Intracellular drug bioavailability: a new predictor of system dependent drug disposition

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Intracellular drug exposure is influenced by cell- and tissue-dependent expression of drug-transporting proteins and metabolizing enzymes. Here, we introduce the concept of intracellular bioavailability (Fic) as the fraction of extracellular drug available to bind intracellular targets, and we assess how Fic is affected by cellular drug disposition processes. We first investigated the impact of two essential drug transporters separately, one influx transporter (OATP1B1; SLCO1B1) and one efflux transporter (P-gp; ABCB1), in cells overexpressing these proteins. We showed that OATP1B1 increased Fic of its substrates, while P-gp decreased Fic. We then investigated the impact of the concerted action of multiple transporters and metabolizing enzymes in freshly-isolated human hepatocytes in culture configurations with different levels of expression and activity of these proteins. We observed that Fic was up to 35-fold lower in the configuration with high expression of drug-eliminating transporters and enzymes. We conclude that Fic provides a measurement of the net impact of all cellular drug disposition processes on intracellular bioavailable drug levels. Importantly, no prior knowledge of the involved drug distribution pathways is required, allowing for high-throughput determination of drug access to intracellular targets in highly defined cell systems (e.g., single-transporter transfectants) or in complex ones (including primary human cells).

Drug-transporting proteins and metabolizing enzymes are determinants of intracellular drug disposition1-2: uptake transporters increase the drug concentration in the cell interior3-5, while efflux transporters6-8 and enzymes9,10 reduce the amount of intracellularly available compound (Fig. 1). Therefore, these proteins are of major importance for drug efficacy and toxicity1. The contribution of such drug-disposition proteins has been studied in vitro, e.g., in cell lines expressing specific transporters5,12-15, in hepatocytes (in suspension, monolayer and sandwich cultures)16,17, and in more complex systems, such as tissue slices18 and 3D cultures19. Typically, only the relative impact of transporters on the intracellular compound concentration is reported, i.e., fold-differences in intracellular concentration when the transporter or metabolic enzymes are active (or overexpressed) compared to when these proteins are inhibited (or absent). Quantitative determination of the intracellular unbound drug concentration, which is the relevant concentration for binding to intracellular target proteins, either relies on experimentally demanding setups in freshly isolated cells3-5,20,21 or tissues22,23, or on a priori knowledge of transport and metabolic mechanisms for the compound being studied24. To overcome these limitations, we recently developed a simple methodology to measure intracellular unbound drug concentrations in cultured cells25, compatible with high-throughput formats26,27. We also showed that our methodology correlates with binding to an intracellular target (thymidylate synthase)28.

In this work, we apply this methodology to study the impact of drug-transporting proteins and metabolizing enzymes on intracellular unbound compound concentrations. For this purpose, we introduced the concept of intracellular compound bioavailability (Fic), which is the fraction of the externally added compound concentration that is available to bind targets in the cell interior. Throughout this study, this concept is equivalent to the
unbound drug accumulation ratio (Kp,un), a term commonly used in pharmacokinetic studies of blood-to-tissue concentration ratios. However, there are situations in which the Kp,un term is not applicable (e.g., when experiments are performed in the presence of serum proteins). First, we evaluated how Fic is affected by a single uptake or a single efflux transporter. We used well-defined cell lines overexpressing the organic anion-transporting polypeptide 1B1 (OATP1B1; SLCO1B1*1a) and P-glycoprotein (P-gp; ABCB1), two of the transporters most commonly involved in drug disposition. We then studied the parallel impact of multiple transporters and enzymes in a more complex system, freshly isolated human hepatocytes. The hepatocytes were used in two configurations known to display different gene expression and activity of transporters and enzymes: directly after isolation and after 24 h of culture in monolayer format. In conclusion, our simple methodology provides a good estimate of compound available to bind targets in the cell interior, even in complex cell systems.

Results

Impact of OATP1B1 on intracellular bioavailability (Fic) of compounds. We investigated the impact of drug-transporting proteins on intracellular compound bioavailability (Fic). First, we studied the influence of an important uptake transporter (Fig. 1) by using HEK293 cells transfected with human OATP1B1 or with an empty vector (mock-transfected cells; Fig. 2a). We measured the intracellular fraction of unbound compound (fucell) in cell homogenates and cellular compound accumulation (Kp) in live cells (see ‘Methods’ for details). From these parallel experiments, Fic was then calculated as the product of fucell and Kp, providing a measure of how much of an extracellularly applied drug concentration is freely available in the cell interior.

Impact of P-gp on intracellular bioavailability (Fic) of compounds. We next evaluated how efflux transporters affect Fic (Fig. 1) through measurements in MDCK cells transfected with human P-gp. To determine baseline Fic in these cells in the absence of P-gp, we used wild-type MDCK cells where background endogenous
canine P-gp was knocked out using CRISPR-Cas9 technology (cP-gp-KO)\textsuperscript{44} (Fig. 3a)—see ‘Supplementary results’ for a discussion on the choice of these cells as a control.

