Molecular Determinants of Ligand Selectivity for the Human Multidrug And Toxin Extrusion Proteins, MATE1 and MATE2-K

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Abbreviations: MATE, Multidrug And Toxin Extruder; RPT, renal proximal tubule; OC, organic cation; MPP, 1-methyl-4-phenylpyridinium; OCT, organic cation transporter.

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Abstract

The present study compared the selectivity of two homologous transport proteins, Multidrug And Toxin Extruders 1 and 2-K (MATE1 and MATE2-K), and developed 3D pharmacophores for inhibitory ligand interaction with hMATE1. The human orthologs of MATE1 and MATE2-K were stably expressed in Chinese Hamster Ovary cells and transport function was determined by measuring uptake of the prototypic organic cation (OC) substrate 1-methyl-4-phenylpyridinium (MPP). Both MATEs had similar apparent affinities for MPP, with $K_{\text{app}}$ values of 4.4 µM and 3.7 µM for MATE1 and MATE2-K, respectively. Selectivity was assessed for both transporters from IC$_{50}$ values for 59 structurally diverse compounds. Whereas the two transporters discriminated markedly between a few of the test compounds, the IC$_{50}$ values for MATE1 and MATE2-K were within a factor of three for most of them. For hMATE1 there was little or no correlation between IC$_{50}$ values and the individual molecular descriptors LogP, total polar surface area, or pK$_{a}$. The IC$_{50}$ values were used to generate a common features pharmacophore, quantitative pharmacophores for hMATE1, and a Bayesian model suggesting molecular features favoring and not favoring the interaction of ligands with hMATE1. The models identified hydrophobic regions, H-bond donor and H-bond acceptor sites, and an ionizable (cationic) feature as key determinants for ligand binding to MATE1. In summary, using a combined in vitro and computational approach, MATE1 and MATE2-K were found to have markedly overlapping selectivities for a broad range of cationic compounds, including representatives from seven novel drug classes of FDA-approved drugs.
Introduction

A key physiological function of the kidneys is clearing the body of a structurally diverse array of organic compounds, the majority of which are exogenous, i.e., xenobiotic, in origin. These include plant-derived compounds found in typical diets and, increasingly, clinically relevant synthetic pharmaceuticals. So-called ‘organic cations’ (OCs), i.e., molecules that carry a net positive charge at physiological pH, are a particularly significant subset of pharmaceuticals as they make up about 40% of all prescribed drugs (e.g., cimetidine, procainamide, pindolol and metformin) (Neuhoff, et al., 2003). The basic cellular model of renal OC secretion in renal proximal tubule (RPT) cells, described first by Holohan and Ross (1981), includes the sequential activity of (i) a basolateral ‘entry step,’ from blood to cell, that involves an electrogenic organic cation transporter; and (ii) an apical ‘exit step,’ from cell to tubular filtrate (that is both the active and rate-limiting step in secretion; Wright and Dantzler, 2004), mediated by electroneutral OC/H+ exchange. Following the cloning in 1994 of the first organic cation transporter, OCT1 (Gründemann, et al., 1994), there is now a broad consensus that, in the human kidney, the basolateral step in this process is dominated by activity of organic cation transporter 2, (OCT2; Wright and Dantzler, 2004; Motohashi, et al., 2002). However, establishing the molecular basis of the apical element in renal OC secretion, i.e., OC/H+ exchange, proved to be more elusive.

The cloning in 2005 of the first mammalian members of the Multidrug And Toxin Extrusion (MATE) family of transport proteins (Otsuka, et al., 2005) provided the first viable candidates for the molecular identity of the apical OC/H+ exchanger. MATE1 and
MATE2 proved to display the ‘physiological fingerprint’ of the apical element of renal (and hepatic) OC secretion: (i) substantial expression in the luminal membrane of RPT cells (and, for MATE1, canilicular membrane of hepatocytes); (ii) support of OC/H+ exchange; and (iii) transport of structurally diverse OCs. The quantitative link between MATE activity and renal OC secretion was then firmly established by the observation that elimination of Mate1 in mice significantly reduces renal clearance of metformin (Tsuda, et al., 2009) and cephalexin (Watanabe, et al., 2010).

A primary focus of studies of MATE function has been establishing the interaction of MATE transporters (typically MATE1) with specific structural classes of drug (e.g., Cutler, et al., 2011; Yokoo, et al., 2007; Ohta, et al., 2009; Watanabe, et al., 2010). However, lacking in these observations is an effort to identify the molecular determinants of ligand (substrate/inhibitor) interaction with MATE transporters, including establishing the differential selectivity of MATE1 versus MATE2 (or with its kidney-specific isoform, MATE2-K; Masuda, et al., 2006; Komatsu, et al., 2011). The multispecificity of the MATEs makes them important potential targets for unwanted drug-drug interactions (Yonezawa and Inui, 2011), so understanding the transport mechanisms that underlie the processes of renal and hepatic OC clearance, including the determinants of selectivity, is particularly relevant to efforts to predict and pre-empt the unwanted outcomes of drug exposure.

Previously, combining in vitro data with computational modeling of transporters enabled the development of pharmacophores and quantitative structure/activity relationships (QSARs) that have facilitated understanding the molecular basis of ligand interaction with transport proteins (e.g., Zolk, et al., 2008; Kido, et al., 2011; Bednarczyk,
et al., 2003; Suhre, et al., 2005). In the current study we used sequential rounds of pharmacophore development and searching of a comprehensive set of FDA approved drugs to: (i) characterize the relative selectivity of MATE1 and MATE2-K for a set of clinically important OCs; (ii) identify novel inhibitors of these two transporters; and (iii) develop initial predictive models of MATE1 selectivity using an in vitro/in silico (IVIS) method that involves successive, iterative steps in the model-building process.
Methods

Reagents. Platinum® High Fidelity DNA polymerase, Zeocin, hygromycin, Flp recombinase expression plasmid (pOG44), Chinese Hamster Ovary cells containing a single integrated Flp Recombination Target (FRT) site (CHO Flp-In), and the mammalian expression vector pcDNA5/FRT/V5-His TOPO, were obtained from Invitrogen Corporation (Carlsbad, CA). Ham’s F12 Kaign’s modification cell culture medium was obtained from Sigma Chemical (St. Louis, MO), as were the test inhibitors of MATE transport activity. [3H]1-Methyl-4-phenylpyridinium ([3H]MPP; 80 Ci/mmol) was synthesized by the Department of Chemistry and Biochemistry, University of Arizona.

