In Vitro Metabolism of Oprozomib, an Oral Proteasome Inhibitor: Role of Epoxide Hydrolases and Cytochrome P450s

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ABSTRACT

Oprozomib is an oral proteasome inhibitor currently under investigation in patients with hematologic malignancies or solid tumors. Oprozomib elicits potent pharmacological actions by forming a covalent bond with the active site N-terminal threonine of the 20S proteasome. Oprozomib has a short half-life across preclinical species and in patients due to systemic clearance via metabolism. Potential for drug–drug interactions (DDIs) could alter the exposure of this potent therapeutic; therefore, a thorough investigation of pathways responsible for metabolism is required. In the present study, the major drug-metabolizing enzyme responsible for oprozomib metabolism was identified in vitro. A diol of oprozomib was found to be the predominant metabolite in human hepatocytes, which formed via direct epoxide hydrolysis. Using recombinant epoxide hydrolases (EHs) and selective EH inhibitors in liver microsomes, microsomal EH (mEH) but not soluble EH (sEH) was found to be responsible for oprozomib diol formation. Coincubation with 2-nonylsulfanyl-propionamide, a selective mEH inhibitor, resulted in a significant decrease in oprozomib disappearance (>80%) with concurrent complete blockage of diol formation in human hepatocytes. On the contrary, a selective sEH inhibitor did not affect oprozomib metabolism. Pre-treatment of hepatocytes with the pan-cytochrome P450 (P450) inhibitor 1-aminobenzotriazole resulted in a modest reduction (∼20%) of oprozomib metabolism. These findings indicated that mEH plays a predominant role in oprozomib metabolism. Further studies may be warranted to determine whether drugs that are mEH inhibitors cause clinically significant DDIs with oprozomib. On the other hand, pharmacokinetics of oprozomib is unlikely to be affected by coadministered P450 and sEH inhibitors and/or inducers.

Introduction

The proteasome is a multicatalytic enzyme complex and key to the ubiquitin-dependent pathway for intracellular protein degradation (Kisselev and Goldberg, 2001; Ciechanover, 2005). The highly regulated ubiquitin proteasome system affects a wide variety of cellular processes and is critical for cellular homeostasis. Inhibition of proteasome leads to accumulation of misfolded proteins and induction of cell cycle arrest and apoptosis. Transformed cells, particularly multiple myeloma (MM) cells, possess elevated levels of proteasome activity. Consequently, these actively proliferating malignant cells are more sensitive to proteasome inhibitors than normal cells (Kumatori et al., 1990; Loda et al., 1997; Li and Dou, 2000; Alnond and Cohen, 2002). Proteasome inhibition has emerged as an effective therapeutic strategy for the treatment of MM and some lymphomas. Bortezomib (Velcade), the first-generation proteasome inhibitor, was approved in 2003 (Bross et al., 2004; Chauhan et al., 2005; O’Connor et al., 2005). Carfilzomib (Kyprolis), a peptide epoxyketone (Fig. 1), represents a new generation of proteasome inhibitors. It selectively binds to the N-terminal threonine of the proteasome via two covalent bonds to form a morpholino ring (Harshbarger et al., 2015), and has demonstrated favorable clinical safety and efficacy profiles compared with bortezomib (Kuhn et al., 2007). Carfilzomib has been approved for use as a single agent and in combination with lenalidomide and dexamethasone in patients with relapsed and/or refractory MM (Thompson, 2013). Notably, carfilzomib in combination with dexamethasone has doubled progression-free survival in patients with relapsed and/or refractory MM compared with bortezomib and dexamethasone in the phase 3 ENDEAVOR clinical trial (Dimopoulos et al., 2016). The introduction of proteasome inhibitors and immunomodulatory drugs has revolutionized MM treatment with the current overall survival of MM patients increased by 2- to 3-fold (Anderson, 2013).

The success of bortezomib and carfilzomib has inspired efforts to develop oral proteasome inhibitors with improved flexibility in dose...
administration, specifically oral administration, for improved patient compliance and convenience. Ixazomib (Ninlaro) and oprozomib, representing peptide boronate and peptide epoxyketone, respectively, are two orally bioavailable proteasome inhibitors under clinical investigation (Rajan and Kumar, 2016). Ixazomib in combination with lenalidomide and dexamethasone administered once weekly demonstrated efficacy in a phase 3 study (Moreau et al., 2016). Oprozomib (Fig. 1) is being evaluated in phase 1/2 clinical trials in patients with hematologic malignancy or solid tumors (Infante et al., 2016). Similar to carfilzomib, oprozomib inhibits primarily the chymotrypsin-like activity of the constitutive and immunoprotoasomes via covalent modification of the N-terminal threonine (Zhou et al., 2009). It has shown a promising clinical activity in patients with relapsed and/or refractory MM either as a single agent or in combination with pomalidomide and dexamethasone (Prada et al., 2015; Shah et al., 2015; Vij et al., 2015).

