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### Effect of CFTR correctors on the traffic and the function of intracellularly retained ABCB4 variants

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**Abbreviations:** ABC, ATP-binding cassette; ABCB4<sup>cc</sup>, ABCB4-closed conformation; ABCB4<sup>if</sup>, ABCB4-inward facing conformation; ATP, adenosine triphosphate; CFTR, cystic fibrosis transmembrane conductance regulator; Cp, coupling helix; ER, endoplasmic reticulum; HEK, human embryonic kidney; NBD, nucleotide binding domain; PC, phosphatidylcholine; PFIC3, progressive familial intrahepatic cholestasis type 3; TM, transmembrane helix; UDCA, ursodeoxycholic acid; WT, wild type.

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#### ABSTRACT

**Background & aim:** ABCB4 is expressed at the canalicular membrane of hepatocytes. This ATPbinding cassette (ABC) transporter is responsible for the secretion of phosphatidylcholine into bile canaliculi. Missense genetic variations of ABCB4 are correlated with several rare cholestatic liver diseases, the most severe being progressive familial intrahepatic cholestasis type 3 (PFIC3). In a repurposing strategy to correct intracellularly retained ABCB4 variants, we tested 16 compounds previously validated as cystic fibrosis transmembrane conductance regulator (CFTR) correctors.

**Methods:** The maturation, intracellular localization and activity of intracellularly retained ABCB4 variants were analyzed in cell models after treatment with CFTR correctors. In addition, *in silico* molecular docking calculations were performed to test the potential interaction of CFTR correctors with ABCB4.

**Results:** We observed that the correctors C10, C13 and C17, as well as the combinations of C3+C18 and C4+C18, allowed the rescue of maturation and canalicular localization of four distinct traffic-defective ABCB4 variants. However, such treatments did not permit a rescue of the phosphatidylcholine secretion activity of these defective variants and were also inhibitory of the activity of wild type ABCB4. *In silico* molecular docking analyses suggest that these CFTR correctors might directly interact with transmembrane domains and/or ATP-binding sites of the transporter.

**Conclusion:** Our results illustrate the uncoupling between the traffic and the activity of ABCB4 since the same molecules can rescue the traffic of defective variants while they inhibit the secretion activity of the transporter. We expect that this study will help to design new pharmacological tools with potential clinical interest.

#### **K**EYWORDS

ABC transporters; Bile secretion; Cell models; Cholestatic liver diseases; Molecular docking; Targeted pharmacotherapy.

#### LAY SUMMARY

- CFTR correctors rescue the maturation and the *in vitro* localization of four distinct ERretained ABCB4 variants identified in patients.
  - CFTR correctors do not rescue the function of these variants even if they are relocalized at the plasma membrane.
  - CFTR correctors inhibit the function of wild type ABCB4.
- *In silico* molecular docking analyses suggest direct interactions of CFTR correctors with functional domains of ABCB4.

#### INTRODUCTION

Bile secretion is an essential function of the liver for lipid digestion and absorption as well as the elimination of xenobiotics and endogenous metabolites. This function mainly depends on transporters localized at the apical (or canalicular) plasma membrane of hepatocytes such as ATPbinding cassette subfamily B member 4 (ABCB4), also known as multidrug resistance protein type 3 (MDR3). ABCB4 belongs to the superfamily of ATP-binding cassette (ABC) transporters which are transmembrane proteins able to bind and hydrolyze ATP in order to fulfil their biological functions<sup>1,2</sup>. The expression of ABCB4 is restricted to the canalicular membrane of hepatocytes<sup>3</sup> and its role is to ensure secretion of phosphatidylcholine (PC) into bile<sup>4</sup>. With the co-secreted cholesterol and bile salts, PC forms mixed micelles in the aqueous environment of bile, thus avoiding the formation of cholesterol gallstones in the biliary tract as well as cell membrane damage by free bile acids on the canalicular membrane of hepatocytes and the apical membrane of cholangiocytes (for reviews, see<sup>1,5</sup>). Until now, more than 220 distinct variations of the ABCB4encoding gene have been reported, mostly in patients with cholestatis and cholelithiasis (see https://evs.gs.washington.edu/EVS/ and http://abcmutations.hegelab.org/). These genetic variations of ABCB4 can affect the expression, the traffic, the function or the stability of the protein. Indeed, we have previously proposed a classification of these variants into five distinct classes: class I with no protein expression, class II with intracellular retention; class III with functional defects, class IV with stability impairment and class V with no apparent defect<sup>6,7</sup>. More details about ABCB4 function, genetic disorders and subsequent diseases can be found in the recent review by Kroll et al.8.

The most severe ABCB4-related disease is progressive familial intrahepatic cholestasis type 3 (PFIC3), which is a rare autosomal recessive disease affecting homozygous or compound heterozygous patients during childhood<sup>9</sup>. PFIC3 appears during the first months of life and is characterized by chronic cholestasis, jaundice and pruritus<sup>10</sup>. The only pharmacological treatment for PFIC3 patients is the administration of ursodeoxycholic acid (UDCA), a bile acid with low hydrophobicity which renders the bile less toxic<sup>11,12</sup>. However, more than half of PFIC3 patients display no or little response to UDCA treatment and worsening of the disease most often requires liver transplantation<sup>10,13</sup>. Therefore, the therapeutic challenge is to identify new targeted

pharmacotherapies as an alternative to liver transplantation for patients with severe forms of ABCB4-related diseases.