As in the experiments with HEK293 cells, we first measured $f_{u,\text{cell}}$ in MDCK cells transfected with human P-gp (Supplementary Table 3). We used this value of $f_{u,\text{cell}}$ for all MDCK cell types used in this study, based on our observation above that the low levels at which transporters are expressed (compared to total protein) do not affect binding (Fig. 2b; P-gp contributes only 0.1% of all protein in P-gp-transfected MDCK cells (6.0 pmol/mg total protein, in-house data)).

We then combined $f_{u,\text{cell}}$ and $K_p$ measurements to calculate $F_{ic}$ in P-gp-transfected MDCK cells and in cP-gp-KO cells (Supplementary Table 3). Because of the active efflux of substrates when the transporter is present, the $F_{ic}$ of compounds previously reported to be P-gp substrates was on average 2.0-fold lower in P-gp-expressing than in cP-gp-KO cells at 0.1 μM compound concentration (range: 0.94–20-fold lower in P-gp transfected than cP-gp-KO cells; $p < 0.001$ in Wilcoxon matched-pairs signed rank test; Fig. 3b). For compounds not previously described as substrates, $F_{ic}$ was more similar in the two cell lines (range: 1.1–3.0-fold difference between cP-gp-KO cells and P-gp transfected cells), but significantly different ($p = 0.001$; see discussion below). Nelfinavir, an established P-gp substrate, displayed a remarkably low $F_{ic}$ in P-gp-transfected cells (20-fold lower than in cells not expressing P-gp). This indicates that P-gp contributes strongly to the cellular elimination of nelfinavir, in particular in cells that do not metabolize this compound. Overall, lipophilic molecules showed lower $F_{ic}$ in P-gp transfected cells ($r_S = −0.36$, $p = 0.05$ for the rank-order correlation between difference in $F_{ic}$ and log D of all compounds; Supplementary Figure 2b), independently of being previously described as substrates or not. This possibly reflects that compounds bind P-gp from within the inner leaflet of the plasma membrane\textsuperscript{45,46}, and suggests that additional compounds in our dataset may be substrates of P-gp, besides those previously reported as such.

As for the OATP1B1 substrates, we measured the concentration-dependence of $F_{ic}$ in P-gp-expressing and cP-gp-KO cells (Supplementary Table 3). Because of the active efflux of substrates when the transporter is present, the $F_{ic}$ of compounds previously reported to be P-gp substrates was on average 2.0-fold lower in P-gp-transfected than cP-gp-KO cells (range: 0.94–20-fold lower in P-gp transfected than cP-gp-KO cells; $p < 0.001$ in Wilcoxon matched-pairs signed rank test; Fig. 3b). For compounds not previously described as substrates, $F_{ic}$ was more similar in the two cell lines (range: 1.1–3.0-fold difference between cP-gp-KO cells and P-gp transfected cells), but significantly different ($p = 0.001$; see discussion below). Nelfinavir, an established P-gp substrate, displayed a remarkably low $F_{ic}$ in P-gp-transfected cells (20-fold lower than in cells not expressing P-gp). This indicates that P-gp contributes strongly to the cellular elimination of nelfinavir, in particular in cells that do not metabolize this compound. Overall, lipophilic molecules showed lower $F_{ic}$ in P-gp transfected cells ($r_S = −0.36$, $p = 0.05$ for the rank-order correlation between difference in $F_{ic}$ and log D of all compounds; Supplementary Figure 2b), independently of being previously described as substrates or not. This possibly reflects that compounds bind P-gp from within the inner leaflet of the plasma membrane\textsuperscript{45,46}, and suggests that additional compounds in our dataset may be substrates of P-gp, besides those previously reported as such.

As for the OATP1B1 substrates, we measured the concentration-dependence of $F_{ic}$ in P-gp-expressing and cP-gp-KO cells. For simvastatin acid, $F_{ic}$ in P-gp-expressing cells was lower than that in cP-gp-KO cells at low concentrations and approached it at higher concentrations (Fig. 3c). Surprisingly, $F_{ic}$ decreased with increasing compound concentration in both the P-gp expressing cells and in the cP-gp-KO cells, which was not caused by a saturation of binding sites\textsuperscript{47} (Supplementary Figure 4). Similar results were observed for loperamide (Supplementary Figure 3). When inhibiting active transport with 10 μM cyclosporine A (a general transporter inhibitor\textsuperscript{47}), we observed that $F_{ic}$ of simvastatin acid was constant across all concentrations and lower than
previously observed without inhibition (Fig. 3c). This suggests that additional active transport mechanisms (including uptake transporters) are present in these cells and affect the cellular disposition of this compound.

