutamic acid raloxifene adducts were observed herein; albeit the transient nature of this complex could preclude the isolation of the unstable peptide adduct. On the basis of the ability of raloxifene diquinone methide to readily react with cysteine, it serves as an excellent probe to explore the potential of P450 enzyme architecture for accessible cysteine residues.

Raloxifene incubations with all eight P450s led to the formation of the diquinone methide except for CYP2E1 (Fig. 2). In contrast, incubations set up to evaluate inactivation kinetics showed that only CYP2C8 and CYP3A4 are susceptible to TDI (Table 1). An overlay of CYP2C8 and CYP3A4 structures revealed the presence of cysteine residues in Q1 for CYP2C8 similar to those for CYP3A4 (Fig. 5). For CYP2C8, two cysteine residues were identified in Q1. Cys225 resides in helix F’ and serves as the upper boundary to the CYP2C8 active site. Helix F’ also is responsible for the interactions with substrates in the active site (Schoch et al., 2008). The second cysteine, Cys216, resides at the start of helix G, also having the potential to interact with substrates and metabolites. In comparison, in CYP3A4, a single cysteine (Cys239) resides in the flexible loop preceding helix G.

Inactivation of CYP3A4 by raloxifene occurs through apooduct formation with Cys239 as demonstrated previously in reconstituted CYP3A4 enzyme (Baer et al., 2007). Confirmation of Cys239 adduction by raloxifene from CYP3A4 Supersomes is unique compared with previous efforts to identify P450 adducts that used purified, reconstituted enzyme (Lightning et al., 2000; Bateman et al., 2004; Wen et al., 2005; Yukinaga et al., 2007). A challenge with the reconstitution of recombinant P450 is the complete incorporation of the enzyme into the lipid membrane, whereby any P450 left unincorporated into the lipid could trap reactive metabolites akin to an exogenously added nucleophile. Corroboration of the previously generated recombinant data with CYP3A4 tij;2Supersomes confirms that Cys239 is the prominent residue involved in CYP3A4 inactivation. Therefore, the use of membrane-prepared recombinant P450 enzymes should reduce the potential for falsely identified adducts while simultaneously increasing the speed and accuracy of adduct characterization.

The mechanism(s) behind raloxifene-mediated CYP2C8 inactivation did not occur through heme adduction or metabolic intermediate complex formation (D. A. Rock, unpublished observations). Furthermore, given the structural alert and known propensity of raloxifene to form the diquinone methide, an apo-adduct(s) with CYP2C8 was hypothesized. A single raloxifene adduct was identified at Cys225 with a mass consistent with the addition of the diquinone methide.
metabolite. Not surprisingly, Cys225 from CYP2C8 and Cys239 from CYP3A4 approximate the same spatial arrangement upon the overlay of the two structures (Fig. 5). Cys216 in CYP2C8 is in close proximity to Cys225. A noteworthy finding is that both CYP2C9 and CYP2C19 also possess Cys216 similar to that found in CYP2C8. Upon the overlay of all of the CYP2C structures, Cys216 resides in a similar space for each enzyme. The lack of adduct formation at Cys216 with raloxifene in CYP2C8 is also consistent with the lack of TDI observed in both CYP2C9 and CYP2C19.

The surrounding protein environment also greatly impacts the nucleophilicity of the cysteine thiols (Ferrer-Sueta et al., 2011). In addition, many protein surface thiols often are targets for oxidation (Requejo et al., 2010). On the basis of the uncoupling of the P450 catalytic cycle and ability to generate hydrogen peroxide, there is a potential for the cysteines near the active site to undergo oxidation to sulfenic acid. Sulfenic acid is highly reactive and capable of undergoing nucleophilic attack or additional oxidations ultimately to a stable sulfonic acid derivative (Romero et al., 1992; Liu et al., 1996; Claiborne et al., 2001). Analysis of peptide digests for oxidized cysteines revealed that all of the CYP2C8 digests possessed a peptide containing oxidized Cys216 (SO3H). Therefore, the oxidation of Cys216 could easily compromise reactivity toward various electrophiles.