Looking for correctors of class II endoplasmic reticulum (ER)-retained ABCB4 variants, we have recently shown that structural analogues of roscovitine are able to partially rescue the traffic, localization and function of these variants<sup>14</sup>. In the present study, in order to pursue this repositioning strategy, we were interested in correctors previously shown to rescue the plasma membrane targeting of the F508del variant of ABCC7/CFTR (cystic fibrosis transmembrane conductance regulator), the ABC transporter mutated in patients with cystic fibrosis (see Supplementary Table S1, and references therein). While we observed that CFTR correctors are able to rescue the maturation and the localization of several ER-retained ABCB4 variants, we report that they are not able to rescue their PC secretion activity and that they also inhibit the activity of the wild type transporter. These two effects might be due to direct interactions of CFTR correctors with functional domains of ABCB4 such as transmembrane domains and/or nucleotide binding domains, as suggested by *in silico* molecular docking calculations.

Accepted

#### MATERIALS AND METHODS

#### Plasmids, cell culture and transfection

Plasmids encoding WT and ER-retained ABCB4 missense variants (I541F, I490T, L556R) have been described<sup>14-16</sup>. The R545H variant was previously reported<sup>17</sup> and site-directed mutagenesis was performed as described<sup>14</sup> using the following primers (Eurogentec, Angers, France): 5'-GATCGCCATTGCACATGCCCTGGTTCGCA-3'(forward) and 5'-TGCGAACCAGGGCATGTGCAATGGCGATC-3' (reverse). Human embryonic kidney (HEK-293, herein referred to as HEK; ATCC®-CRL-1573<sup>TM</sup>) cells and human hepatocellular carcinoma HepG2 (ATCC®- HB-8065<sup>TM</sup>) cells were obtained from ATCC (Manassas, VA). Cells were grown at 37°C with 5% CO<sub>2</sub> in Dulbecco's Modified Eagle Medium (Thermo Fisher Scientific, Villebon-sur-Yvette, France) containing 4.5 g/L D-glucose and supplemented with 10% heatinactivated fetal bovine serum (Sigma, Saint-Quentin Fallavier, France), 2 mM L-glutamine, 2 mM sodium pyruvate, 100 units/mL penicillin and 100 µg/mL streptomycin (Thermo Fisher Scientific). Transient transfections were performed using Turbofect (Thermo Fisher Scientific) at a ratio of reagent:DNA of 2:1 for HEK cells, and JetPrime (PolyPlus Transfection, Illkirch, France) at a ratio of reagent:DNA of 2:1 for HepG2 cells, according to manufacturers' instructions. To maintain similar ABCB4 expression and avoid fluctuation of protein expression levels from one condition to another, cells were split in separated wells only after transfection of the whole population in 10 cm Petri dishes and overnight expression of the transgene.

#### Chemicals and cell treatments

CFTR correctors (see details in Supplementary Table S1) were kindly provided by the CFTR Chemical Compound Program of the Cystic Fibrosis Foundation (Chicago, IL). All compounds were solubilized in dimethylsulfoxide (DMSO) as 1000X concentrated stock solutions in order to treat cells with 10  $\mu$ M final concentration, using DMSO as control vehicle at the same dilution (0.1% DMSO for all conditions). Twenty-four hours post-transfection (except for cytotoxicity assays), cells were treated during 16 hours with these drugs. After drug treatment, cells were used for cytotoxicity assays, immunoanalyses, or PC secretion assays.

#### Cytotoxicity assays

Cytotoxicity of CFTR correctors was assessed by the conversion of MTT (3-[4,5-dimethylthiazol-2-yl]-2,5 diphenyl tetrazolium bromide; Sigma) into formazan crystals by living cells, as described<sup>18</sup>. In brief, subconfluent HEK cells were plated in 96-well plates in triplicate for each tested condition, including controls (no cells, treatment with vehicle only). Eight hours after cell seeding and 16 hours after drug treatment, 125  $\mu$ g/mL MTT (final concentration) was added in each well and cells were re-incubated at 37°C for 2 hours. Then culture media were gently washed out, cells were lysed in 100  $\mu$ L of DMSO and absorbance at 540 nm was measured using a multiplate cytofluorimeter SpectraFluor from Tecan (Männedorf, Switzerland). Cytotoxicities were calculated for each triplicate and after background subtraction, means were expressed as percentages of the mean for cells treated with vehicle only.

## Immunoblots, immunofluorescence and measurement of ABCB4-mediated phosphatidylcholine secretion

Immunoblots and indirect fluorescence analyses were performed as previously described<sup>14</sup>, using the following primary antibodies: mouse monoclonal anti-ABCB4 (clone P<sub>3</sub>II-26) and anti-ABCC2 (clone M2I-4) from Enzo Life Sciences (Villeurbanne, France); anti- $\alpha$ -tubulin (clone 1E4C11) from ProteinTech (Manchester, United Kingdom). Peroxidase- and fluorochromeconjugated secondary antibodies were from Sigma and Thermo Fisher Scientific, respectively. Immunoblots were quantified in the linear range of detection using ImageJ 1.50i software (U.S. National Institutes of Health, Bethesda, MD). Immunofluorescence images were acquired using a confocal microscope (Eclipse TE-2000-Nikon-C2) equipped with a 60X objective, serial xy optical sections with a z-step of 0.3  $\mu$ m were taken using Nikon NIS-Elements software version AR 4.50 with constant settings (laser powers and correction of signal intensities) and treated using Adobe Photoshop version 8.0.1. Measurement of ABCB4-mediated PC secretion using a fluoroenzymatic assay was performed as described<sup>19</sup>. Each condition was analysed in triplicate and the secreted PC was calculated as follows:

$$N_{x} = \frac{(V_{x+} - V_{bgrd+}) - (V_{x-} - V_{bgrd-})}{A}$$
(1)

with  $N_x$  the amount of PC in nmol;  $V_{x-}$  and  $V_{x+}$  the fluorescence means for the tested condition in the absence or presence of phospholipase D, respectively;  $V_{bgrd-}$  and  $V_{bgrd+}$  the background fluorescence means in the absence or presence of phospholipase D, respectively; A the slope of the standard curve (in AU.nmol<sup>-1</sup>). Then, results were expressed as percentages of wild type ABCB4 (ABCB4-WT) activity as follows:

$$M_x = \frac{N_x - N_{mock}}{N_{WT} - N_{mock}} \times \frac{D_{WT}}{D_x} \times 100$$
 (2)

with  $M_x$  the PC secretion activity as a percentage of ABCB4-WT activity;  $N_x$ ,  $N_{mock}$  and  $N_{WT}$  the amounts of secreted PC for the tested condition, mock-transfected cells and ABCB4-WT expressing cells, respectively, according to (1);  $D_x$  and  $D_{WT}$  the expression levels of mature ABCB4 determined by densitometry analysis from immunoblots with the corresponding cell lysates for the tested condition and ABCB4-WT expressing cells, respectively.

#### In silico molecular docking and calculations

Two conformations of human ABCB4 (*h*ABCB4) were considered for molecular docking calculations, namely inward-facing and closed conformations (ABCB4<sup>if</sup> and ABCB4<sup>cc</sup>, respectively). ABCB4<sup>if</sup> was built by homology modelling using the inward-facing human ABCB1 (*h*ABCB1) protein data bank structure (PDB ID: 6QEX)<sup>20</sup>, given the high sequence identity between *h*ABCB1 and *h*ABCB4 (76.8%, see Supplementary Figure S1). ABCB4<sup>cc</sup> model was built using the recently resolved cryo-EM structure of ABCB4<sup>cc</sup> structure trapped in ATP-bound state (PDB ID: 6S7P)<sup>21</sup>. The non-resolved extracellular loop (residues 85-104) of ABCB4<sup>cc</sup> was built based on the resolved *h*ABCB1 structure. For both models, L<sub>1</sub>-linker connecting nucleotide binding domain (NBD) 1 to transmembrane helix (TM) 7, and the N- and C-terminal domains were not included in the present study owing to their absence in both resolved *h*ABCB1 and *h*ABCB4 structures (Supplementary Figure S1). Present models did not include the ATP molecules. Homology modeling were performed using the Modeller software version 9.23<sup>22-24</sup>. Prior to docking calculations, both models were minimized in solvated membrane lipid bilayers that were removed for docking calculations (for further details, see *Supplementary Materials and Methods*).

Molecular docking calculations were carried out using the Autodock Vina software<sup>25</sup>. Given the absence of *a priori* knowledge about the interaction mechanism of CFTR correctors, the following procedure was performed to extensively sample plausible binding sites of CFTR correctors to both ABCB4<sup>if</sup> and ABCB4<sup>cc</sup> conformers. First, so-called "blind" docking calculations were performed with all ligands in which the whole protein was considered as intentionally too large volume search space centered on the protein center of mass for both ABCB4<sup>if</sup> and ABCB4<sup>cc</sup> conformations (Supplementary Table S2A). A set of 20 replicas per CFTR corrector was performed, using different random seeds, providing a total of approximately 6400 poses per ABCB4 conformation. These molecular poses exhibited clusters located at different spatial regions (see Supplementary Figure S2). Refined docking calculations were then performed in which smaller space search volumes were used (Supplementary Figure S3), which do not exceed the recommended search volume of 27 000 Å<sup>3</sup> to ensure the reliability of docking calculation results. Again, 20 replicas of refined molecular docking calculations were performed for each CFTR corrector. For blind and refined docking calculations, exhaustiveness was set to 40 and 100, respectively, in order to increase the computational effort used during molecular pose search<sup>26</sup>. The maximum number of poses was set to 100 by replica and the initial affinity cutoff was defined at 15 kcal.mol<sup>-1</sup> with respect to the top pose of a given replica. The list of flexible residues allowed for each volume search space is reported in Supplementary Table S2B. Given the large number of plausible flexible residues, only sidechain  $C_{\beta}$ - $C_{\gamma}$  bonds were allowed to rotate as a compromise within the limit of 32 rotatable bonds allowed by Autodock Vina software.

Additional information about ABCB4 and ligand preparations as well as analyses are provided in Supplementary Materials and Methods.

#### Statistics

Graphics and one-way ANOVA tests were performed using Prism version 7.00 (GraphPad Software, La Jolla, CA). A *P* value of less than 0.05 was considered significant with \*: p<0.05; \*\*: p<0.01; \*\*\*: p<0.005; ns: not significant. Symbols always indicate the comparison between the control (vehicle-treated) and the other tested conditions.