Oprozomib is a tripeptide analog (Fig. 1) of carfilzomib that was developed to improve absorption. It was believed that smaller peptides were likely to be more effectively delivered across the small intestine (Hamman et al., 2005; Zhou et al., 2009). When dosed via oral gavage to Balb/C mice at 30 mg/kg, more than 80% proteasome inhibition was observed in blood, liver, and adrenal gland 1 hour post-administration (Zhou et al., 2009). Furthermore, oprozomib demonstrated improved metabolic stability in vitro across multiple species compared with carfilzomib. Nonetheless, oprozomib still displayed a high systemic clearance and short half-life (Fang et al., 2015). This is not surprising because both the peptide backbone and the epoxyketone warhead could be susceptible to metabolism (Yang et al., 2011; Wang et al., 2013). The rapid clearance of oprozomib was primarily mediated by metabolism, supported by the observation that the parent compound was a minor component in urine and bile samples from rats, as well as urine samples from human (in-house unpublished data). In addition, oprozomib has shown a relatively large interpatient pharmacokinetic (PK) variability (Infante et al., 2016).

The objective of the present study was to characterize the major metabolic pathways and key enzymes responsible for oprozomib metabolism in vitro using cryopreserved human hepatocytes, recombinant enzymes, and liver cellular fractions in the presence and absence of selective enzyme inhibitors. Deciphering the role of various enzymes responsible for oprozomib elimination is important in drug-drug interaction (DDI) risk assessment and mitigation of exaggerated pharmacology of this potent therapeutic. Particularly, it is important for cytochrome P450 (P450)—mediated DDIs since P450 isoforms are responsible for the primary metabolism of the majority of small molecule drugs on the market (Wienkers and Heath, 2005). Defining the primary metabolic routes could also help shed light on the observed oprozomib PK variabilities in patients.

Materials and Methods

Materials. All chemicals and reagents were obtained from Sigma (St. Louis, MO) and liquid chromatography (LC) solvents were obtained from Fisher Scientific (Pittsburgh, PA). Oprozomib and authentic metabolite standards were synthesized and characterized at Onyx Pharmaceuticals, an Amgen subsidiary (South San Francisco, CA). Deuterated oprozomib (d5-oprozomib, five deuterium on the benzyl ring) was obtained from PepTech Corp. (Bedford, MA). A selective microsomal epoxide hydrolase (EH) inhibitor, 2-nonylsulfonyl-propionamide (NSPA) was previously synthesized (Morissee et al., 2008), and a selective soluble EH (sEH) inhibitor 1-trifluoromethoxyphenyl-3-(1-propionylpiperidin-4-yl) urea (TPPU) was previously characterized (Rose et al., 2010). The structures of these two inhibitors are shown in Supplemental Fig. 1. Human recombinant microsomal EH (mEH, purity 80%) and sEH (purity 95%) were prepared and characterized as described previously (Morissee et al., 2000, 2011). Supersomes containing cDNA-expressed human P450s, P450 reductase, and cytochrome b5 were purchased from BD Bioscience (San Jose, CA). Pooled human cryopreserved hepatocytes (20 donors, mixed gender) and media were purchased from BioreclamationIVT (Baltimore, MD). Liver subcellular fractions, including male or female liver microsomes, and cytosols (10 donors per gender, age: 21–58) were purchased from Sekisui Xenotech (Kansas City, KS). Pooled, gender-mixed liver microsomes were prepared by mixing an equal volume of male (10 donors) and female (10 donors) liver microsomes.

Metabolic Profiling of Oprozomib in Human Hepatocytes. Pooled cryopreserved human hepatocytes were thawed at 37°C, centrifuged at 65g in 45 mL thawing medium, and resuspended in maintenance medium. Cell viability and density were measured by a trypan blue exclusion assay. For metabolic profiling, suspensions (1.0 mL) containing hepatocytes (0.5 million cells/mL) were incubated with 10 μM of oprozomib for 1 hour. Reactions were conducted in a 37°C cell culture incubator with 100% relative humidity, 95% air, and 5% CO2. Final incubation contained 0.2% of dimethylsulfoxide (DMSO). Reactions were terminated by addition of an equal volume of acetonitrile with 0.1% formic acid (FA). Control incubation was prepared as follows: a mixture containing hepatocytes only was incubated for 1 hour, quenched with acetonitrile with 0.1% FA, and then spiked with 10 μM of oprozomib. Supernatants (600 μL) were concentrated under a nitrogen stream and then reconstituted in 100 μL of 40% acetonitrile/water (0.1% FA). Metabolite identification was conducted using a Thermo Orbitrap Q Exactive high-resolution mass spectrometer equipped with a Thermo Accela LC system from Thermo Fisher Scientific (Waltham, MA). Metabolite structures were assigned based on the accurate mass obtained from full scan mode at a resolution of 70,000 and tandem mass spectrometry (MS/MS) fragmentation pattern from data-dependent acquisition at a resolution of 17,500. For the quantification of oprozomib diol (PR-176) and (S)-2-amino-1-(4R)-2-methylxiran-2-yl)-3-phenylpropyn-1-one (PR-025) diol formation in human hepatocytes, cells (0.25 mL, 0.5 million cells/mL) were incubated with 2 μM of oprozomib in 12-well, collagen-coated plates (Thermo Scientific). At each time point (0, 10, 20, 30, and 60 minutes), reactions were terminated by the addition of an equal volume of acetonitrile containing 100 ng/mL internal standard (IS) d5-oprozomib. After centrifugation, supernatants were collected and analyzed for oprozomib and the major metabolites by LC-MS/MS as described subsequently.