Cell culture and stable expression of hMATE1 and hMATE2-K. The full length human MATE1 sequence used in this study was generously provided by Dr. Kathleen Giacomini (UC, San Francisco; Chen, et al., 2007). The full length human MATE2-K sequence used in this study was generously provided by Dr. Ken-ichi Inui (Kyoto University; Masuda, et al., 2006). Chinese Hamster Ovary cells (CHO) containing the Flp recombination target site were grown in Ham’s F12 Kaign’s modification medium supplemented with 10% fetal calf serum and 100 µg/ml Zeocin. Cultures were split every 3 days. Aliquots of 5 x 10^6 cells were transfected by electroporation (BTX ECM 630, San Diego; 260 volts and time constant of ~25 ms) with 10 µg of salmon sperm, 18 µg of pOG44, and 2 µg of pcDNA5/FRT/V5-His TOPO containing the open reading frame of either hMATE1 or hMATE2-K construct. Cells were seeded in a T-75 flask following transfection and maintained under selection pressure with hygromycin (100 µg/ml) for at least two weeks before use in transport studies.
**Transport experiments.** CHO cells expressing hMATE1 or hMATE2-K were grown to confluence in multi-well (typically 24-well) plates. Prior to transport experiments, the media was aspirated and the cells rinsed twice, briefly, with room temperature Waymouth buffer (WB) containing in mM: 135 NaCl, 28 D-glucose, 5 KCl, 1.2 MgCl₂, 2.5 CaCl₂, 0.8 MgSO₄, and 13 HEPES-NaOH, pH 8.5. Transport was measured at room temperature and was initiated by adding transport solution containing WB with 1 μCi/ml[^3]H]MPP (~10-20 nM) and, in studies assessing the kinetics of transport, increasing concentrations of unlabeled substrate or inhibitor. To reduce the inhibitory effect of extracellular H⁺ on MATE transport activity the pH of the transport buffer in these studies was typically 8.5 (the impact of pH on the kinetics of MATE transport is discussed in the Results section). The solubility of some of the test agents required that stock solutions be prepared in dimethyl sulfoxide (DMSO), ethanol or methanol resulting in the presence of these solutes in some experimental solutions. Preliminary experiments revealed that 2% DMSO, ethanol or methanol (the highest concentration required to maintain solubility of selected agents) reduced the rate transport of[^3]H]MPP by up to 40%. Consequently, for those agents that required the presence of DMSO, ethanol or methanol in the test solutions, all solutions, including parallel control experiments (when no inhibitor was present) contained 2% DMSO, ethanol or methanol in WB. Since initial experiments showed that uptake of[^3]H]MPP was linear for ~10 min (see Dangprapai and Wright, 2011), 5 min uptakes were used to approximate the initial rate of transport for use in kinetic analyses. After the transport period, the solution was aspirated and the wells were rinsed three times with 1 ml of ice-cold WB. The cells were solubilized in 0.2 ml of 0.5N NaOH with 1% SDS (v/v), and the resulting lysate was
neutralized with 0.1 ml of 1N HCl. Accumulated radioactivity was determined by liquid scintillation spectrometry (Beckman model LS3801). Mediated (i.e., inhibitable) accumulation of [3H]MPP into wild-type CHO cells is typically less than 2% of the mediated uptake into cells stably expressing MATE1 (Zhang and Wright, 2009) and, so, was ignored. Individual transport observations were typically performed in duplicate for each experiment, and observations were usually confirmed at least three times in separate experiments using cells of a different passage.

**Physicochemical descriptors.** LogP (log of the octanol–water partition coefficient) values were calculated with the ALOGPS 2.1 package (Tetko, et al., 2005), which displays values calculated with ALOGPs, MLogP (Moriguchi octanol–water partition coefficient, Dragon 5.4, Talete, Milano, Italy), ALogP (Ghose–Crippen octanol–water partition coefficient, Dragon 5.4, Talete, Milano, Italy), and KowWin logP (Syracuse Research Corporation, Syracuse, NY, USA). The average value of these LogP calculations was used for our analysis. The pK_a values were calculated by the SPARC On-Line Calculator ([http://archemcalc.com/sparc](http://archemcalc.com/sparc)) (Hilal, et al., 1996). Topological polar surface area (TPSA) was calculated using the Interactive PSA calculator ([http://www.molinspiration.com/services/psa.html](http://www.molinspiration.com/services/psa.html)).

**Computational modeling**

**Common features hMATE1 pharmacophore development.** A common features pharmacophore was developed using Accelrys Discovery Studio vers 2.5.5. (San Diego, CA) following the approach taken previously with other transporters (e.g., Diao, et al., 2010; Diao, et al., 2009). Template molecule structures were downloaded from ChemSpider ([www.chemspider.com](http://www.chemspider.com)), and conformer generation was carried out using
the CAESAR algorithm (Conformer Algorithm based on Energy Screening and Recursive Buildup; Li, et al., 2007) applied to the selected template molecules (maximum of 255 conformations per molecule and maximum energy of 20 kcal/mol).

3D-QSAR development used the Hypogen method in Discovery Studio. hMATE1 IC₅₀ values were used as the indicator of biological activity. In the HypoGen approach (Bednarczyk, et al., 2003; Ekins, et al., 2002), ten hypotheses were generated using hydrophobic, hydrogen bond acceptor, hydrogen bond donor, and the positive and negative ionizable features, and the CAESAR conformer generation method was used. After assessing all generated hypotheses, the hypothesis with lowest energy cost was selected for further analysis, as this model possessed features representative of all the hypotheses and had the lowest total cost. The total energy cost of the generated pharmacophore was calculated from the deviation between the estimated activity and the observed activity, combined with the complexity of the hypothesis (i.e. the number of pharmacophore features). A null hypothesis, which presumed that there was no relationship between chemical features and biological activity, was also calculated. Therefore, the greater the difference between the energy cost of the generated and null hypotheses, the less likely the generated hypothesis reflects a chance correlation. Also, the quality of the structure-activity correlation between the predicted and observed activity values was estimated via correlation coefficient.

**Quantitative model update with variable weights and tolerances.** We selected hydrogen bond acceptor, hydrogen bond donor, hydrophobicity, positive ionizable and negative ionizable features for model building (again using CAESAR for conformation generation). Variable weights and tolerances were employed and a
maximum of 10 pharmacophores were selected. The pharmacophore with the best correlation (lowest RMS error) was used for further analysis.