#### RESULTS

#### **CFTR** correctors rescue the traffic of the ER-retained I541F variant of ABCB4

In order to identify new pharmacological correctors for ER-retained ABCB4 variants, we tested the possibility to repurpose CFTR correctors previously reported to rescue the plasma membrane targeting of F508del-CFTR. We tested the effect of 16 CFTR correctors (Supplementary Table S1). Using MTT assay, we first ascertained that none of these molecules were cytotoxic after 16 hours of treatment at 10 µM in HEK cells (Figure 1). As a model of ER-retention of ABCB4, we used the well-characterized I541F variant, for which plasma membrane targeting could be restored at low temperature (27°C) and by pharmacological means<sup>14-16</sup>. On immunoblots from HEK cell lysates, ABCB4-I541F is mainly present at an apparent molecular weight of 140 kDa due to its incomplete glycosylation, by opposition to ABCB4-WT mainly present as a fully glycosylated protein with an apparent molecular weight of 160 kDa (Figure 2A), as previously demonstrated<sup>15</sup>. After 16 hours of cell treatment with 10 µM of the different CFTR correctors, we observed that some of them were able to correct ABCB4-I541F maturation, considering the increased amounts of its mature form (Figure 2A). The quantification of these results indicated that C10 (KM11057), C13 (Corr-4C) and C17 (15jF) were the best ABCB4 corrector candidates, as well as combinations of C3+C18 (VRT-325+VRT-534) and C4+C18 (Corr4A+VRT-534) (Figure 2B), previously reported to be able to rescue F508del-CFTR as well as defective ABCA4 variants<sup>27,28</sup>. We then investigated the potential of these correcting molecules to restore canalicular targeting of ABCB4-I541F in HepG2 cells, a human hepatoma-derived cell line forming pseudo-bile canaliculi in culture<sup>29</sup>. In these polarized cells, ABCB4-WT colocalized with ABCC2 at bile canaliculi, which was not the case of ABCB4-I541F displaying a diffuse cytoplasmic staining (Figure 2C). However, after 16 hours of treatment with 10 µM of C10, C13, C17, or the C3+C18 and C4+C18 combinations, canalicular staining of ABCB4-I541F was rescued, as shown by its colocalization with ABCC2 (Figure 2C). Altogether, these results indicate that C10, C13, C17, and C3+C18 and C4+C18 combinations are able to partially rescue the maturation and the canalicular localization of ABCB4-I541F.

## CFTR correctors also rescue the maturation and the localization of three other ER-retained ABCB4 variants

In order to determine if these CFTR corrector candidates might be envisioned as general correctors for ER-retained ABCB4 variants, we investigated their effect on three other ABCB4 variants (1490T, R545H, L556R), previously reported in patients with liver diseases<sup>9,17,30</sup>, and already shown to be class II ER-retained for the I490T and L556R variants<sup>6,14</sup>. Our results give evidence that C10, C13, C17, and C3+C18 and C4+C18 combinations are also able to rescue the maturation of these three immature variants (Figure 3A-C). The quantifications of these experiments indicate that rescuing efficiencies were all statistically significant, even though efficiencies vary among the three variants (Figure 3D-F). As performed for ABCB4-I541F, we analysed the intracellular localization of these three variants in HepG2 cells. After vehicle treatment, ABCB4-I490T, -R545H and -L556R do not colocalize with ABCC2 (Figure 3G-I, upper panels). But after treatment with C10, C13 or C17, as well as with C3+C18 or C4+C18 combinations, the localization of the three variants at bile canaliculi was rescued (Figure 3G-I). Thus, these CFTR correctors are able to rescue the maturation and the localization of four distinct ER-retained ABCB4 variants, suggesting that they trigger common correcting molecular mechanisms.

### CFTR correctors do not restore the function of ER-retained ABCB4 variants and inhibit ABCB4-WT function

The aim of correcting the intracellular traffic of ER-retained ABCB4 variants is to obtain sufficient amounts of functional transporters present at the plasma membrane in order to rescue PC secretion in the extracellular space. Using a previously described fluoro-enzymatic approach<sup>19</sup>, we measured ABCB4-mediated PC secretion of the four variants transiently expressed in HEK cells with or without treatment with the different CFTR correcting drugs. As expected, we observed a strongly reduced activity (less than 10-15% of ABCB4-WT activity) for the four variants in the absence of treatment by comparison to ABCB4-WT (Figure 4A-D). Nevertheless, none of the CFTR correctors (used alone or in combination) was able to significantly rescue the function of any of the four variants (Figure 4A-D), despite their partial rescue of maturation and canalicular localization (Figures 2 and 3). Instead, we observed a trend towards less activity in treated cells,

even if differences did not reach statistical significance (Figure 4A-D). We also tested these CFTR correctors in HEK cells expressing ABCB4-WT. Surprisingly, we observed that 10  $\mu$ M of C10, C13, C17, C3+C18 or C4+C18 dramatically inhibited ABCB4-WT activity to less than 10% of its maximal activity (Figure 4E). Similar inhibitory effects of these CFTR correctors were also observed on ABCB4-WT when used at 5  $\mu$ M (data not shown). A possible interaction of CFTR correctors with membrane lipids leading *in fine* to their release into the extracellular medium is unlikely or negligible since our results indicate that ABCB4-WT-mediated PC secretion is dramatically reduced to 1.6–6.0% of control activity after treatment with these compounds (Figure 4E). Altogether, our results provide evidence that, despite their capacity to restore ABCB4 targeting at the canalicular membrane, the CFTR correctors investigated here also inhibit PC secretion function mediated by the transporter.