To better understand these observations, we simulated cellular drug accumulation using three simple kinetic models: 1) one including both active uptake and active efflux mechanisms (corresponding to P-gp transfected cells), 2) one including only the uptake mechanisms (corresponding to cP-gp-KO cells), and 3) one including only the passive mechanism (corresponding to inhibition of all active mechanisms with cyclosporine A). We simulated all possible combinations of low, medium and high values of passive permeability (0.1, 1, and $10 \times 10^{-6}$ cm/s), $V_{\text{max,uptake}}$ (10, 100, and 1000 pmol/min/mg protein) and $K_{m,\text{uptake}}$ (1, 10, and 100 μM) for each of the transporters (see Matsson et al. and the 'Methods' section for details on the simulations). The simulations were in best agreement with the experimental observations for a compound with high passive permeability ($10 \times 10^{-6}$ cm/s), medium affinity ($K_{m,\text{uptake}} = 10$ μM) for an uptake transporter and lower affinity ($K_{m,\text{efflux}} = 100$ μM) for an efflux transporter, with

Figure 3. Impact of the efflux transporter P-gp on intracellular bioavailability ($F_{lc}$). (a) Schematic representation of cell types used in this study: MDCK cells where canine P-gp has been knocked-out using CRISPR/Cas 9 (cP-gp-KO); MDCK cells transfected with human P-gp, expressing both canine and human P-gp. (b) Comparison between $F_{lc}$ in cP-gp-KO and in P-gp-transfected MDCK cells at 0.5 μM compound concentration. Negatively charged compounds at pH 7.4 are represented as triangles, neutral and zwitterionic species are represented by circles, and positively charged compounds are represented by squares. Substrates of P-gp are highlighted in yellow. Nelfinavir is represented in parentheses as its $F_{lc}$ is outside the range of this plot in P-gp-transfected cells. (c) Relationship between simvastatin acid concentration and $F_{lc}$ in P-gp-transfected MDCK cells (yellow filled squares), P-gp-KO MDCK cells (blue circles), and P-gp-KO MDCK cells pre-incubated (15 min) followed by co-incubation with 10 μM cyclosporine A (gray triangles). (d) Kinetic cell model simulations that best described experimental observations of concentration-dependence $F_{lc}$ for simvastatin acid (conditions: passive permeability = $10 \times 10^{-6}$ cm/s; $V_{\text{max,uptake}} = 1000$ pmol/min/mg protein; $K_{m,\text{uptake}} = 10$ μM; $V_{\text{max,efflux}} = 1000$ pmol/min/mg protein; $K_{m,\text{efflux}} = 100$ μM).
both transporters having comparable (high) transport capacities \((V_{\text{max}})\) (Fig. 3d). This matches previous observations for simvastatin acid, which show that this compound, in addition to being an OATP1B1 substrate, has a high passive permeability, and a low affinity for efflux transporters\(^{49-51}\).

**Intracellular bioavailability \((F_{\text{ic}})\) in complex cell systems.** Our results in MDCK cells revealed that unknown endogenous transport mechanisms can influence \(F_{\text{ic}}\) to a larger extent than previously recognized. To further investigate how multiple transporters impact \(F_{\text{ic}}\), we performed measurements in freshly isolated human hepatocytes. These cells also express metabolizing enzymes, which can further affect \(F_{\text{ic}}\) (Fig. 1), and better represent an *in vivo* situation where multiple processes occur simultaneously. Hepatocytes were used in two commonly employed culture conditions (Fig. 4a): 1) freshly isolated cells (in suspension) that express similar levels of transporters and metabolizing enzymes as those observed in the human liver\(^{29-31,52}\); and 2) cells cultured for 24 h post-isolation (in monolayer format) with a significant down-regulation of many important transporters and enzymes due to cell dedifferentiation\(^{29-31}\). Thus, without knowing the exact composition of the contributing transporters and enzymes, their relative impact in two complex hepatocyte systems could be compared. For these studies, we selected a subset of 16 compounds from the initial dataset (Supplementary Table 4). The compounds were chosen to be structurally diverse and to be substrates of different transporters and enzymes with altered expression (Supplementary Table 5).