Classification Bayesian models. Laplacian-corrected Bayesian classifier models were generated using Discovery Studio. Molecular function class fingerprints of maximum diameter 6 (FCFP_6), AlogP, molecular weight, number of rotatable bonds, number of rings, number of aromatic rings, number of hydrogen bond acceptors, number of hydrogen bond donors, and molecular fractional polar surface area were calculated from input sdf files using the “calculate molecular properties" protocol. The “create Bayesian model" protocol was used for model generation (Diao, et al., 2010).
Results

The kinetics of MPP and H⁺ interaction with hMATE1 & hMATE2-K. The kinetics of MATE1- and MATE2-K-mediated MPP transport (Figure 1A, 1B) were adequately described by the Michaelis-Menten equation for competitive interaction of labeled and unlabeled substrate (Malo and Berteloot, 1991).

\[ J^* = \frac{J_{\text{max}} [S^*]}{K_{\text{tapp}} + [S^*] + [S]} + D_{\text{ns}} [S^*] \]

where \( J^* \) is the rate of transport of the radiolabeled substrate (in this case, \([^3\text{H}]\text{MPP}\)) from a concentration of the labeled substrate equal to \([S^*]\); \( J_{\text{max}} \) is the maximal rate of mediated substrate transport; \( K_{\text{tapp}} \) is the apparent Michaelis constant of the transported substrate; \([S]\) is the concentration of unlabeled substrate; and \( D_{\text{ns}} \) is a first-order rate constant that describes the nonsaturable component of labeled substrate accumulation (reflecting the combined influence of diffusion, nonspecific binding, and incomplete rinsing of \([^3\text{H}]\text{MPP}\) from the cell culture well). In four separate experiments \( K_{\text{tapp}} \) and \( J_{\text{max}} \) values for MATE1 and MATE2-K, respectively, were 4.37 ± 0.32 µM and 3.72 ± 0.45 µM; with \( J_{\text{max}} \) values of 2.14 ± 0.27 and 1.86 ± 0.28 pmol cm⁻² min⁻¹, respectively. Expressed per mg of membrane protein, these \( J_{\text{max}} \) values become 21.4 and 18.6 pmol mg⁻¹ min⁻¹ for MATE1 and MATE2-K, respectively.

It is important to acknowledge that MATEs are sensitive to the intra- and extracellular concentration of protons and therefore we characterized the kinetics of H⁺ inhibition of \([^3\text{H}]\text{MPP}\) uptake. We previously showed that elevated concentrations of H⁺ in the extracellular solution inhibit transport mediated by hMATE1 (Dangprapai and Wright, 2011). Figure 1C, D compares the pH sensitivity of MPP transport mediated by hMATE1 and hMATE2-K. As anticipated, transport activity of both proteins was
inhibited by increasing concentrations of H⁺ in the extracellular solution, and was described by the following relationship:

\[
J^* = \frac{J_{\text{app}}[S^*]}{IC_{50} + [H^+]_o} + D_{ns}[S^*]
\]

eq 2

where \(J_{\text{app}}\) is the product of the maximum rate of \(S^*\) (i.e., \([3H]MPP\) uptake (\(J_{\text{max}}\)) and the ratio of the \(K_i\) (\(IC_{50}\)) of H⁺ and \(K_{\text{tapp}}\) for MPP transport; and \(IC_{50}\) is the concentration of [H⁺]₀ (or other test inhibitor) that reduced mediated (i.e., blockable) [3H]MPP transport by 50%. In three experiments, the \(IC_{50}\) for H⁺-inhibitable hMATE1-mediated MPP uptake was 19.6 ± 0.7 nM (pH 7.73), similar to the value of 12.4 nM reported previously (Dangprapai and Wright, 2011). MATE2-K proved to be substantially more sensitive to H⁺, displaying an \(IC_{50}\) value of 3.5 ± 0.6 nM (pH 8.5, n=3).

In the upcoming examination of the kinetics of ligand interaction with MATE transporters we elected to maximize control rates of MATE-mediated transport by running experiments at an external pH of 8.5, rather than employ the ‘ammonia pre-pulse’ method to acidify the cytoplasm and thereby create an outwardly-directed pH gradient and a stimulation of OC uptake (e.g., Yasujima, et al., 2010;Tanihara, et al., 2007;Kajiwara, et al., 2007;Otsuka, et al., 2005;Masuda, et al., 2006). The acidification of the cytoplasm following an ammonia pulse is generally short-lived and constantly changing (e.g., Kapus, et al., 1994) during the several minute time courses used to measure the rate of MATE-mediated transport, and these ill-defined conditions complicate the interpretation of kinetic measurements. Importantly, we previously showed that cytoplasmic pH is effectively constant (at a \(pH_{in}\) of 7.5-7.6) during exposure of CHO cells to an external pH of 8.5 (Dangprapai and Wright, 2011), so transmembrane H⁺ gradients were both (i) outwardly-directed and (ii) unchanging during
our transport measurements. The rank-order of ligand selectivity at pH 8.5 and 7.4 is similar, if not identical, for the two transporters, as supported by the similar rank order of uptake ratios for transport of a structurally diverse set of organic cations into hMATE1 and hMATE2-K at these two pH values (Tanihara, et al., 2007). However, given the apparent pKa values for the interaction of the two MATE transporters with H+ evident in Figure 1B, the absolute IC50 values for inhibition of MATE1 and MATE2-K activity measured at pH 8.5 can be expected to underestimate the values anticipated at pH 7.4, by approximately 3- to 6-fold, respectively.

**Inhibitory selectivity of hMATE1 and hMATE2-K test set selection.** The test compounds (Table 1) were selected to represent a structurally diverse collection of drug and drug-like compounds, the intention being to interrogate the complex chemical space expected to influence interaction with the binding region(s) of multiselective organic cation transporters, i.e., MATE1 and MATE2-K. Weak bases and cations dominated the battery of test compounds; neutral compounds and those having a net negative charge at physiological pH were largely excluded. This ‘bias’ toward cations reflected the existing database from the early literature on transport in isolated renal membranes (Holohan and Ross, 1981; Holohan and Ross, 1980; Wright, et al., 1995) and in intact renal tubules (McKinney, 1983; Dantzler, et al., 1989; David, et al., 1995), and from more recent work with MATE transporters (Tanihara, et al., 2007), showing that cationic charge is a key criterion of ligand interaction with these processes (a conclusion supported by the present study, as documented below). Indeed, it was a specific goal of this study to identify molecular determinants of interaction of organic cations with MATE1 and MATE2-K. The final battery of organic compounds (Table 1) included 23
from the list of compounds generated by Ahlin et al. (Ahlin, et al., 2008), in their study of selectivity of OCT1; 13 selected because of previous evidence of their interaction with OC/H⁺ exchange activity in either native renal membranes, intact tubules, BBMV or heterologous expression systems expressing MATE1 or MATE2-K (Tanihara, et al., 2007; Wright and Wunz, 1999; Wright and Wunz, 1998; Wright, et al., 1995; Ullrich and Rumrich, 1996; David, et al., 1995); and 23 selected from lists of target ‘hits’ from databases of compounds that were interrogated by the pharmacophore model(s) developed during the course of this study.