## Molecular docking calculations reveal different plausible binding regions of CFTR correctors within ABCB4

We hypothesized that the inhibition of ABCB4 activity by CFTR correctors could be related to their direct interaction with functional domains of the transporter. To test this possibility, we performed in silico molecular docking simulations. Blind calculations were first performed using large box parameters (Supplementary Table S2A) with inward-facing ABCB4 (ABCB4<sup>if</sup>) and its closed conformation (ABCB4<sup>cc</sup>) wherein no a priori plausible structural binding regions were defined (for details, see *Materials and Methods*). This provided 6398 and 6395 molecular poses for ABCB4<sup>if</sup> and ABCB4<sup>cc</sup>, respectively. Molecular poses were then spatially clustered, revealing a single binding site for CFTR correctors within ABCB4<sup>if</sup> while three binding sites were obtained for ABCB4<sup>cc</sup> (Supplementary Figure S2). In ABCB4<sup>if</sup>, 95% of molecular poses are located in ABCB4 protein chamber (Supplementary Table S3), expected to be the canonical phospholipid binding site region<sup>21</sup>. Regarding ABCB4<sup>cc</sup>, 81% of molecular poses can be initially divided into three distinct regions, namely an alternative site at the lipid-protein interface (30% of molecular poses) and the two known ATP-binding sites at both NBD interfaces (15 and 36% for ATPbinding sites 1 and 2, respectively - see Supplementary Table S3). Given the low sensitivity of blind docking calculations, regions obtained from blind docking calculations on ABCB4<sup>if</sup> were also explored for refined docking calculations on ABCB4<sup>cc</sup> and vice versa (Supplementary Figure S3). It is worth mentioning that no molecular poses were observed for ATP-binding site regions on ABCB4<sup>if</sup>, which makes sense considering the large distance between the two NBDs in this conformation. Altogether, refined molecular docking calculations on ABCB4<sup>if</sup> highlights the relevance of the so-called protein chamber (Figure 5A). Regarding ABCB4<sup>cc</sup>, molecular docking calculations suggested six possible regions for CFTR corrector binding; namely protein chamber, ATP-binding sites 1 and 2 and three alternative sites 1, 2 and 3 at the lipid protein interface (Supplementary Figure S3 and Figure 5A).

Thorough analysis over all molecular poses were then performed for each binding site to stress out key residues possibly involved in CFTR corrector binding to ABCB4. Only poses exhibiting binding affinity difference with respect to top-ranked molecular poses below 2.0 kcal.mol<sup>-1</sup> were considered as already described<sup>31</sup>. In total, a set of 38670 molecular poses over 50482 were obtained from refined molecular docking calculations (Supplementary Table S4). Accounting aforementioned in vitro results, special attention was paid to selected CFTR correctors, namely C3, C4, C10, C13, C17 and C18 (Figure 5A). Contact analyses using a 4.5 Å cutoff were performed allowing to decipher key residues to CFTR corrector binding in the four regions of interest in ABCB4 (Figure 5B). It is worth mentioning that contact analyses performed over the whole set of CFTR correctors exhibit similar profiles with respect to selected CFTR correctors (Supplementary Figure S4). Eleven residues are in contact with the selected CFTR correctors in at least 50% of calculated molecular poses, in ABCB4<sup>if</sup> protein chamber (Figure 5B). Most of these residues are either aromatic (histidine, phenylalanine and tyrosine) or aliphatic (alanine, leucine), and presumably involved in substrate binding (Figure 6A), as proposed recently for His989 and Ala990<sup>21</sup> in ABCB4<sup>if</sup> protein chamber (underlined residues in Figure 6A). Molecular docking calculations in ABCB4<sup>cc</sup> alternative site 1 clearly suggested the key role of Val192, Phe195, Ile354 and Phe357 for which contacts are higher than 90% (Figures 5B and 6A). Likewise, binding to alternative sites 2 and 3 involved mostly contacts with aliphatic residues (Leu842, Val864, Ile867, Ala868, Leu987, Ala990, Ser991 and Ala994 for alternative site 2 and Leu724, Ala727 and Leu761 for alternative site 3). Contact analyses of ABCB4<sup>cc</sup> ATP-binding sites and protein chamber exhibited more residues per site owing to smaller volumes as well as more important steric hindrance. Indeed, molecular docking calculations revealed seventeen, ten and seven residues for which contact rate is higher than 90% in ABCB4<sup>cc</sup> protein chamber, ATPbinding sites 1 and 2, respectively (Figure 5B). Regarding ATP-binding sites, residue profile again included aromatic amino acids (e.g. NBD A-loop residues Tyr403 and Tyr1043) but also polar residues such as arginine (Arg406 and Arg1046) (Figure 6A). Residue profile for ABCB4<sup>cc</sup> protein chamber included mostly polar and aliphatic residues (see Figures 5B and 6A). For all ABCB4 binding sites, thorough analyses of non-covalent interactions (H-bond and van der Waals interactions) were performed, highlighting the key role of  $\pi$ -stacking interactions between CFTR correctors' aromatic rings and ABCB4 aromatic residues (Supplementary Table S5). This was clearly shown for ABCB4<sup>cc</sup> alternative site 1 and ATP-binding sites 1 and 2 for which Phe195, Phe357, Tyr403 or Tyr1043 are involved in all  $\pi$ -stacked conformations. To a lesser extent, H-bonding were also shown to be relevant. For instance, H-bonding with Gln725 were involved in 21% of molecular poses in ABCB4<sup>cc</sup> ATP-binding site 1), Arg1046, or Glu1080 (ABCB4<sup>cc</sup> ATP-binding site 2) are involved in 27, 21 and 21% of molecular poses, respectively (Supplementary Table S5).