Epoxide Hydrolysis by Recombinant Human mEH and sEH In Vitro. Recombinant human EHs were kept at −80°C until use. Briefly, incubations (0.1 mL) were conducted in 0.1 M Tris-HCl buffer (pH 9.0 for mEH; pH 7.5 for sEH) containing 0.1 mg/mL fatty acid-free bovine serum albumin (BSA) (Morissee et al., 2000, 2011). Enzyme activities of mEH and sEH (5 μg/mL) toward epoxide hydrolysis were confirmed using their corresponding probe substrates (50 μM) cis-stilbene oxide (cis-SO) and trans-stilbene oxide (trans-SO), respectively. To determine whether oprozomib is a substrate for recombinant mEH and/or sEH, enzymes were prewarmed for 5 minutes at 37°C before the addition of oprozomib (2 μM) to initiate the reaction. The enzyme concentration used was from 1 to 10 μg/mL for mEH and up to 100 μg/mL for sEH. After 0, 10, 20, or 30 minutes as indicated, reactions were terminated by addition of 2 volumes of acetonitrile containing 0.1% FA and supernatants were collected for LC-MS/MS analysis.

Kinetics of Oprozomib Epoxide Hydrolysis in Recombinant mEH and Human Liver Microsomes (HLMs). Reaction kinetic parameters (Vmax and Km) were determined using recombinant mEH (2 μg/mL, calculated based on 80% purity) and oprozomib (10–600 μM) in 0.1 M Tris-HCl buffer (pH 9.0) containing 0.1 mg/mL BSA. Kinetic studies in male and female HLMs were conducted in 0.1 M potassium phosphate buffer (pH 7.4) containing 0.1 mg/mL BSA. The final microsomal protein concentration was optimized to be 0.5 mg/mL and oprozomib was 10–600 μM. The final DMSO concentration was 0.5%. Oprozomib was
added to initiate the reaction after preheating enzymes for 5 minutes. Incubations (0.2 ml) were conducted for 30 minutes and terminated by addition of 2 volumes of acetonitrile containing 100 ng/ml of IS d5-oprozomib. For kinetic studies of cis-SO, incubations (0.4 ml) were optimized and performed using 4 μg/ml recombinant mEH or 0.1 mg/ml microsomal proteins and cis-SO (1–200 μM) at 37°C for 10 minutes. The reaction was terminated by addition of one volume acetonitrile containing 50 μM trans-SO as IS. Supernatants were collected by centrifugation and analyzed by either ultra-performance LC-MS/MS or ultra-performance LC linked with a photodiode array (PDA) detector as described subsequently. Each set of data was fit to a simple Michaelis-Menten kinetics model using nonlinear regression data analysis (GraphPad Prism version 6).

NSPA was used as the selective mEH inhibitor to inhibit mEH enzyme activities in HLMs (Morisseau et al., 2008). Briefly, prewarmed pooled HLMs (0.5 mg/ml) were incubated with oprozomib (10 μM) in the absence or presence of NSPA (0.02–100 μM) in 0.1 M potassium phosphate buffer (pH 7.4, 0.1 mg/ml BSA) at 37°C. After 15 minutes, reactions were quenched with 2 volumes of acetonitrile containing 0.1% FA and supernatants were isolated for quantitative analysis. The inhibitory effects of NSPA on cis-SO (50 μM) hydrolysis were also determined under similar conditions except for a lower microsomal protein concentration (0.1 mg/ml) and shorter incubation time (10 minutes). The IC50 values were estimated using nonlinear regression data analysis (GraphPad Prism version 6).

**Inhibition of Oprozomib Epoxide Hydrolysis in Human Hepatocytes in the Presence of EH Inhibitors.** Pooled cryopreserved human hepatocytes were thawed and resuspended in hepatocyte maintenance medium, and cell viability and density were measured. Incubation mixtures (0.1 ml) contained human hepatocytes (0.5 million cells/ml), oprozomib (2 μM), or cis-SO (50 μM), and inhibitors NSPA or TPPU (10 μM). The final solvent DMSO concentration was 0.2%. At each time point (0, 2, 5, 10, 20, 40, and 60 minutes), reactions were terminated by the addition of 2 volumes of acetonitrile containing either IS d5-oprozomib or trans-SO. After centrifugation, supernatants (150 μl) were collected and analyzed using LC-MS/MS or LC-PDA detection as described subsequently.

**Inhibition of P450-Mediated Oprozomib Metabolism in Human Hepatocytes by 1-Aminobenzotriazole (1-ABT).** The role of P450s on oprozomib metabolism in human hepatocytes was investigated using a pan-P450 inhibitor 1-ABT. Pooled cryopreserved human hepatocytes were thawed at 37°C, centrifuged, and resuspended in hepatocyte maintenance medium. Incubations (0.25 ml) containing hepatocytes (0.5 million cells/ml) were preincubated with 0.5 mM 1-ABT or solvent (0.1% DMSO) for 30 minutes. Oprozomib was added at a final concentration of 2 μM to initiate the reactions. Reactions were conducted in a 37°C cell culture incubator with 100% relative humidity, 95% air, 5% CO2. At each time point (0, 2, 5, 10, 20, 40, and 60 minutes), reactions were terminated by the addition of an equal volume of acetonitrile containing IS d5-oprozomib. After centrifugation, supernatants were collected for the quantification of oprozomib disappearance and PR-176 formation by LC-MS/MS.