In addition, the absence of a Cys216 raloxifene adduct could be a result of specific egress channel of the diquinone methide that traverses proximal to Cys225. Using CAVER, access channels in CYP2C8 were modeled in an attempt to gain further insight to the regioselectivity demonstrated by raloxifene diquinone methide. Three primary channels were observed for CYP2C8. The largest channel in CYP2C8 overlaps with that determined in CYP3A4. In addition, the second most prominent channel traverses a similar path between CYP2C8 and CYP3A4. The second channel passes in close proximity to Cys225 and Cys239 in CYP2C8 and CYP3A4, respectively. No access channels were shown to pass directly by Cys216, which is consistent with the unmodified peptide observed in the digest. The combined data support the notion that CYP2C8 TDI is mediated through the raloxifene diquinone methide predominantly, if not exclusively, by Cys225.

The lack of raloxifene-mediated inactivation of CYP1A2, CYP2D6, and CYP3A5 is consistent with the lack of cysteine residues present in Q1. However, numerous examples of TDI have been documented with the aforementioned enzymes. Many of the examples cause TDI through metabolic intermediate complex formation and are not applicable to this discussion (VandenBrink and Isoherranen, 2010). However, examples where TDI is likely to occur by apo-adduct formation exist. For example, CYP2D6 inactivation was observed with 1-[(2-ethyl-4-methyl-1H-imidazo[5-yl]methyl)-4-(trifluoromethyl)-2-pyrindinyl]piperazine as an apo-adduct (Hutzler et al., 2004). The exact site and residue of adduct formation was never pursued. CYP1A2-mediated metabolism of zileuton also leads to TDI, for which the mechanism is hypothesized to occur through apo-adduct formation (Lu et al., 2003). There are two interesting concepts conserved across these TDI examples: 1) GSH adducts for the drugs have been identified implicating the potential for apo-protein formation and 2) there are a lack of cysteine residues in Q1. A potential explanation for the disconnect between the observed inactivation and the absence of Q1 cysteines would be the potential for novel egress channels beyond Q1. For CYP1A2, there are two cysteine residues that reside in close proximity to the active site, Cys405 and Cys406. However, the CYP1A2 egress channels produced in CAVER do not converge upon residues Cys405 or Cys406 or any other cysteine residues within the enzyme. Conversely, the reactive metabolites may have sufficient electrophilic properties to react with other less commonly described nucleophiles; additional research is warranted to link these reactive metabolites to P450 inactivation. Moreover, on the basis of these structures and predicted access channels, other drugs prone to bioactivation and reactivity toward soft electrophiles also may be capable of escaping the active site without enzyme inactivation. Reactive metabolites with similar reactivities suggest that P450 clearance pathways could influence exposure to reactive metabolites and exacerbate drug-induced liver injury. Exploration of this with more relevant drugs, for example, drugs that are cleared in vivo by different P450s that concurrently show the potential for hepatotoxicity, will ultimately enable further testing of this hypothesis.

In conclusion, all of the P450 enzymes examined (except CYP2E1) formed the diquinone methide, thus providing a probe of each active site for solvent-accessible cysteines capable of enzyme inactivation. Similar to CYP3A4, an active site cysteine residue in CYP2C8 is alkylated upon raloxifene bioactivation to the diquinone methide to inactivate the enzyme. The inability for inactivation by the diquinone methide with CYP1A2, CYP2D6, and CYP3A5 is consistent with the crystal structure data wherein no cysteines are present in Q1. Additional support from the CAVER-derived access channels clearly demonstrates that the primary metabolite exit channels are void of cysteine residues capable of trapping the raloxifene diquinone methide. The inability to inactive CYP2C9 and CYP2C19 could be due to oxidation of Cys216 similar to reports here with CYP2C8. Finally, the combined data presented suggest the extent of reactive metabolite formation by a P450 enzyme does not correlate with P450 inactivation. Multiple factors, including active site architecture, exit channels, and chemical structure of the reactive metabolite all may contribute to the ability to form apo-adducts with a P450 enzyme.

Authorship Contributions

Participated in research design: VandenBrink, Davis, Pearson, Foti, Wienkers, and Rock.

Conducted experiments: VandenBrink, Davis, Pearson, Foti, and Rock.

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Wrote or contributed to the writing of the manuscript: VandenBrink and Rock.

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