Finally, molecular docking calculations and aforementioned atomic-scaled analyses allowed to define putative binding sites in terms of secondary structure (Supplementary Table S6). ABCB4<sup>if</sup> protein chamber involved the lipid-embedded regions of all TMs but TM11, (Figure 6B). ABCB4<sup>cc</sup> alternative site 1 is defined by TM1, TM3, TM4 and TM6 (Figure 6B). Alternative sites 2 and 3 are located on the other side of ABCB4<sup>cc</sup>, involving TM1, TM4-6, TM7-9 and TM12 for site 2 and TM9, TM10-12 for site 3. Alternative sites 1 and 2 are interfacing with lower leaflet membrane while alternative site 3 is in contact with upper leaflet membrane. In addition to known residues involved in ATP binding, ABCB4<sup>cc</sup> ATP-binding site regions were also defined, including the coupling helices (CH), namely CHs between TM2 and 3 and TM10 and 11 for ATP-binding site 1, and CHs between TM4 and 5 and TM8 and 9 for ATP-binding site 2 (Figure 6B). Altogether, these *in silico* molecular docking analyses suggest that CFTR correctors might directly interact with functional domains of ABCB4, which could explain their effect on ABCB4 activity.

#### DISCUSSION

Genetic variations of the phospholipid floppase ABCB4 are correlated with rare cholestatic liver diseases, the most severe form being PFIC3. Some of these genetic variations can cause ABCB4 misfolding, its retention in the ER and, as a result, the loss of its phospholipid floppase function<sup>6</sup>. In previous studies, we and others have demonstrated that small molecules such as cyclosporins<sup>6,16</sup>, 4-phenylbutyrate (4-PBA), curcumin<sup>32</sup> and structural analogues of roscovitine<sup>14</sup> are able to partially rescue the intracellular traffic and the cell surface localization of ER-retained ABCB4 variants. However, cytotoxicity, inhibitory effect on ABCB4 function<sup>33</sup>, bioavailability or the high used concentration of these molecules are major limitations for their therapeutic use. In the present study, in order to identify new correctors for ER-retained ABCB4 variants, we explored the potential effect of several molecules, developed as CFTR correctors, on four distinct ER-retained ABCB4 variants identified in patients (I490T, I541F, R545H, L556R mapped in Supplementary Figure S5). CFTR correctors, which were used in this study are small molecules, mostly identified by high-throughput screening strategies to correct the traffic of F508del-CFTR, the most frequent genetic variation detected in patients with cystic fibrosis<sup>34</sup>. The extended use of these correctors is an attractive option for multiple rare diseases associated with protein misfolding and missorting. Interestingly, some of these molecules have been shown to successfully correct the intracellular localization and the function of other defective ABC transporters such as ABCA335 and ABCA4<sup>28</sup> and also ER-retained variants of ATP8B1<sup>36</sup> and  $\alpha$ -sarcoglycan<sup>37</sup>, proteins without CFTR similarity. Here, we show that the correctors C10, C13 and C17 as well as the combinations of C3+C18 and C4+C18 can partially rescue the maturation and the plasma membrane targeting of four defective ABCB4 variants. Further investigations at the molecular level will be required to determine the capacity of these CFTR correctors to rescue a proper folding of the class II ERretained ABCB4 variants studied here, thus allowing their plasma membrane targeting. The fact that the floppase activity of the I541F, I490T and L556R variants of ABCB4 can be partially rescued by roscovitine analogues<sup>14</sup> indicates that these missense variants have intrinsic activity once rescued at the plasma membrane. However, the CFTR correctors tested here are not able to restore the function of the ER-retained ABCB4 variants and they also inhibit the floppase activity of ABCB4-WT. While none of the compounds used in this study have been tested in vivo or in clinics, it is interesting to note that rare cases of cholestasis have been reported in patients taking sildenafil <sup>38,39</sup>, the parent molecule of C9 and C10, as well as cystic fibrosis patients treated with Orkambi, which contains VX-809/Lumacaftor – a derivative from C18/VX-534 (https://www.ema.europa.eu/en/documents/product-information/orkambi-epar-product-information\_en.pdf), arguing for a potential inhibitory effect of these molecules on canalicular ABC transporters.