**Metabolic Stability of Oprozomib in HLMs and Recombinant P450s.** Metabolic stability of oprozomib in pooled HLMs (mixed gender) was conducted in the presence and absence of cofactor NADPH. The value of intrinsic clearance was determined by the rate of parent compound disappearance (Obach, 1999). Briefly, oprozomib (1 μM) was mixed with liver microsomes (0.25 mg/ml) with or without 1 mM NADPH in 0.1 M potassium phosphate buffer (pH 7.4) containing 3.3 mM of MgCl2. The incubation mixtures were kept at 37°C for 1, 5, 10, 20, 30, and 40 minutes. The disappearance of oprozomib was monitored using a LC-MS/MS method described subsequently. In addition, a preliminary in vitro P450 phenotyping study was conducted by incubating oprozomib (1 μM) with six major recombinant P450 isoforms (125 pmol/ml) 1A2, 2D6, 3A4, 2C8, 2C9, and 2C19 individually, in the presence of 1 mM NADPH at 37°C for 0, 15, 30, and 60 minutes. The percentage of oprozomib remaining after incubation was calculated by comparing the peak response with that from the zero-minute incubation.

**Inhibition of P450 Activities by Oprozomib.** The relative potential of oprozomib to inhibit six major P450 isoforms was assessed in pooled HLMs (mixed gender) with and without 30-minute preincubation. The probe substrates used were phenacetin (CYP1A2), amodiaquine (CYP2C8), diclofenac (CYP2C9), S-mephenytoin (CYP2C19), dextromethorphan (CYP2D6), midazolam, and testosterone (CYP3A4/5). The incubation mixtures (0.2 ml) contained oprozomib at various concentrations (0–30 μM), a probe substrate, microsomal proteins, and 1.5 mM NADPH in 0.1 M potassium phosphate buffer (pH 7.4) with 3 mM MgCl2 for 15 minutes. To examine time-dependent inhibition, oprozomib was preincubated with liver microsomes for 30 minutes in the presence of NADPH, followed by the addition of probe substrates. The concentrations of P450 probe substrates were approximately equal to their reported Km values. The

![Fig. 2.](image-url)
micromosomal protein concentration was 0.05 or 0.1 mg/mL. The low protein concentrations were chosen to minimize possible binding of oprozomib to micromosomal proteins or lipids, as well as to reduce oprozomib metabolism via epoxide hydrolysis.

**Quantification of Oprozomib and Its Metabolites Using LC-MS/MS.**

Quantification of oprozomib and its major metabolites was performed using either an AB Sciex QTrap 4500 or 5500 mass spectrometer equipped with an electrospray ionization source (Redwood City, CA). Chromatographic separation was achieved using a Waters (Milford, MA) XBridge BEH C18 column (3.5 μm, 2.1 mm × 50 mm) and water (A)/acetonitrile (B) containing 0.1% FA as mobile phases at a flow rate of 0.5 mL/min. The mobile phase started at 10% B for 0.5 minute, linearly increased to 65% B within 1 minute, held for 0.3 minute before increasing to 95% B within 0.2 minute, held at 95% B for 1.5 minutes, and returned to 10% B within 0.2 minute. Multiple reactions monitoring of the transitions m/z 533.0 > 126.0, 551.3 > 325.2, 206.0 > 120, 224.0 > 150.0, and 538.2 > 126.1 was used to detect oprozomib, PR-176, PR-025, PR-025 diol, and IS ds-oprozomib, respectively. Calibration curves were generated by plotting the peak area ratio for each compound and IS versus the corresponding concentrations of synthetic metabolite standards, and fitting with a linear regression equation.

**Quantification of cis-SO, trans-SO, R,R-Hydroxybenzoin, and meso-Hydroxybenzoin Using LC-PDA Detection.** Formation of R,R-hydroxybenzoin and meso-hydroxybenzoin from cis-SO and trans-SO hydrolysis was used for measuring mEH and sEH activities in vitro, respectively. Quantification of these two diol products was performed using a Shimadzu LC-20AD system coupled with a PDA detector (Columbia, MD). Chromatographic separation was achieved using a Waters XBridge BEH C18 column (2.1 mm × 100 mm, 3.5 μm) and water (A)/acetonitrile (B) containing 0.1% FA as mobile phases at a flow rate of 0.3 mL/min. The mobile phase started at 10% B for 0.5 minute, linearly increased to 95% B at 5 minutes, and held for 2 minutes before returning to initial 10% B within 0.5 minute. The total running time was 9 minutes. Quantification of the diol metabolites was calculated by fitting their peak area ratio (metabolites/IS) to the calibration curves that were generated by using authentic metabolite standards R, R-hydroxybenzoin and meso-hydroxybenzoin, respectively.

**Statistical Analysis.** GraphPad Prism version 6 was used for the statistical analysis. Data represent mean ± S.D. The mean values of the measured values of each treatment group were compared using Student’s t test. Means were considered statistically different from one another if P < 0.05.

**Results**

**Oprozomib Metabolism in Human Hepatocytes.** Incubation in cryopreserved human hepatocytes with oprozomib for 60 minutes led to the formation of several metabolites, which were detected by LC-MS/MS (Fig. 2). A diol of oprozomib (PR-176) was formed from the initial incubation. It is worth mentioning that in rat hepatocyte incubations, two oprozomib conjugates via direct GSH conjugation were identified as major metabolites, in addition to PR-176 (Supplemental Fig. 2).