Figure 2 shows the range of inhibition of MATE-mediated transport activity produced by the 59 organic test compounds. At inhibitor concentrations of 10 μM, MPP transport was reduced by ≥50% by 20 (MATE1) or 14 (MATE2-K) of these compounds. Figure 3 shows inhibitory profiles against transport activity of MATE1 and MATE2-K produced by four compounds (quinidine, agmatine, nialamide and allopurinol) selected to emphasize the spectrum of inhibitory effectiveness of this battery of test agents, with IC₅₀ values (determined using equation 2) that ranged from low micromolar (e.g., quinidine), through near millimolar (e.g., nialamide), to no effective interaction at all (e.g., allopurinol). Table 1 lists the IC₅₀ values (as measured at pH 8.5, and as calculated for pH 7.4 using the apparent IC₅₀ values for H⁺ interaction with the transporters shown in Figure1C/D) for all 59 organic molecules used to inhibit transport activity of one or both MATE transporters.

As inferred from the results presented in Figures 1 through 3, there was substantial overlap in the interaction of the test compounds with MATE1 and MATE2-K. The extent of this overlap is evident in the comparison of the MATE1 and MATE2-K IC₅₀
values for the 59 organic molecules used to probe both transporters (Figure 4A); 75% of these agents had IC$_{50}$ values for the two transporters that differed by less than a factor of 3 at pH 8.5 (61% of the compounds, based on the calculated IC$_{50}$ values at pH 7.4). MATE1 and MATE2-K did, however, markedly discriminate (ratio of IC$_{50}$ values >5) between 8 of the compounds examined in this study. For example, as shown in Figure 4B, the apparent affinities of hMATE1 for atropine (IC$_{50}$ of 5.90 ± 1.31 µM) and amantadine (7.50 ± 1.49 µM) were ~10-times greater than those displayed by MATE2-K (52.8 ± 13.7 µM and 88.9 ± 9.0 µM, for atropine and amantadine, respectively); whereas the apparent affinity of hMATE2-K for azidoprocainamide (Mol, et al., 1989; APMI; IC$_{50}$ of 0.50 ± 0.22 µM) was ~10 times greater than that displayed by hMATE1 (6.2 ± 0.3 µM; Figure 4B). Despite these differences, the data in hand support the view that MATE1 and MATE2-K show far more similarities in selectivity, than differences.

Modeling of MATE1 selectivity

Influence of selected molecular descriptors on MATE1-mediated transport activity. Figure 5 shows the log of the IC$_{50}$ values for inhibition of hMATE1 activity graphed as a function of several commonly applied molecular descriptors for the test agents in this study. There was a significant, albeit weak correlation between hMATE1 IC$_{50}$ values and LogP ($r$ value of 0.332, $p < 0.05$; Figure 5A). Interestingly, when the IC$_{50}$ values for inhibition of MATE1 were restricted to a structurally constrained subset of the test agents of the present study, i.e., an n-tetraalkylammonium series, the influence of LogP was more evident ($r = 0.97$ for TEA through TPeA; Figure 5B). There was no correlation between TPSA and hMATE1 IC$_{50}$ ($r$ value of 0.045, $p > 0.05$; Figure 5C), and
a modest, albeit significant, correlation between pK<sub>a</sub> and hMATE1 IC<sub>50</sub> values (r = 0.423, p < 0.01; Figure 5D).

**Computational analysis of hMATE1 inhibition.** The value of the approach offered by computational assessment of structure/activity relationships (SARs) is that it may enable more insight into the molecular basis of ligand interaction than a view restricted to the influence of single physicochemical parameters alone. Following previous in silico modeling efforts on drug transporters, (e.g., Diao, et al., 2010; Diao, et al., 2009; Suhre, et al., 2005) we extended our in vitro/in silico (IVIS) strategy in this study the use of multiple iterations of model development. The IVIS approach uses a comparatively small amount of in vitro data for development of an initial model that is then used to screen a database of potential additional compounds for testing. The results obtained from these initial tests feed into a further round(s) of model building and database searching, and so on. The advantage of such an approach is the model is co-developed with data acquisition, and is validated and tuned with each additional set of test compounds. The approach does not require a large library of compounds to be tested and can save reagents and money (associated with testing many inactive compounds). Using this pharmacophore approach may suggest non-intuitive compounds as inhibitors because while they include one or more of the initially mapped features, they may still prove to be low affinity ligands (owing to the absence of what proves to be a missing critical feature). This in particular may be another potentially valuable side effect of the approach, enabling us to find novel compounds that may have a dissimilar 2D structure, but similar 3D shape to known inhibitors or different mapping to pharmacophore features.
**Initial Round: hMATE1 Common Features Pharmacophore.** There were 26 compounds in the initial round of inhibitors studied (Table 1, underlined), from which five were selected to generate a common features pharmacophore: two high affinity compounds, pyrimethamine (IC\textsubscript{50} of 0.04 µM) and quinidine (IC\textsubscript{50} of 1.57 µM) and three low affinity compounds, histamine (IC\textsubscript{50} of 761 µM), caffeine (IC\textsubscript{50} of 1096 µM) and chloramphenicol (IC\textsubscript{50} of 1115 µM; Figure 6A). By restricting the initial set of test compounds to these extremes (i.e., very high affinity versus very low affinity), the intent was to identify key features that may influence effective interaction with the transporter. In other words, common molecular and chemical features of the high affinity substrates were included in the pharmacophore, whereas the molecular features of the low affinity substrates were excluded from the pharmacophore. The resulting 'common features pharmacophore' had 2 hydrophobic regions (cyan), 1 H-bond donor (magenta), and 1 H-bond acceptor (green). The pharmacophore is depicted an overlay of the structure of pyrimethamine (PYR; the highest affinity substrate; Figure 6B).

**1st Iteration: hMATE1 Common Features Pharmacophore Testing.** The common features pharmacophore, with the van der Waals surface of PYR to provide a shape restriction (note: the 2D molecule structures and Discovery Studio pharmacophores are available upon request from the authors), was used to search a 3D database of 2690 FDA approved compounds (www.collaborativedrug.com), and identified 126 molecules as potential inhibitors (see Supplemental Table 1, which is available online). Fifteen commercially available compounds were selected from this list and tested as inhibitors of MATE1 (and MATE2-K), and the resulting IC\textsubscript{50} values are presented in Table 1 (identified with the superscript a). Nine compounds in this test set proved to be
comparatively high affinity inhibitors of MATE1 (IC$_{50}$ values of 0.9-25 µM), whereas four displayed modest affinity (IC$_{50}$ values of 26-300 µM), two had weak interactions (<80% inhibition at 1000 µM) and one exerted no inhibition (NI) of either transporter at a concentration of 1 mM.