It is noteworthy to mention that this is not the first time that inhibitory effects of correctors on ABCB4 function are reported. Indeed, cyclosporin A, roscovitine and triazole compounds such as itraconazole were shown to inhibit ABCB4 floppase activity<sup>14,33,40</sup>. Sildenafil, the C10 parent molecule, has also been shown to inhibit the transport function of ABCB1<sup>41</sup>, which exhibits 86%  $ABCB4^{42}$ . sequence-based similarity with The PgpRules prediction tool (https://pgprules.cmdm.tw/)<sup>43</sup> indicates that C3, C4, C10, C13, C17 and C18 might inhibit ABCB1 (data not shown), suggesting similar inhibitory mechanisms on ABCB4. It is important to note that PgpRules prediction tool is initially designed to predict ABCB1 inhibition. Given (i) the high sequence-based similarity between both transporters and (ii) the inhibitor overlaps for both transporters, one can expect that they might, at least partially, share inhibition mechanisms, despite the divergence of substrates translocated by these two transporters<sup>8</sup>. Furthermore, even if CFTR correctors belong to different chemical classes (quinolines, bithiazoles, quinazolines, pyrimidines)<sup>34</sup>, they share features with pharmacophore models of ABCB1 inhibitors previously described as hydrophobic, hydrogen bond acceptor, aromatic ring centered with positive ionizable moieties<sup>44</sup>.

To explain how these CFTR correctors could correct missorting of ABCB4 variants and at the same time inhibit the floppase activity of the transporter, we suggest that these small molecules may directly interact with ABCB4. Such an interaction has already been reported for C18 with CFTR<sup>45</sup>. To virtually test this hypothesis, we investigated the putative binding sites of CFTR correctors in ABCB4 by *in silico* docking simulations. Using two models of human ABCB4 in inward-facing and closed conformations (ABCB4<sup>if</sup> and ABCB4<sup>cc</sup>, respectively), we identified seven distinct plausible binding regions for CFTR correctors: protein chambers in both ABCB4<sup>iff</sup> and ABCB4<sup>cc</sup> models; and three alternative sites at the protein-lipid interface and two ATP-binding sites in ABCB4<sup>cc</sup>. Analysis of contact residues with a 4.5 Å cutoff around the docked poses of CFTR correctors indicated that several key residues could be involved in CFTR corrector binding. In ABCB4<sup>iff</sup>, we identified His989 and Ala990 as potential interactors with CFTR correctors, these residues having been recently shown to be crucial for PC binding and ABCB4

function<sup>21</sup>. We also detected Gln725, a residue conserved in ABCB1 and implicated in the binding of tariquidar<sup>46</sup>. Concerning ABCB4<sup>cc</sup>, we identified Phe357, Tyr403 and Arg1046 located in the alternative site 1, the ATP binding sites 1 and 2, respectively. These three residues were previously reported to be mutated in patients suffering from PFIC3 and low-phospholipid associated cholelithiasis syndrome<sup>47-49</sup>, suggesting their important role in ABCB4 expression and function. Moreover, Tyr403 and Arg1046 were also reported to be essential for ATP binding and hydrolysis<sup>21,50</sup>. It is thus tempting to speculate that the predicted direct interaction of CFTR correctors with key residues of ABCB4 may preclude its phospholipid floppase activity. This view is supported by the fact that sildenafil (the C10 parent molecule) and C3 impair the function of ABCB1 and CFTR, respectively<sup>41,51</sup>. It is important to note that present molecular docking calculations should be considered as an "idea generator" rather than a strong prediction. Thus, these aspects will require further joint experimental and theoretical confirmations, including µs-scaled molecular dynamics simulations. However, *in silico* findings might help in the design of site-directed mutagenesis variants to decipher small molecule-dependent inhibition mechanisms of ABCB4.

We propose that the trafficking rescue of ABCB4 variants mediated by CFTR correctors may be conferred through their direct interaction with critical domains of ABCB4, as suggested by our *in silico* docking analyses. We expect that our findings will help to design new non-inhibitory ABCB4 correctors that may lead to the preclinical development of pharmacological alternative to UDCA treatment and liver transplantation. Indeed, if our hypothesis is confirmed, the next challenge will be to identify small molecules able to promote folding and trafficking of defective variants without compromising their activity. Thus, further investigations are eagerly needed in order to optimize bioavailability and affinity of these correctors, not only to promote traffic correction but also to maintain the function of the transporter.

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#### **FIGURE LEGENDS**

Figure 1. CFTR correctors are not cytotoxic. HEK cells were treated with 10  $\mu$ M of the indicated compounds during 16 hours. Then cytotoxicity was assessed by MTT assay and expressed as the percentage of means for control vehicle-treated cells. Means (± SD) of at least three independent experiments per condition performed in triplicate are shown.

Figure 2. CFTR correctors partially rescue the maturation and the localization of ABCB4-1541F. (A) ABCB4-I541F was transiently expressed in HEK cells. After 16 hours of treatment with vehicle (ctrl, control) or 10 µM of the indicated compounds, cell lysates were prepared and analysed by immunobloting using the indicated antibodies. ABCB4-WT is shown as reference. The mature and immature forms of ABCB4 are indicated (arrows), as well as molecular weight markers. This panel is representative of at least five independent experiments for each condition. (B) Densitometry analysis of A. The amount of ABCB4-I541F mature form was quantified, normalized to the amount of tubulin, and then expressed as fold change compared to vehicletreated cells (ctrl). Dark grey bars indicate compounds for which means are significantly different from control. Means  $(\pm SD)$  of at least five independent experiments per condition are shown. (C) ABCB4-WT or ABCB4-I541F were expressed in HepG2 cells. After 16 hours of treatment with vehicle (control) or 10 µM of the indicated correctors, cells were fixed and permeabilized. Then, after immunolabelling, ABCB4 (red) and endogenous ABCC2 (green) were visualized by confocal microscopy. Nuclei shown in the merged images were labelled with Hoechst 33342 (blue). Asterisks in the lower frames indicate bile canaliculi. This panel is representative of at least three independent experiments per condition. Bars: 10 µm.