**Oprozomib Hydrolysis by Recombinant Human mEH but Not sEH.** Since epoxide hydrolysis was identified to be the major metabolic pathway, recombinant human mEH and sEH were used to investigate the capacity of the different EH isoforms to hydrolyze oprozomib epoxide. Formation of PR-176 increased linearly with both incubation time (up to 30 minutes, data not shown) and mEH concentration (0.5–10 μg/mL) as shown in Fig. 3A. In contrast, no significant formation of PR-176 was observed when oprozomib was incubated with recombinant sEH for 30 minutes, at enzyme concentrations up to 100 μg/mL. In a parallel experiment, both mEH and sEH can readily convert their corresponding probe substrates, cis-SO or trans-SO, respectively, in a time-dependent manner (Fig. 3B).

The kinetic properties of oprozomib epoxide hydrolysis were subsequently evaluated using recombinant mEH. The highest oprozomib concentration (up to 600 μM) was chosen based on its aqueous solubility in the phosphate buffer system. An optimized protein concentration (2 μg/mL) and incubation time (30 minutes) were applied in the study to ensure the linearity over the course of the reaction with sufficient amounts of PR-176 formation for accurate quantification. As shown in Fig. 3C, formation of PR-176 increased with oprozomib concentration up to 600 μM without reaching a plateau. Fitting the data into the Michaelis-Menten equation estimated the values of $K_m$ and $V_{max}$ of oprozomib at 974 ± 112 μM and 109 ± 8.6 nmol/min/mg proteins,

![Fig. 3. Epoxide hydrolysis by recombinant microsomal and soluble EHs. (A) Epoxide hydrolysis of oprozomib by recombinant human mEH and sEH. Formation of PR-176 was quantified after incubation of oprozomib (2 μM) with various concentrations of enzymes for 30 minutes. A pilot study has shown a linear formation of PR-176 within 30 minutes under the current conditions. (B) Epoxide hydrolysis of cis-SO and trans-SO (50 μM) by mEH and sEH (5 μg/mL), respectively. Significant diol product formation was observed after incubation for 10 or 20 minutes, confirming the hydrolysis activities of EHs. (C) Kinetics of the formation of PR-176 using recombinant mEH. A range of oprozomib concentrations were incubated with mEH (2 μg/mL) at 37°C for 30 minutes. (D) Kinetics of cis-SO hydrolysis using recombinant mEH. A series of cis-SO concentrations were incubated with mEH (4 μg/mL) at 37°C for 10 minutes. Data represent mean ± S.D. from three replicate incubations. The $K_m$ and $V_{max}$ values were estimated by fitting the curve into the Michaelis-Menten equation.](https://www.aspetjournals.org/content/doi/10.1124/dmd.117.0715)
respectively. While the values of \( K_m \) and \( V_{\text{max}} \) could not be more accurately determined under the current assay conditions, the estimation suggests a relatively high \( K_m \) value for oprozomib epoxide hydrolysis with recombinant mEH. As a control, the values of \( K_m \) and \( V_{\text{max}} \) of cis-SO were determined to be 56.9 ± 3.8 \( \mu \)M and 64.8 ± 17 nmol/min/mg proteins, respectively, which were comparable to previously reported values (Morisseau et al., 2011).

**Inhibition of Oprozomib Epoxide Hydrolysis by a Selective mEH Inhibitor in HLMs.** NSPA is a selective mEH inhibitor that has been previously characterized with recombinant rat and human mEH for its potency (Morisseau et al., 2008, 2011). Here, we evaluated its potency in inhibiting hydrolysis of cis-SO and oprozomib in HLMs. As seen in Fig. 4A, NSPA significantly inhibited cis-SO hydrolysis with a mean IC\(_{50}\) value of 0.35 \( \mu \)M, and oprozomib diol formation with a mean IC\(_{50}\) value of 0.51 \( \mu \)M in HLMs. The kinetic properties of oprozomib diol formation in both male and female HLMs were also determined (Fig. 4B). Similar to what was observed in the kinetic study with recombinant mEH (Fig. 3), formation of PR-176 essentially increased linearly with oprozomib concentration without reaching a plateau. As a result, the values of \( K_m \) and \( V_{\text{max}} \) could not be accurately estimated using nonlinear regression analysis. Considering the \( K_m \) values were likely over 5-fold higher than 50 \( \mu \)M, the apparent ratios of \( V_{\text{max}}/K_m \) values were roughly calculated from the slopes of the PR-176 formation rate over a low oprozomib concentration range (10–50 \( \mu \)M). As a result, the mean \( V_{\text{max}}/K_m \) values were estimated as 8.43 and 3.20 \( \mu \)l/min/mg protein in male and female HLMs, respectively. In addition, the mean values of \( V_{\text{max}}/K_m \) toward cis-SO and oprozomib epoxide hydrolysis in liver microsomes from rats and dogs were also estimated (Supplemental Table 1). As a control, the estimated mean \( V_{\text{max}}/K_m \) values for cis-SO were comparable to values reported in the literature (Kitteringham et al., 1996).