On average, \(F_{\text{ic}}\) was 5.7-fold lower in freshly isolated suspension hepatocytes than in monolayer hepatocytes cultivated for 24 h (Fig. 4b; \(p = 0.0002\) in Wilcoxon matched-pairs signed rank test). No apparent correlation was observed between the physicochemical characteristics of the compounds and the differences in \(F_{\text{ic}}\) in the two types of hepatocytes (Supplementary Figure 5). However, compounds that are reported as substrates of multiple efflux transporters and enzymes displayed larger differences between the two culturing conditions \(\left(r_S = 0.72\right)\) for the rank-order correlation between differences in \(F_{\text{ic}}\) and the number of targeted enzymes and transporters). This was in agreement with the reported higher mRNA transcript levels in suspension hepatocytes for these proteins\(^{29-31}\), which are associated with the removal of compounds from the cell interior (leading to a lower \(F_{\text{ic}}\)). In agreement with these results, compounds that were metabolically cleared at higher rates (e.g., atorvastatin, propranolol, and ritonavir; Fig. 4c; \(p < 0.05\) in Mann-Whitney test) showed lower \(F_{\text{ic}}\) in suspension hepatocytes. In analogy, compounds with a similar metabolic clearance under both culture conditions (e.g., astemizole, indomethacin, and ketoconazole) also had similar \(F_{\text{ic}}\) (Fig. 4c). These results strongly support our hypothesis that \(F_{\text{ic}}\) is a general predictor of intracellular drug disposition. We also measured the protein concentrations of the major transporters and metabolic enzymes involved in the clearance of the studied compounds using state-of-the-art methods.
mass spectrometry-based proteomics (Supplementary Figure 6). To our surprise, the agreement between mRNA transcript levels28,30, enzymatic activity (Fig. 4c) and $F_c$ (Fig. 4b) did not translate to the protein level, since we did not observe significant differences in protein concentrations between the two culturing conditions ($p = 0.6$ in Wilcoxon matched-pairs signed rank test). This can be explained by drug-transporting proteins being redistributed from the plasma membrane to intracellular compartments under certain culturing conditions, or as a result of external stimuli53–55. Similarly, the activity of metabolic enzymes can be reduced by post-translational modifications56 or lower levels of co-factors57. In these cases, these proteins will contribute to the whole-cell protein levels but will not affect $F_c$. This underscores the importance of functional studies to assess the impact of drug-transporting proteins and metabolizing enzymes on intracellular drug concentrations.

Discussion

Intracellular compound exposure is of paramount importance for the pharmacodynamics of drugs acting on intracellular targets, and for interactions with intracellular drug-metabolizing enzymes11,28. In this study, we introduce the concept of intracellular compound bioavailability ($F_c$) and demonstrate its utility in describing the impact of drug-transporting proteins and metabolizing enzymes on cellular drug disposition.

Our methodology is based on the parallel determination of the fraction of intracellular compound that is not bound to cellular components ($f_{\text{cell}}$) and cellular compound accumulation ($K_p$). Our results indicate that over-expression of single proteins (e.g. OATP1B1) has little impact on $f_{\text{cell}}$ (Fig. 2b). Further, good correlations were observed for $f_{\text{cell}}$ across all cell types used in this study, despite different proteomes in these cells ($r_p > 0.8$ for all comparisons of between HEK293 cells, MDCK cells and freshly isolated hepatocytes). This supports previous observations that $f_{\text{cell}}$ is mostly dominated by membrane partitioning52,53,33,34, and that specific proteins contribute negligibly to this parameter.

In contrast, clear effects were observed for $F_c$ when HEK293 cells were transfected with the cellular uptake-mediating transporter OATP1B1, with intracellular exposure increasing for all investigated substrates of OATP1B1. Interestingly, some statins, such as atorvastatin and pitavastatin, showed $F_c < 0.1$ when the transporter was lacking, suggesting that they have limited cell penetration in the absence of transporters.

Our studies on the concentration-dependent uptake of the OATP1B1 substrates showed a greater impact of the transporter at lower substrate concentrations. At the lowest measured concentrations (approaching unbound clinical plasma concentrations), $F_c$ was up to 10-fold higher for atorvastatin and pitavastatin, and up to 2-fold higher for fluvastatin and simvastatin acid in the presence of the transporter. These results are in agreement with clinical drug-drug interaction studies12,13,37,38, in which the pharmacokinetics of atorvastatin and pitavastatin are more affected than fluvastatin and simvastatin acid when co-administered with OATP1B1 inhibitors. For example, cyclosporine A increases the AUC of pitavastatin and atorvastatin by 5- to 15- fold, and that of simvastatin and fluvastatin by 3- to 4-fold in human subjects59. Furthermore, pharmacogenomic studies have also shown that reduced-function haplotypes of SLCO1B1 contribute to higher plasma concentrations of these compounds60–62. With the exception of simvastatin acid, which (based on only one study62) has a larger clinical effect than the other statins from a reduced function polymorphism of OATP1B1, our results accurately reproduce the rank order of the clinically observed AUC change (pitavastatin > atorvastatin > fluvastatin).

Contrary to HEK293 cells, MDCK cells express significant levels of endogenous drug-transporting proteins, including canine P-gp in the present study, we used our recently established MDCK cell line where this transporter was completely knocked out using CRISPR-Cas9 (cP-gp-KO cells)44. This enabled the study of baseline transport into MDCK cells without the need for chemical inhibition of P-gp. This was considered important, since we have shown that P-gp inhibitors such as elacridar also inhibit other endogenous transporters35,47. Since efflux transporters are driven by intracellular (or intramembraneous) substrate concentrations, it is desirable to avoid simultaneous inhibition of the uptake transporters that some compounds depend on to reach the cell interior. Thus, studies that rely on chemical inhibition can misclassify compounds as non-substrates. In fact, we observed that some compounds not previously annotated as P-gp substrates (e.g. astemizole, ketoconazole, lovastatin and rosiglitazone) showed large differences in $F_c$ between P-gp-expressing and cP-gp-KO cell lines. These compounds have previously been reported as P-gp inhibitors64–66. Our findings suggest that they are also transported substrates. In follow-up experiments, we have confirmed that at least two of these compounds show asymmetrical transport across a monolayer of MDCK cells (manuscript in preparation). This indicates that the setup used here can replace more demanding transport assays to identify substrates of proteins involved in drug disposition and resistance to chemotherapy67.