The inhibitory profiles produced by two of the 15 compounds in this ‘test set,’ cinchonidine and ethohexadiol (Figure 7A, B), provided particular insight into the molecular determinants associated with ligand interaction with MATE1. Although both molecules were good fits for the common features pharmacophore (Figure 7C, D; fit values of 3.1 and 2.7, respectively, data not shown), cinchonidine was an effective, high affinity inhibitor of hMATE1 (IC$_{50} =$ 0.93 µM; Figure 7A), whereas ethohexadiol (Figure 7B) had a weak interaction with hMATE1 (~20% inhibition at a concentration of 1 mM). Thus, the presence in ethohexadiol of the common structural features of the initial pharmacophore, i.e., the 2 hydrophobic regions, 1 H-bond donor site and 1 H-bond acceptor site, was not sufficient for an effective inhibitory interaction with MATE1. It is noteworthy that ethohexadiol is not an organic cation; its presence in the ‘hit list’ reflected the absence of a cationic feature in the common features pharmacophore that, in turn, reflected the presence of such a feature in all three of the ‘weak inhibitors’ of MATE1 activity used to generate the pharmacophore. In other words, whereas a cationic feature was not sufficient to insure a strong interaction with MATE1, these data argue that charge does exert a strong, permissive influence on the binding interaction.

**1st Iteration: Quantitative Pharmacophore Development for hMATE1.** We generated a quantitative pharmacophore in parallel to the common features hMATE1 pharmacophore, taking advantage of the broad range of activities (IC$_{50}$ values from 40
nM to >>1 mM) displayed by the initial round of inhibitors. Twenty-four of the initial 26 compounds (H⁺, due to its restricted size, and verapamil, because it was a racemic mixture, were not included) were used in an analysis that resulted in a model containing two hydrophobic features (cyan), one hydrogen-bond donor (magenta), and, unlike the common features pharmacophore, one positive ionizable feature (red: Figure 8A). The model had a small cost difference as total cost = 125.97 and null cost = 137.57, suggesting a modest quality model. Nevertheless, unlike the modeling efforts based on single physical descriptors (Figure 5), the correlation between observed and predicted IC₅₀ values resulted in r = 0.68 (p < 0.0001; Figure 8B).

2nd Iteration: Quantitative Pharmacophore Development for hMATE1. Of the 39 compounds (the initial 24 and the “test set” of additional compounds derived from searching the database of FDA approved compounds) used to generate and validate the two pharmacophores, PYR was the most potent inhibitor of hMATE1 (and MATE2-K). Consequently, we chose to probe two structural analogs of PYR: (5-(4-chlorophenyl)-6-ethyl-2,4-pyrimidinediamine):1-(2-chlorophenyl)-6,6-dimethyl-1,6-dihydro-1,3,5-triazine-2,4-diamine (PYR-2); and 1-(3-chlorophenyl)-6,6-dimethyl-1,6-dihydro-1,3,5-triazine-2,4-diamine (PYR-3). The IC₅₀ values of 0.04, 0.14, and 0.20 µM for PYR, PYR-2 and PYR-3, respectively (Table 1), showed that the modest differences in structure between these three compounds had comparatively little impact on their inhibitory interactions with MATE1, and suggested that the structural features of this series of compounds may provide insight into molecular characteristics that optimize ligand interactions with the binding site/surface transport of the protein. A second quantitative pharmacophore model for hMATE1 reflecting these data (a total of 43
compounds) was generated (Figure 8C), and also included 2 hydrophobes (cyan), 2 hydrogen-bond acceptors (green), and an ionizable feature (red; Figure 8C). The correlation between observed and predicted IC₅₀ values resulted in an r value of 0.71 (p < 0.0001; Figure 8D).

**Final Iteration: Quantitative Pharmacophore Development for hMATE1.** We ultimately screened 59 compounds, adding several novel structural groups including the n-tetraalkylammonium series mentioned earlier. To minimize interpretational issues associated with compounds that were weakly ionized at the experimental pH of 8.5 we chose to eliminate for modeling analysis all compounds with pKᵦ values below 8.0 (a total of 13). The result was a final pharmacophore based on 46 structures and depicted in Figure 8E. This ‘N46’ model had several features in common with the previous iteration models, in that it included two hydrophobes (cyan), a hydrogen bond-acceptor (magenta), and an ionizable feature (red), though the spatial distribution of these elements differed somewhat from the previous models. Figure 8F displays the relationship between measured and predicted IC₅₀ values based on the N46 model (r value of 0.73, p < 0.0001).

**Bayesian Model.** A Bayesian model for MATE 1 (at pH 8.5) was generated using the N46 set of molecules; the receiver operator characteristic (ROC) was 0.88. Following leave out 50% x 100 this value is 0.82. (Concordance = 82.6 ± 4.7%, Specificity = 83.9 ± 5.5%, Selectivity = 66 ± 7.5%). These results suggest the model is stable.

Use of the molecular ‘function class fingerprints of maximum diameter 6’ (FCFP_6) descriptors allowed the identification of molecular features that favored
inhibition (Figure 9A), as well as features that did not promote inhibition (Figure 9B). Particularly noteworthy were the inclusion of nitrogen-containing 6-membered rings as ‘Bayesian good features,’ and the exclusion of nitrogen-containing 5-membered rings (pyrrole). The probable distinguishing characteristic between the two groups is the typically low basicity of the pyrrole moiety. In other words, pyrrole-containing structures generally are not cationic at physiological pH, again underscoring the importance of charge for increasing the effectiveness of ligand interaction with MATE transporters.
Discussion

The current study is the most complete to date on the selectivity of the human orthologs of MATE1 and MATE2-K, both of which are likely to play key roles in the renal secretion of organic cations. Previous data on selectivity of human MATEs are sufficiently sparse that it is difficult to compare our observations with those in the literature. Nevertheless, a few comparisons are noteworthy for their agreement with the observations reported here. The antimalarial drug, pyrimethamine (PYR), the highest affinity inhibitor in the present study (IC$_{50}$ of 42 nM; Table 1), was reported to have an IC$_{50}$ of 93 nM against hMATE1-mediated metformin transport (Kusuhara, et al., 2011), very similar to the IC$_{50}$ value we predicted for inhibition at pH 7.4 of MATE1-mediated MPP transport (IC$_{50}^{pH7.4}$ of 109 nM). Also, both hMATE1 and hMATE2-K are known to transport the antidiabetic drug, metformin, with K$_{tapp}$ values of 238 μM (Meyer zu Schwabedissen, et al., 2010) and 1.1 mM (Masuda, et al., 2006), respectively, not unlike the calculated IC$_{50}^{pH7.4}$ values we measured here (123 μM and 581 μM). But not all the comparisons of the present observations corresponded so closely to those observed in previous studies. For example, whereas tacrine had an IC$_{50}$ value of 0.6 μM for inhibition of hMATE1-mediated MPP transport, which agreed well with the 1.1 μM value recently reported (Kido, et al., 2011) for inhibition of hMATE1-mediated transport of 4-4-dimethlamminostyryl)-N-methyl-pyridinium (ASP), the IC$_{50}$ of 1.1 μM for tacrine inhibition of hMATE2-K-mediated MPP transport contrasted sharply with the value of >100 μM for inhibition of ASP transport by MATE2-K (Kido, et al., 2011). The experimental conditions used in the aforementioned study (i.e. HEK-293 cells at pH 7.4) differed from those employed here and could account for some of the difference.
However, we suggest it is more plausible that these differences are due to the use of MPP as the transported substrate, versus ASP (Kido, et al., 2011), and provide support for this suggestion later in this discussion.