**Effect of the mEH Inhibitor NSPA on Oprozomib Metabolism in Human Hepatocytes.** Both NSPA and TPPU were used to further determine the importance of mEH and sEH, respectively, on oprozomib metabolism in human hepatocytes (Fig. 5). The metabolic stability of the two inhibitors, TPPU and NSPA, in human hepatocytes was first determined. TPPU was stable in human hepatocytes over a 1-hour incubation, whereas NSPA was labile with a half-time value of ~15 minutes. Thus, a concentration of NSPA (10 \( \mu \)M) 20-fold higher than the measured in vitro IC\(_{50}\) values on mEH activity was used in the human hepatocyte study. The inhibitory effect of NSPA in human hepatocytes was first confirmed using cis-SO as the probe substrate, where over 80% inhibition of cis-SO hydration was observed within 10-minute incubation; however, inhibition was reduced to 60% with a longer incubation time (up to 20 minutes; data not shown). The rate of oprozomib metabolism in human hepatocytes was significantly inhibited (>80%) by NSPA cotreatment (9.7 ± 0.9 \( \mu \)m/min/10\(^6\) cells) compared with the vehicle (DMSO) control (40.7 ± 1.4 \( \mu \)m/min/10\(^6\) cells). Consistently, formation of PR-176 was also significantly decreased (>90% inhibition) by NSPA treatment up to the 60-minute incubation (Fig. 5B). In contrast, the sEH inhibitor TPPU did not significantly inhibit oprozomib metabolism (35.0 ± 1.5 \( \mu \)m/min/10\(^6\) cells) and PR-176 formation (Fig. 5).

**Metabolic Stability of Oprozomib in HLMs in the Presence and Absence of NADPH.** Metabolic identification and quantitation results (Figs. 2 and 5) in human hepatocytes have suggested a minor role of P450s on oprozomib metabolism. HLM contains high hepatic mEH activity, and is also a common in vitro system for assessment of P450-mediated metabolism. Therefore, microsomes were expected to be a relevant in vitro system to study the metabolism of oprozomib. Interestingly, in pooled HLMs, the presence of NADPH significantly increased the intrinsic clearance of oprozomib by 3.5-fold compared with that in the absence of NADPH (70.7 ± 2.9 versus 20.2 ± 1.8 \( \mu \)l/min/mg protein) (Fig. 6A). These results indicated a significant role of oxidation pathways over epoxide hydrolysis for oprozomib metabolism in HLMs, which is not consistent with what was observed in human hepatocytes (Fig. 2).

**Oprozomib Showed Time-Dependent Inhibition of CYP3A4 in HLMs.** In vitro inhibition potential of oprozomib on major P450 isoforms (CYP1A2, 2C8, 2C9, 2C19, 2D6, and 3A4/5) was also evaluated using pooled HLMs (Table 1). Oprozomib did not inhibit the activities of CYP1A2, 2C8, 2C9, 2C19, and 2D6 at concentrations up to 30 \( \mu \)M. As a result, the estimated mean IC\(_{50}\) values were calculated using nonlinear regression data analysis. Considering the IC\(_{50}\) values were likely over 5-fold higher than 50 \( \mu \)M, the apparent ratios of IC\(_{50}/K_m\) values were roughly calculated from the slopes of the PR-176 formation rate over a low oprozomib concentration range (10–50 \( \mu \)M). As a result, the mean IC\(_{50}/K_m\) values were estimated as 20.2 and 18 \( \mu \)l/min/mg protein (data not shown). The ratios of the maximum inactivation rate constant (\( k_{\text{inac}} \)) to the concentration of the inactivator that gives half the maximum rate of inactivation (\( K_i \)) were
about 2.0 and 1.4 mM/min/µmol, respectively, with midazolam and testosterone as marker substrates. In addition, a preliminary P450 phenotyping study using recombinant P450 isoforms showed oprozomib was a substrate of CYP3A4 (Fig. 6B).

Effect of the pan-P450 Inhibitor 1-ABT on Oprozomib Metabolism in Human Hepatocytes. In this experiment, the rates of oprozomib metabolism in human hepatocytes in the absence and presence of pan-P450 inhibitor 1-ABT were compared. After preincubation with 1-ABT (0.5 mM) to inactivate P450 enzymes (in presence of NADPH. Initial oprozomib concentration was 1 µM. (B) P450 phenotyping on oprozomib oxidation using six major P450 isoforms. Oprozomib was incubated with six recombinant P450 isoforms for various incubation times, individually. The loss of oprozomib was quantified using LC-MS/MS assays. Data represent mean ± S.D. from triplicate incubations.

Discussion

Peptide epoxyketone drugs represent a new class of proteasome inhibitors that are highly efficacious for the treatment of patients with MM. Since these inhibitors act via covalent modification, they induce prolonged pharmacodynamic effects in spite of a short duration of systemic exposure (Yang et al., 2011; Wang et al., 2013). Proteasome activity recovery primarily relies on the rate of de novo proteasome synthesis, with a half-life of about 20–72 hours across a variety of tissues in preclinical species, except for blood after dosing with oprozomib (Zhou et al., 2009; Dou and Zonder, 2014). Different from the first-generation epoxyketone proteasome inhibitor carfilzomib, which is administered intravenously due to the lack of oral bioavailability, oprozomib demonstrated improved metabolic stability and was orally bioavailable (Zhou et al., 2009). However, oprozomib still displayed high systemic clearance, a short half-life, and relatively high PK variability in preclinical species and patients (Fang et al., 2015; Infante et al., 2016).