The possibility of studying the involvement of multiple transport processes with this methodology, as observed with simvastatin acid in MDCK cells (Fig. 3c), prompted us to study $F_c$ in the more complex system of freshly isolated human hepatocytes. Primary human hepatocyte models are extensively used in drug discovery and are available in many different configurations, which display marked differences in gene expression and activity of drug-transporting proteins and drug-metabolizing enzymes29–31. Our measurements of $F_c$ reflected these differences in that lower $F_c$ values were observed in suspension hepatocytes, where the elimination pathways via efflux transporters and drug-metabolizing enzymes are more active29–31. Interestingly, only compounds reported to be substrates of multiple drug-transporting proteins and metabolizing enzymes were affected (Supplementary Table 5), while compounds reported to be substrates of mainly one or two metabolizing enzymes (e.g. astemizole, indinavir, ketoconazole, lopinavir, and rosiglitazone) displayed similar $F_c$ in both culturing conditions. For example, ritonavir, a compound with an $F_c$ of 1.0 in mock-transfected HEK293 cells (where passive mechanisms are the major determinants of drug accumulation23), showed an $F_c$ of 0.26 in freshly isolated suspension hepatocytes (indicative of a dominance of elimination mechanisms) and an $F_c$ of 2.3 in monolayer hepatocytes (indicative of reduced elimination and a dominance of active uptake mechanisms). This was confirmed by functional in vitro clearance studies, which showed that the metabolic CLint of ritonavir was in fact
Previous approaches to determine intracellular unbound drug concentrations based on mathematical modeling require extensive collection of experimental data (at multiple time points and concentrations)\(^3\), which limits their compound throughput. Alternative approaches rely on a separate measurement of drug binding to the cells that is combined measurements of the total intracellular drug concentrations. In such approaches, binding can be estimated by inhibiting all active processes (with chemical inhibitors\(^7\) or reduced temperature\(^21\)) or using equilibrium dialysis of tissue or cell homogenates\(^22,25\). The use of chemical inhibitors assumes that all transport and metabolism mechanisms are known and can be inhibited, while lowering the temperature assumes that changes in membrane fluidity\(^69\) do not affect drug binding. Our approach uses equilibrium dialysis of cell homogenates and assumes that binding is not affected by the disruption of the cellular context during homogenization. This allows the processing of large numbers of compounds without prior knowledge of compound elimination kinetics or metabolic pathways. Despite the utility of a simple estimate of intracellular available compound, the \(F_{ic}\) parameter does not provide information on the subcellular distribution of the compound. Instead, it provides an average bioavailability in the whole cell. As previously shown, mathematical modeling of pH\(^22,25\) or electrochemical gradients\(^5\) can offer insight into the accumulation of compounds in lysosomes or mitochondria. In addition, inhibition of ion channels responsible for the maintenance of these gradients can be used to validate such models\(^25\). Alternatively, subcellular fractionation can be performed prior to\(^5\) or after\(^26\) incubation with the drug. However, fractionation approaches might alter compound distribution, since the organelle is no longer in its native environment. All of these approaches are compatible with the overall methodology presented in this study and can be used in specific cases, when the target of interest is located in a particular subcellular compartment.

In conclusion, we introduce \(F_{ic}\) as a new approach to study the impact of proteins involved in drug disposition on the intracellular drug concentration available for target and off-target (e.g. drug-metabolizing enzymes) interactions. We show how single and multiple drug-transporting proteins and metabolizing enzymes influence \(F_{ic}\) in various cellular systems of different complexity. We also show that our setup enables identification of potential substrates of transporters and could constitute an alternative to state-of-the-art in vitro permeability assays. Thus, \(F_{ic}\) provides a true value of intracellular accumulation that is unbiased from membrane interactions. \(F_{ic}\) should therefore not only be applicable in predictive pharmacokinetics, but in particular during compound profiling in drug discovery aiming at intracellular targets\(^{11,27,28,71,72}\).