The present observations support several conclusions concerning the molecular basis of selectivity of the mammalian MATEs. First, they support the hypothesis that no single physicochemical parameter of ligand structure is likely to provide an adequate predictor of interaction with MATE1 or MATE2-K. Previous studies of inhibition of OC/H⁺ exchange activity in isolated renal brush border membrane vesicles (Wright and Wunz, 1998; Wright and Wunz, 1999; Wright, et al., 1995) and intact microperfused renal proximal tubules (David, et al., 1995; Ullrich and Rumrich, 1996; Somogyi, et al., 1996; Ullrich, et al., 1991; Ullrich, et al., 1992) sought to correlate inhibitor effectiveness with selected, single physicochemical characteristics of the test agents included in these studies, such as LogP (hydrophobicity) and pKₐ (basicity). In fact, the IC₅₀ for inhibition of OC/H⁺ exchange activity in rabbit renal BBMV (Wright and Wunz, 1999), and intact rat RPT (David, et al., 1995), was shown to be strongly correlated with LogP, but the inhibitors used in those studies were more structurally constrained than those used in the present work and that may have masked the influence of steric (and other physicochemical) parameters on ligand interaction with the transporter(s). In support of this interpretation, the correlation between IC₅₀ and LogP for the n-tetraalkylammonium series was clearly evident (Figure 5B). We were not surprised that single molecular descriptors were not particularly effective predictors of inhibitory interaction with the MATEs. Given the broad structural diversity of compounds that interact effectively with the MATEs, it is probable that binding is a more complex process requiring multiple
molecular interactions and therefore single physicochemical properties alone are likely to be of limited predictive value.

The second set of conclusions arising from the present work stems from the application of computational methods to identify several physicochemical parameters that do influence ligand binding to MATE1, namely, the presence and location of multiple hydrophobic moieties; hydrogen donors; and an ionizable (i.e., cationic) feature. With respect to these two latter points, a recent study by Kido et al. (2011) that screened some 900+ compounds for inhibitory interaction with hOCT2 noted that inhibitory effectiveness was particularly influenced by (i) ligand lipophilicity and (ii) average charge, as well as (iii) molecular volume, (iv) TPSA, and (v) the number of hydrogen bond donors and acceptors. Third, in our study the several iterations of pharmacophore development led to the identification and subsequent verification of several novel clinical classes of compounds as MATE ligands. Table 2 identifies 12 drug classes not previously shown to interact with the human MATEs; seven of these displayed IC₅₀ values less than 25 µM, three of which (ketoconazole, proguanil, and imiquimod) represented a ratio of maximum plasma concentration (C_max) vs. IC₅₀ of <0.1.

The final pharmacophore (Figure 8 E/F) provides insight concerning the molecular basis of ligand interaction with the MATEs. Whereas the agreement between ‘predicted’ and ‘measured’ IC₅₀ values for 65% of the test compounds was within a factor of 5, for others the model displayed substantially less predictive capability. For example, the prototypic substrates of MATE1, MPP and TEA, were predicted to have IC₅₀ values some 10-15 times greater than their measured values of 5 and 50 µM,
respectively. These ‘misses’ may reflect an underlying assumption of pharmacophore analysis, namely, that there is a unique ‘most effective structure’ for interaction of ligand with a binding ‘site,’ with the pharmacophore representing both the location and physicochemical character of chemical features important for interaction with that singular site. However, the physiological role of MATE transporters requires that they interact effectively with a multitude of structurally diverse compounds, a characteristic that is, arguably, inconsistent with the existence of a single site for substrate/inhibitor interaction. Instead, we suggest our data are consistent with inhibitory ligand interactions at several structurally distinct sites that ‘overlap’ with the area(s) most favored for interaction with MPP. In this view, the pharmacophores represent a ‘statistical average’ of the influence of selected structural features of test ligands on inhibition of transport of a specific test probe. Thus, the inhibitory effectiveness of these ligands is likely to be influenced by the structural features of several distinct and potentially very different sites, a situation that would be difficult to describe with a single pharmacophore. A corollary to this suggestion is the hypothesis that the profiles of inhibition of structurally distinct substrates could result in distinct pharmacophores. Evidence in support of this idea was found in a preliminary analysis of data reported in the recent study by Kido et al. (2011) that included IC$_{50}$ values for inhibition of MATE1-mediated transport of the fluorescent OC, ASP. We used these IC$_{50}$ values (six compounds, spanning 2.5 orders of magnitude) to generate for hMATE1 a quantitative pharmacophore that proved to have a fundamentally different structure than those we generated based on inhibition of [³H]MPP transport (Figure 10). The quantitative pharmacophore based on inhibition of ASP transport had 3 hydrophobes (cyan), 2
hydrogen bond acceptors (green) and 3 excluded volumes (grey; Figure 10A) arranged in a spatial configuration that differed substantially from that of our model (Figure 10B). These data suggest that a comprehensive assessment of selectivity of MATEs (and other multidrug transporters) may require use of several structurally distinct substrates. This situation is analogous to what has been observed for the enzyme CYP3A4 (Kenworthy, et al., 1999).