To understand the parameters governing the elimination of oprozomib, we conducted a number of in vitro metabolism studies. As summarized in Scheme 1, the in vitro metabolism data showed that hydrolysis of oprozomib epoxide to form the diol PR-176 by EH is the major metabolic pathway in human hepatocytes, with peptide cleavage by peptidases, oxidation by P450s, and direct GSH conjugation representing minor alternative metabolic pathways. Consistent with the results in human hepatocytes, PR-176 was also found to be a major circulating metabolite in dogs, the non-rodent toxicity species, as well as human plasma samples from a first-in-human clinical study (in-house unpublished data). Interestingly, GSH conjugation, an alternative pathway to detoxify xenobiotic epoxides, was minor in human hepatocytes. Although cryopreservation may reduce the in vitro GSH-conjugating capacity in both human and rat hepatocytes (Sohlenius-Sternbeck and Schmidt, 2005), two GSH conjugates of oprozomib were identified as major metabolites in addition to diol in cryopreserved rat hepatocytes. In addition, multiple GSH conjugates were observed for clozapine, a positive control for GSH adduct formation in both rat and human hepatocytes (Supplemental Fig. 3). Consistent with the in vitro observation, no detectable GSH conjugates were observed in human plasma and urine samples; on the other hand, GSH conjugates were observed in vivo in rat plasma samples (in-house unpublished data). These data indicate GSH conjugation plays a minor role in oprozomib metabolism in humans.

The predominant role of EHs on oprozomib metabolism could have implications on potential DDIs and therefore may influence development strategy. Several EHs are found in mammals; among them, two major isoforms, mEH and sEH, have been extensively characterized (Freitland and Omiecinski, 2000; Kodani and Hammock, 2015). Both

### TABLE 1

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>P450 Reaction</th>
<th>IC₅₀</th>
<th>Inhibition at 30 µM</th>
<th>Time Dependent (30-Minute Preincubation)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>µM</td>
<td>%</td>
<td>µM</td>
</tr>
<tr>
<td>CYP1A2</td>
<td>Phenacetin O-deethylation</td>
<td>&gt;30</td>
<td>21</td>
<td>&gt;30</td>
</tr>
<tr>
<td>CYP2C8</td>
<td>Amiodarone N-dealkylation</td>
<td>&gt;30</td>
<td>20</td>
<td>&gt;30</td>
</tr>
<tr>
<td>CYP2C9</td>
<td>Diclofenac 4’-hydroxylation</td>
<td>&gt;30</td>
<td>6.3</td>
<td>&gt;30</td>
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<tr>
<td>CYP2C19</td>
<td>S-Mephentynol 4’-hydroxylation</td>
<td>&gt;30</td>
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<td>&gt;30</td>
</tr>
<tr>
<td>CYP2D6</td>
<td>Dextromethorphan O-demethylation</td>
<td>&gt;30</td>
<td>0.2</td>
<td>&gt;30</td>
</tr>
<tr>
<td>CYP3A4/5</td>
<td>Testosterone 6β-hydroxylation</td>
<td>&gt;30</td>
<td>45</td>
<td>5.5</td>
</tr>
<tr>
<td>CYP3A4/5</td>
<td>Midazolam 1’-hydroxylation</td>
<td>&gt;30</td>
<td>33</td>
<td>12</td>
</tr>
</tbody>
</table>

NA, not applicable.

*The inhibition (%) was calculated using the data for the highest concentration of test article evaluated and the following formula: inhibition (%) = 100% − % activity of remaining substrate of control.

*No value was obtained since the rates at the highest concentration of oprozomib evaluated (30 µM) were higher than the control rates.
isoforms are expressed ubiquitously throughout the mammalian body with the highest expression generally in the liver (Gill and Hammock, 1980; Pacifici et al., 1988; Coller et al., 2001). Collectively, data from recombinant EHs (Fig. 3), liver microsomes (Fig. 4), and hepatocytes in the presence or absence of selective inhibitors (Fig. 5) indicate that the epoxide ring opening of oprozomib was catalyzed by mEH, but not sEH. Therefore, oprozomib PK is unlikely to be affected by co-administration with sEH inhibitors. On the other hand, oprozomib is primarily metabolized by mEH. Some amide-containing drugs such as valpromide or progabide have been shown to inhibit in vivo mEH activity (Fretland and Omiecinski, 2000; Kodani and Hammock, 2015). Thus, further studies may be warranted to determine whether drugs that are mEH inhibitors cause clinically significant DDIs with oprozomib. Moreover, mEH is involved in the metabolism of numerous xenobiotics, including 1,3-butadiene, naphthalene, carbamazepine, and phenytoin (Decker et al., 2009). It is possible that oprozomib acts as a mEH inhibitor that affects the clearance of coadministered drugs, which are mainly metabolized by mEH. However, a preliminary in vitro study in HLMs showed that oprozomib does not significantly inhibit \( \text{cis}-\text{SO epoxide hydrolysis at a concentration up to 100 } \mu\text{M (data not shown).} \)

The high variability of oprozomib PK observed in preclinical species and patients could be attributed to variability of absorption and/or metabolism by mEH. Expression of mEH in ex vivo tissues was shown to vary considerably among individuals (up to 8-fold) (Hassett et al., 1997), and that expression is affected by age, sex, diseases, and environmental factors (Václavíková et al., 2015). In addition, two human polymorphisms have been identified in the coding region of the mEH gene: exon 3 (a Tys113His mutation), resulting in a 50% decrease in enzyme activity in vitro, and exon 4 (a His139Arg mutation), resulting in a 25% increase in enzyme activity in vitro (Pinarbasi et al., 2010).