### Methods

#### Theoretical background: intracellular compound bioavailability (\(F_{ic}\)).

To determine intracellular unbound compound concentrations, we measured intracellular fraction of unbound compound (\(f_{u,cell}\)) in parallel with total intracellular compound accumulation (\(C_{cell}\)), as described below. Intracellular compound bioavailability (\(F_{ic}\)) was calculated as the ratio between intracellular unbound compound concentration (\(C_{u,cell} = f_{u,cell}C_{cell}\)) (the concentration available to bind intracellular targets) and the externally added compound concentration (\(C_{medium}\)):

\[
F_{ic} = \frac{f_{u,cell} \cdot C_{cell}}{C_{medium}} = f_{u,cell} \cdot \frac{C_{cell}}{C_{medium}} = f_{u,cell} \cdot Kp
\]

where \(Kp\) is the ratio \(C_{cell}/C_{medium}\).

In our previous work\(^25\), this ratio was termed unbound drug accumulation ratio (\(Kp_{u,cell}\)), in analogy to previous work on unbound drug exposure in brain tissue\(^5\). We introduce the more descriptive and general \(F_{ic}\) parameter here, since it is applicable also to situations where only a fraction of the extracellular compound is unbound. This is of interest in drug discovery settings where cellular screens are often performed in the presence of serum proteins. In such assay setups, the \(Kp_{u,cell}\) term is not applicable, since it assumes extracellular drug to be completely unbound.

Contrary to oral bioavailability, which is defined in terms of dose (i.e., amount), \(F_{ic}\) is defined in concentration terms. For that reason, \(F_{ic}\) can take any positive value and is not limited to the range between zero and one. However, \(F_{ic}\) can be interpreted in similar terms, as the fraction of the extracellular concentration that is available intracellularly.

#### Materials and compound set.

Cell culture reagents were acquired from Thermo Fisher Scientific Inc. or Sigma-Aldrich. HEK293 cells were originally obtained from Thermo Fisher Scientific and MDCK cells from ATCC. Test compounds of analytical grade (\(\geq 95\%\) purity) were obtained from Sigma-Aldrich, Toronto Research Chemicals or OChem Inc. Compounds were dissolved in dimethyl sulfoxide (DMSO) and stored at \(-20^\circ C\).

The compound set used in this study included 34 drug-like molecules (Supplementary Table 1), and was enriched in compounds previously described as OATP1B1 (n = 11)\(^7\) or P-gp substrates (n = 22)\(^5,56\), with ten of the compounds included being substrates of both drug-transporting proteins. ADMET Predictor, version 7.0 (Synamtics Plus, Lancaster, CA) was used to predict physicochemical characteristics of the compound set.

The compounds were generally within the boundaries of Lipinski’s rule of 5 (64% of molecules within all rule-of-5 boundaries\(^7\)), with the major violation being MW > 500 Da (83% of violations). At physiological pH (pH = 7.4), eight of the included compounds were predicted to be predominantly negatively charged, fourteen as mostly unchanged, nine as mostly positively charged, and three as zwitterions (Supplementary Table 1). In agreement with the literature\(^24\), substrates of OATP1B1 were generally negatively charged and more polar than non-substrates (mean PSA of 105 vs. 74 Å\(^2\); \(p = 0.03\)). Compounds previously described as substrates of P-gp...
were more polar (mean PSA of 99 vs. 56 Å²; p = 0.002) and larger (mean MW of 497 vs. 335; p < 0.001) than compounds lacking reports of P-gp-mediated transport.

**Cell culture and preparation of experiments.**  **Cell line maintenance.**  HEK293 cells stably transfected with OATP1B1 (SLCO1B1*1a allele) or with the corresponding empty vector were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 2 mM L-glutamine and 75 μg/ml hygromycin B. CRISPR-Cas9 canine P-gp knock-out MDCK cells (P-gp–KO) were generated as described elsewhere. MDCK cells were grown in DMEM (glucose 1 g/l) supplemented with 10% FBS, GlutaMAX, penicillin (100 units/ml), and streptomycin (100 μg/ml). For human P-gp (ABCB1) transfected MDCK cells, hygromycin B (375 μg/ml) was included as a selection antibiotic. All cell lines were kept at 37 °C in a humidified 5% CO₂ atmosphere and subcultivated twice a week with a 1:6 ratio.

**Isolation of primary human hepatocytes.**  Hepatocytes were isolated from human liver tissue obtained from two donors undergoing surgical resections at the Department of Surgery, Uppsala University Hospital. Ethical approval was granted by the Uppsala Regional Ethics Committee (ethical approvals no. 2009/028 and 2011/037), donors gave informed consent and all studies were performed in accordance with the current national regulations and ethical guidelines. Isolation was performed using a two-step collagenase procedure, as previously described.

**Preparation of cell homogenates for cell binding experiments.**  For intracellular fraction of unbound compound experiments, cell lines or primary cells were suspended to a concentration of 10 × 10⁶ cells/ml in Hank’s balanced salt solution (HBSS). Cell suspensions were homogenized for 10 s with a VCX-500 ultrasonic processor (Sonics & Materials) at 20% intensity and used immediately for binding experiments.