In summary, we generated the first relatively large database for ligand inhibition of hMATE1 and hMATE2-K. Applying these data over the course of several computational modeling iterations using the IVIS approach resulted in a series of pharmacophores for hMATE1. The hMATE1 pharmacophores identified key structural features strongly correlated with ligand binding to hMATE1. The observations also supported the view that inhibitory profiles derived from the use of structurally distinct transported substrates can result in distinct pharmacophores, consistent with the contention that hMATE1 may have a complex binding surface for ligand interaction, rather than a single binding site.
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Authorship Contributions

Participated in research design: Astorga, Ekins, and Wright

Conducted experiments: Astorga and Morales

Contributed new reagents or analytic tools: N/A

Performed data analysis: Astorga, Morales, Ekins, and Wright

Wrote or contributed to the writing of the manuscript: Astorga, Ekins, and Wright
References


Footnotes

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Conflict of interest statement

SE consults for Collaborative Drug Discovery, Inc.
Legends for Figures

Figure 1. Kinetic characteristics of transport mediated by hMATE1 and hMATE2-K expressed in CHO cells. Kinetics of MPP transport mediated by hMATE1 (A) or hMATE2-K (B). Effect of extracellular [H+] on MPP transport mediated by hMATE1 (C) or hMATE2-K (D). In all experiments, 5 min uptakes of [3H]MPP (~13 nM) were measured in the presence of increasing concentrations of unlabeled MPP (A, B) at an external pH of 8.5; or in the presence of increasing extracellular H+ concentration (C,D). Each point is the mean (±SEM) of uptakes measured in three wells of a 24-well plate from single, representative experiments. Kinetic values shown in the figures represent the average of 3-5 experiments.

Figure 2. Range of inhibition of transport mediated by the human orthologs of MATE1 and hMATE2-K produced by the battery of test compounds used in this study. The height of the gray bars indicates the degree of inhibition of mediated uptake (5 min) of [3H]MPP (~13 nM) produced by 10 µM concentrations of 59 test compounds. The horizontal dashed line indicates 50% inhibition of transport.

Figure 3. The effect on the rate of hMATE1- and hMATE2-K-mediated MPP transport of increasing concentration of representative test inhibitors. Solid circles represent hMATE1, open circles hMATE2-K, and depict inhibition produced by: (A) quinidine, (B) agmatine, (C) nialamide, and (D) allopurinol. Each point represents the mean value (±SEM; determined in three separate experiments) of the 5 min uptake of [3H]MPP (~13nM) measured in the presence of increasing concentration of test inhibitor; uptakes were normalized to that measured in the absence of inhibitor. NI = No interaction.
Figure 4. Comparison of hMATE1 & hMATE2-K IC\textsubscript{50} Values. (A) hMATE1 IC\textsubscript{50} values were graphed as a function of hMATE2-K IC\textsubscript{50} values for the 59 test compounds measured at pH 8.5 (left) and as calculated for pH 7.4 (right). The solid line represents equal IC\textsubscript{50} values for the two transporters; the dashed lines indicate 3-fold +/- differences in these values. (B). Inhibitory profiles for several test ligands (atropine, amantadine and azidoprocainamide (APMI)) against MPP transport mediated by hMATE1 (solid circles) and hMATE2-K (open circles). Each point is mean of 5 min uptakes (±SEM; normalized to uptake measured in the absence of inhibitor) determined in three separate experiments each run in duplicate. IC\textsubscript{50} values indicated in the figures are average values from three separate experiments.

Figure 5. Relationship between hMATE1 IC\textsubscript{50} Values and the molecular descriptors (A) LogP, (C) TPSA and (D) pK\textsubscript{a}. Panel (B) shows the relationship between LogP and the IC\textsubscript{50} values for a structurally constrained n-tetraalkylammonium series (TMA, tetramethylammonium; TEA, tetraethylammonium; TPrA, tetrapropylammonium; TBA, tetrabutylammonium; TPeA, tetrapentylammonium).

Figure 6. (A) Inhibitory profiles for the five test ligands used to generate (B) a Common Features Pharmacophore. Each point is the mean of 5 min uptake (normalized to the transport measured in the absence of inhibitor) determined in single representative experiments. PYR (open triangles), quinidine (solid triangles), histamine (solid diamonds), caffeine (solid squares) and chloramphenicol (solid circles). The Common Features Pharmacophore (displayed with the structure of PYR) includes 2 hydrophobic regions (cyan), 1 H-bond donor (magenta), and 1 H-bond acceptor (green).
Figure 7. Kinetics of inhibition of hMATE1-mediated MPP transport produced by (A) cinchonidine and (B) ethohexadiol. Each point is mean (±SEM) of 5 min uptake measured in triplicate in single representative experiments. IC50 values indicated in the figures are average values from three separate experiments. IC50 values were calculated as an average of three experiments. Below each graph the Common Features Pharmacophore is shown with the respective test molecule.

Figure 8. (A,B) Quantitative pharmacophore generated from analysis of data obtained using the first round of hMATE1 inhibitors (see text). (A) Twenty four of the initial twenty six compounds were used in an analysis that resulted in a model containing two hydrophobic features (cyan), one hydrogen-bond donor (magenta), and one positive ionizable feature (red), shown here with the structure of cinchonidine. (B) The relationship between measured and predicted IC50 values based on the model shown in (A) (r = 0.68, p < 0.0001). (C,D) Quantitative pharmacophore generated from analysis of the data that incorporated the second round of hMATE1 inhibitors. (C) Analysis of 43 compounds (including the initial 24 plus the “test set” of 15 compounds that probed the common features model) resulted in a model that included 2 hydrophobes (cyan), 2 hydrogen-bond acceptors (green), and an ionizable feature (red). (D) The relationship between measured and predicted IC50 values based on the model shown in (C) (r = 0.71, p < 0.0001). (E,F) Quantitative pharmacophore generated from analysis of 46 of the 59 test ligands (see text for inclusion criteria). (E) The model included two hydrophobes (cyan), a hydrogen bondacceptor (magenta), and an ionizable feature (red). (F) The relationship...
between measured and predicted IC$_{50}$ values based on the N46 model ($r = 0.73$, $p < 0.0001$). Quinidine is mapped to all pharmacophores.

Figure 9. (A) FCFP_6 features associated with hMATE1 inhibitors – pH 8.5 N46 model. (B) FCFP_6 features associated with hMATE1 non-inhibitors – pH 8.5 N46 model. Each of the 20 panels shows the naming convention for one fragment, the numbers of compounds containing the fragment, and the Bayesian score for the fragment.

Figure 10. hMATE1 quantitative pharmacophores based on inhibition of MATE1-mediated transport of (A) ASP or (B) MPP. The ASP-derived pharmacophore (with ondansetron) had an $r = 0.95$ and was generated using a data set obtained from (Kido, et al., 2011) that consisted of six compounds with IC$_{50}$ values ranging from 0.15 µM to 66 µM. The MPP-derived N46 pharmacophore (with quinidine) was developed as described in the text for the other pharmacophores and in Figure 8E. Pharmacophore features as described in Figure 8 (with the addition of grey features indicating excluded volumes).
Table 1. IC₅₀ values for inhibition of MPP transport into CHO cells stably expressed with either hMATE1- or hMATE2-K.