![Scheme 1. Proposed major metabolic pathways of oprozomib in human hepatocytes. Oprozomib is predominantly metabolized by mEH in human hepatocytes, whereas P450s, peptidases, and glutathione S-transferase (GST)/GSH play minor roles. Oxidative and cleavage products containing an epoxyketone could be further metabolized via epoxide hydrolysis.](image-url)
However, the variant alleles appeared to exert only a modest impact on mEH enzymatic activities (within 2-fold) in vivo (Kroetz et al., 1993; Hosagrahara et al., 2004). Based on this information and the high systemic clearance of oprozomib, interindividual difference in mEH activity may contribute to the high interindividual PK variability of oprozomib in patients (Infante et al., 2016), but it is unlikely to be the main factor. Variable absorption, particularly associated with a narrow absorption window, may be another key factor in oprozomib PK variability (Teague et al., 2013).

In contrast to the major role of mEH, a minor role of P450 on oprozomib metabolism was observed in human hepatocytes (Fig. 7). Using the pan-P450 inhibitor 1-ABT, a modest decrease in oprozomib disappearance (<20%) was observed. This corroborates the result that oxidative products are minor metabolites in human hepatocytes (Fig. 2), as well as in plasma samples from patients (in-house unpublished data). Thus, although oprozomib is a substrate of CYP3A4, coadministration of oprozomib with other drugs that are P450 modulators would be unlikely to significantly alter the PK of oprozomib. In addition, oprozomib did display time-dependent inhibition of CYP3A4/5 in HLMs (Fig. 6); however, the inhibitory effect was not manifested in vivo in a clinical DDI study where oprozomib was administered orally and coadministered with midazolam (Tsimberidou et al., 2016).

Choosing relevant in vitro systems is critical in understanding the metabolism and associated DDI potentials for oprozomib and other peptide epoxyketone inhibitors. When studying oprozomib metabolism in HLMs (Fig. 6), a higher level of P450-mediated metabolism was observed in contrast to the results observed in human hepatocytes (Fig. 7). Moreover, the high levels of oxidative metabolites in HLMs were also not consistent with the in vivo human metabolite profiling results (in-house unpublished data). The distinct observation of oprozomib metabolism in HLMs versus human hepatocytes might be due to significant loss of mEH activity during microsomal subcellular fractionation. In hepatocytes, besides the endoplasmic reticulum membrane, mEH is also present on the plasma membrane facing the extracellular medium (Levy, 1996), thus a significant portion of the hepatocyte mEH activity might be lost when preparing microsomes. On the other hand, P450 activities are concentrated in the endoplasmic reticulum membrane, which could explain why HLM incubation may bias metabolism toward P450 oxidation.

Furthermore, it is worth discussing the limitations of using hepatocytes in evaluation of the metabolism and extrapolation of the in vivo clearance of peptide epoxyketone analogs. Hepatocytes provided consistent metabolic profiles for oprozomib with those from in vivo samples. However, the rates of oprozomib disappearance and diol formation from in vitro hepatocyte incubation would underpredict the in vivo clearance of oprozomib, which was measured higher than hepatic blood flow in preclinical species (Fang et al., 2015). There are several possible reasons. First, the rate of diol formation obtained from in vitro kinetic incubation may underestimate the efficiency of mEH. For EH-mediated hydrolysis, the epoxide is rapidly trapped as a covalent hydroxyl alkyl enzyme intermediate. The rate of diol formation is limited by the slow hydrolysis of the intermediate (Armstrong, 1999). Second, the activity of epoxide hydrolases is generally not well characterized in commercial preparations. The effects of reagents and in vitro preparation procedure on epoxide hydroxylase activity have not been systematically studied to the best of our knowledge. Third, extrahepatic metabolism also plays an important role in the elimination of this series of compounds because EHS and peptidases are ubiquitously expressed throughout the body. Therefore, the rate of metabolism in hepatocytes could be used simply as a tool to rank order analogs for prioritizing in vivo studies instead of extrapolating clearance during lead optimization.

In summary, in vitro characterization of oprozomib metabolism revealed that epoxide hydrolysis mediated by mEH, rather than P450, is the major metabolic pathway for oprozomib metabolism, consistent with findings from in vivo samples. Oxidative metabolism plays a minor role in oprozomib elimination. These data suggest P450 and SEH activity modulators when coadministered with oprozomib would unlikely significantly affect oprozomib PK. On the other hand, the results in the present study indicate that mEH inhibitors administered concomitantly with oprozomib may affect oprozomib PK. Further studies may be warranted to determine whether drugs that are mEH inhibitors cause clinically significant DDIs with oprozomib. Hepatocytes serve as a good in vitro system to assess the metabolic profiles of peptide epoxyketone inhibitors but generally underestimate the rate of in vivo clearance of oprozomib. Metabolic findings using liver microsomes should be interpreted with caution since the microsomal system may be biased toward P450-mediated metabolism and may underestimate non-P450 metabolism for peptide epoxyketones.

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Wrote or contributed to the writing of the manuscript: Zhi. Wang, Fang, Morisseau, Hammock, Rock, Zhe. Wang.

References