**Preparation for intracellular compound accumulation experiments.**  HEK293 cells were grown for two days in 24-well plates, as previously described. MDCK cells were grown for two days in 96-well plates, prior to the intracellular compound accumulation experiments. The growth media were the same as described in the section ‘Cell line maintenance’ without addition of hygromycin B to avoid transporter inhibition.

For the corresponding experiments in human hepatocytes, two cell culture configurations were used: freshly isolated cells in suspension, or cells cultivated as monolayers for 24 h. The freshly isolated cells were suspended directly after isolation at a concentration of 5 × 10⁶ cells/ml in HBSS. The cell suspension (100 μl) was added to 96-well plates, which were immediately used for accumulation experiments. The monolayer-cultured hepatocytes were allowed to attach to collagen-coated 24-well plates for three hours (375 000 cells/well) in DMEM with HEPES supplemented with 5% FBS, penicillin (100 units/ml), streptomycin (100 μg/ml), insulin (4 μg/ml), and dexamethasone (1 μM). After cell attachment, medium was replaced by hepatocyte maintenance medium (Lonza) supplemented with insulin (10 μg/ml), transferrin (5.5 μg/ml), selenium (5 ng/ml), penicillin (100 units/ml), streptomycin (100 μg/ml), and dexamethasone (0.1 μM). Cells were kept in maintenance medium for 24 h, until the start of the intracellular compound accumulation experiments.

**Intracellular fraction of unbound compound.**  Binding to cell homogenates at steady-state was measured using dialysis as previously described. Simultaneous measurement of intracellular binding for multiple compounds (cassette-mode) is made possible by our previous observation that binding to cellular structures at low compound concentration is mainly non-specific and unaffected by the presence of other drug-like molecules.

Briefly, the cell homogenate was spiked with six randomly-chosen compounds. Homogenates were dialyzed for 4 h using a Rapid Equilibrium Dialysis device (Thermo Fisher Scientific Inc.). At the end of the incubation, uniform sample matrices were obtained by addition of blank homogenate to samples from the buffer chamber, and by addition of blank buffer to samples from the homogenate chamber. Dilutions of samples from the homogenate chambers (10-fold and 100-fold) were prepared. Protein was precipitated with acetonitrile/water (60:40) spiked with 50 mM warfarin (internal standard). Samples were centrifuged for 20 min at 2465 × g. Compounds in the supernatant were analyzed by ultra-performance liquid chromatography coupled to tandem mass spectrometry (UPLC-MS/MS), as described below.

The fraction of compound that was not bound to cell homogenate (fu,hom) was calculated according to:

\[
 fu,hom = \frac{PA_{\text{buffer}}}{PA_{\text{hom}}} 
\]

(2)

where PABuffer is the peak area of compound in the buffer chamber and PAHom is the peak area of compound in the homogenate chamber, both corrected for the peak area of the internal standard in the respective chamber. For each compound, the dilution of the homogenate sample where PAHom was closest to PABuffer was used.

The fraction of unbound compound in the cell (fu,cell) was calculated according to:

\[
 fu,cell = \frac{1}{D \cdot (1/fu,hom - 1) + 1} 
\]

(3)

where D was estimated for each homogenate preparation based on a cellular volume of 6.5 μl/mg protein, and on the protein concentration measured using the BCA Protein Assay Reagent Kit (Thermo Fisher Scientific Inc.).

**Intracellular compound accumulation.**  Intracellular compound accumulation at steady-state was measured essentially as described before. Briefly, cells were washed twice with HBSS and incubated for 45 min with solutions of single test compounds (in HBSS) on an orbital shaker (300 rpm) at 37 °C. At the end of the
experiment, the remaining extracellular compound solutions were collected and cells were washed twice with ice-cold phosphate-buffered saline pH 7.4 (PBS). For hepatocytes in suspension, the cells were separated from the extracellular solutions by centrifugation (100 × g for 5 min). Compounds were extracted from the cells using acetonitrile/water (60:40) spiked with 50 nM warfarin (internal standard). Extracellular medium samples were diluted 10-fold with the same extraction solution. All samples were centrifuged for 20 min at 2465 × g and compounds were quantified in the supernatant by UPLC-MS/MS, as described below. Protein concentrations were measured in representative wells of the plates using the BCA Protein Assay Reagent Kit (Thermo Fisher Scientific Inc.).

The ratio between total compound concentrations in the cells and in the medium (Kp) was calculated according to:

\[ K_p = \frac{A_{\text{cell}}}{C_{\text{medium}}} \]

where \( A_{\text{cell}} \) is the amount of compound in the cell samples, \( V_{\text{cell}} \) is the cell volume (6.5 \( \mu \)L/mg protein\(^{\text{80}} \)), and \( C_{\text{medium}} \) is the compound concentration in the medium sample.