<table>
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<tr>
<th>Compound</th>
<th>Drug Class</th>
<th>MATE1 IC₅₀ (µM) pH 8.5</th>
<th>MATE1 Cal. IC₅₀ pH 7.4</th>
<th>MATE2-K IC₅₀ (µM) pH 8.5</th>
<th>MATE2-K Cal. IC₅₀ pH 7.4</th>
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<td>Agmatine</td>
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<td>53.9 ± 1.4</td>
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<td>Atropine</td>
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<td>Caffeine</td>
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<td>451 ± 120</td>
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<td>Chloramphenicol</td>
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<td>2900</td>
<td>1951 ± 45</td>
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<td>0.939 ± 0.043</td>
<td>5.8</td>
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<td>Ni</td>
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<td>Midobutine</td>
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<tr>
<td>Phenotamine</td>
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<td>6.34 ± 0.23</td>
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<td>OA</td>
<td>17.8 ± 1.6</td>
<td>46</td>
<td>74.6 ± 17.6</td>
<td>490</td>
</tr>
<tr>
<td>Trichloromethiazide</td>
<td>DI</td>
<td>249 ± 10</td>
<td>640</td>
<td>679 ± 10</td>
<td>4200</td>
</tr>
<tr>
<td>trimethazin</td>
<td>AB</td>
<td>9.70 ± 1.2</td>
<td>225</td>
<td>281 ± 12.7</td>
<td>17</td>
</tr>
<tr>
<td>Triptophan</td>
<td>AAM</td>
<td>Ni</td>
<td>N/A</td>
<td>Ni</td>
<td>N/A</td>
</tr>
<tr>
<td>Tyramine</td>
<td>MA</td>
<td>86.6 ± 8.2</td>
<td>230</td>
<td>138 ± 11</td>
<td>900</td>
</tr>
<tr>
<td>Valproate</td>
<td>LTCB</td>
<td>41.9 ± 6.9</td>
<td>110</td>
<td>37.9 ± 9.7</td>
<td>370</td>
</tr>
</tbody>
</table>

Each value determined at pH 8.5 is a mean IC₅₀ (µM ± SEM) of 2-4 experiments. IC₅₀ values listed for pH 7.4 were calculated (see text). Underlined compounds indicate the drugs that comprised the initial set of 24 organic molecules used in the first iteration of pharmacophore development; * indicates the 15 compounds used to test the common features pharmacophore; italicized compounds are those identified by the pharmacophores from the list of FDA-approved drugs during the course of model development. Abbreviations: AA, not available; A2A, Alpha-2A Adrenergic Agonist; AS2A, Alpha-2A Adrenergic Receptor Agonist; AA, Antiarrhythmic; AAA, Alpha Adrenergic Agonist; AAC, Amino Acid; AB, Antibiotic; AC, Anticholinergic; ACS, Anticholinesterase; ACT, Anticonvulsant; AD, Antiadipetic; AF, Antifungal; AM, Antimalarial; AMS, Antimuscarinic; AP, Antipaltetate; AS, Antispasmodic; AV, Antiviral; BB, Beta Blocker; CT, Chemotherapy Drug; DI, Diuretic; EI, Endogenous Ion; H, Herbicide; H2RA, Histamine H2-Receptor Agonist; IMR, Immune Response Modifier; IR, Insect Repellent; IRR, Immune Response Regulator; LTCB, L-Type Calcium Blocker; MA, Monoamine; MAOI, Monoamine Oxidase Inhibitor; MBP, Metabolism By-Product; MS, Muscle Relaxant; NT, Neurotransmitter; NTX, Neurotoxin; OA, Opate Agonist; OAN, Opate Antagonist; PAL, Photoaffinity Label; POC, Prototypical Organic Cation; S, Stimulant; Sc, Synthetic Chemical; VP, Vasopressor; XO1, Xanthine Oxidase Inhibitor; and NI, No interaction.
Table 2. MATE1 inhibitors from novel drug classes identified during the course of pharmacophore development.

<table>
<thead>
<tr>
<th>Inhibitor</th>
<th>Class</th>
<th>IC\textsubscript{50} (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ketoconazole</td>
<td>AF</td>
<td>1.3</td>
</tr>
<tr>
<td>Proguanil</td>
<td>PAM</td>
<td>4.4</td>
</tr>
<tr>
<td>Phentolamine</td>
<td>AAA</td>
<td>4.6</td>
</tr>
<tr>
<td>Propranolol</td>
<td>BB</td>
<td>7.8</td>
</tr>
<tr>
<td>Imiquimod</td>
<td>IMR</td>
<td>13.9</td>
</tr>
<tr>
<td>Tramadol</td>
<td>OA</td>
<td>17.8</td>
</tr>
<tr>
<td>Naloxone</td>
<td>OAN</td>
<td>24.1</td>
</tr>
<tr>
<td>Midodrine</td>
<td>VP</td>
<td>109</td>
</tr>
<tr>
<td>Nialamide</td>
<td>MAOI</td>
<td>212</td>
</tr>
<tr>
<td>Topiramate</td>
<td>ACT</td>
<td>1000</td>
</tr>
<tr>
<td>Ticlopidine</td>
<td>AP</td>
<td>1000</td>
</tr>
<tr>
<td>Ethohexadiol</td>
<td>IR</td>
<td>2000</td>
</tr>
</tbody>
</table>

Abbreviations: AF, antifungal; PAM, prophylactic antimalarial; AAA, alpha adrenergic antagonist; BB, beta blocker; IMR, immune response modifier; OA, opiate agonist; OAN, opiate antagonist; VP, vasopressor; MAOI, monoamine oxidase inhibitor; ACT, anticonvulsant; AP, antiplatelet; IR, insect repellant.
Figure 1

(A) [H]MPP Uptake (fmol cm⁻² min⁻¹) vs. [MPP] μM for hMATE1 with $J_{max} = 2.14$ pmol cm⁻² min⁻¹ and $K_{tapp} = 4.37$ μM.

(B) [H]MPP Uptake (fmol cm⁻² min⁻¹) vs. [MPP] μM for hMATE2-K with $J_{max} = 1.86$ pmol cm⁻² min⁻¹ and $K_{tapp} = 3.72$ μM.

(C) [H]MPP Uptake (fmol x cm⁻² min⁻¹) vs. [H⁺] nM for hMATE1 at pH 7.7 with $IC_{50} = 19.6$ nM.

(D) [H]MPP Uptake (fmol x cm⁻² min⁻¹) vs. [H⁺] nM for hMATE2-K at pH 8.5 with $IC_{50} = 3.5$ nM.
Figure 4
Figure 5
Pyrimethamine IC$_{50}$ = 0.04 μM
Quinidine IC$_{50}$ = 1.57 μM
Histamine IC$_{50}$ = 761 μM
Chloramphenicol IC$_{50}$ = 1115 μM
Caffeine IC$_{50}$ = 1096 μM

Figure 6
Figure 7
Figure 8
Figure 9