**Metabolic clearance (CL\(_{\text{int}}\)) in primary human hepatocytes.** Metabolic clearance (CL\(_{\text{int}}\)) was determined, as previously described\(^{\text{81}} \). Briefly, freshly isolated human hepatocytes (in suspension or after 24 h of culture in monolayer format) were incubated with 1 \( \mu \)M compound solution (total incubation volume = 200 \( \mu \)L). At time points 0, 10, 30, and 60 min, the reaction was stopped by addition of acetonitrile (200 \( \mu \)l) spiked with 50 nM warfarin (internal standard). Samples were then centrifuged for 20 min at 2465 × g and compounds were quantified in the supernatant by UPLC-MS/MS, as described below. CL\(_{\text{int}}\) was calculated as the product of the compound disappearance rate and incubation volume, normalized by the cell number.

**Compound quantification.** Compound quantification was performed using UPLC-MS/MS. The system consisted of a Waters Xevo TQ MS with electrospray ionization coupled to a Waters Acquity UPLC. Compounds were chromatographically separated with a 2 min gradient elution (flow rate of 0.3 ml/min) on a Waters BEH C18 column, 2.1 × 50 mm (1.7 \( \mu \)m) at 60 °C, as described elsewhere\(^{\text{35}} \). Mass transitions for each compound and their respective cone voltages and collision energies can be found in Supplementary Table 7.

**Protein quantification of drug-transporting proteins and drug-metabolizing enzymes.** Protein concentrations for drug-transporting proteins and drug-metabolizing enzymes were quantified as previously described\(^{\text{44}} \). Briefly, a cell pellet (10 × 10⁶ cells) of freshly isolated human hepatocytes or of cells cultured in monolayer format was lysed with 100 mM Tris/HCl, pH 7.8, 2% (m/v) SDS and 50 mM DTT. Proteins were denatured at 95 °C for 5 min. The lysates were sonicated and then clarified by centrifugation. Trypsin digestion was performed using the filler-aided sample preparation protocol\(^{\text{82}} \). Peptides were eluted and spiked with stable-isotope labelled standard peptides (JPT Peptide Technologies) and analyzed with UPLC-MS/MS as previously described\(^{\text{44}} \). The sequences of proteospecific peptides have been previously published\(^{\text{83,84}} \).

**Intracellular compound accumulation simulations.** Intracellular compound accumulation was simulated, as previously described\(^{\text{44}} \), with a mechanistic model developed in R version 3.1.1 (http://www.r-project.org) using the deSolve package for differential equation solving. Integration was performed using the ode function, which automatically selects an integration algorithm of appropriate stiffness. The model consisted of two compartments (extracellular and intracellular), with model dimensions set to reflect the setup used in accumulation experiments in MDCK cells (apical surface area = 1 × 10⁻⁴ m², extracellular volume = 200 \( \mu \)l, total intracellular volume = 0.25 \( \mu \)l). In simulations intended to reflect compound accumulation in MDCK P-gp knockout cells, compound transport across the apical membrane was incorporated as a combination of transporter-mediated uptake (assumed to follow regular Michaelis-Menten kinetics) and bidirectional passive Fick’s diffusion. Simulations of compound accumulation in MDCK cells transfected with human P-gp also included a saturable efflux component in the apical membrane, while simulations of compound accumulation in cells where active transport has been inhibited with cyclosporine A included only bidirectional passive Fick’s diffusion. All combinations of low, medium, and high values of passive permeability (0.1, 1, and 10 × 10⁻⁶ cm/s), and \( V_{\text{max}} \) (10, 100, and 1000 pmol/min/mg protein) and \( K_m \) (1, 10, and 100 \( \mu \)M) for each of the transporters were simulated. Each simulation was repeated at 20 concentrations in the range 0.01 \( \mu \)M–100 \( \mu \)M. Drug accumulation was simulated over 120 min, and the resulting equilibrium intra- and extracellular unbound concentrations were used to calculate \( F_{\text{ic}} \). The concentration dependencies of \( F_{\text{ic}} \) were then visually inspected to identify scenarios that matched experimental observations.

**Statistics.** All experiments were performed in triplicate on at least 2 independent occasions. Atorvastatin and lopinavir were used as controls in intracellular fraction of unbound compound experiments (added to all cultures: average coefficient of variation (CV) in a single experiment <20%; day-to-day CV <10%), and in intracellular compound accumulation experiments (in parallel with tested compound: average CV in a single experiment <10%; day-to-day CV <10%). All results are presented as geometric mean ± standard error of the mean (S.E.M.), unless otherwise stated. A p-value < 0.05 was considered significant.

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**Author Contributions**

A.M. and A.T. performed all measurements of intracellular compound bioavailability. C.W. performed the proteomics measurements. M.K. established all cell lines used in this study. A.M., A.T., P.M. and P.A. performed the majority of data analysis and wrote the manuscript with critical input and contributions from all co-authors. A.M., A.T., P.M. and P.A. designed and supervised the study.

**Additional Information**

*Supplementary information* accompanies this paper at http://www.nature.com/srep

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