Figure 3. CFTR correctors partially rescue the maturation and the canalicular localization of other ER-retained ABCB4 variants. (A-C) ABCB4-I490T (A), R545H (B) or L556R (C) were transiently expressed in HEK cells. After treatment with 10  $\mu$ M of the indicated CFTR correctors, the maturation of the missense variants was assessed by immunobloting as in Figure 2A. Arrows indicate mature ABCB4. Molecular weight markers (in kDa) are also indicated. These panels are representative of at least five independent experiments for each condition. (D-F) Densitometry analyses of A-C, as performed in Figure 2B. Means ( $\pm$  SD) of at least five independent experiments per condition are represented. (G-I) After treatment with vehicle (control) or 10  $\mu$ M of the indicated CFTR correctors, HepG2 cells expressing ABCB4-I490T (G), ABCB4-R545H (H) or ABCB4-L556R (I) were processed for indirect immunofluorescence and confocal microscopy, as described in Figure 2C. Dashed squares indicate magnification shown on the right of each merged panel. Asterisks indicate bile canaliculi. Each panel is representative of at least three independent experiments for each condition. Bars:  $10 \mu m$ .

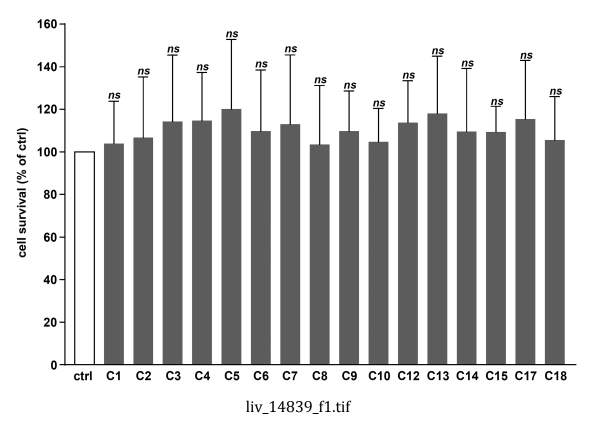
Figure 4. CFTR correctors do not rescue the activity of ER-retained ABCB4 variants and inhibit ABCB4-WT activity. HEK cells expressing ABCB4-I541F (A) ABCB4-I490T (B), ABCB4-R545H (C), ABCB4-L556R (D) or ABCB4-WT (E) were treated with vehicle (ctrl, control) or 10  $\mu$ M of the indicated CFTR correctors, and ABCB4-mediated PC secretion was measured and represented as a percentage of the activity for control vehicle-treated cells expressing ABCB4-WT after background subtraction. Means (± SD) of at least three independent experiments performed in triplicate for each tested condition are shown.

**Figure 5.** *In silico* molecular docking of CFTR correctors suggest their direct interaction with ABCB4. (A) In silico molecular docking of the six color-coded CFTR correctors was performed with ABCB4 in inward-facing (ABCB4<sup>if</sup>) and closed conformation (ABCB4<sup>cc</sup>) models (see *Materials and Methods* for details). For ABCB4<sup>if</sup>, a single site of potential interaction was identified in the protein chamber while three sites were identified for ABCB4<sup>cc</sup> in an alternative site and the two ATP-binding sites. For each CFTR corrector and for each site, the 50 poses with the highest affinities from 20 independent simulations are shown. (B) For each of the six CFTR correctors of interest, the contact rate of ABCB4 residues was calculated for the four sites of interest shown in A (see *Materials and Methods* for details). For the sake of clarity, only residues for which the average contact rates (± SD) were higher than the thresholds indicated by the dashed lines are shown.

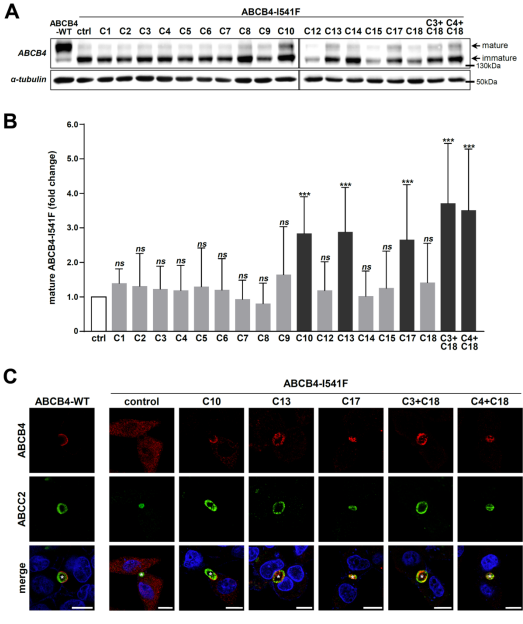
**Figure 6.** *In silico* molecular docking of CFTR correctors reveals putative interaction residues and domains in ABCB4. (A) ABCB4 residues indicated in Figure 5B with dark grey bars for which the contact rate is greater than 50% (ABCB4<sup>if</sup> - Protein chamber) or 90% (ABCB4<sup>cc</sup>) are represented with their lateral chains in the 3D structures of ABCB4<sup>if</sup> and ABCB4<sup>cc</sup>. Each transmembrane domain and nucleotide binding domain are represented in different colors, as well as the coupling helices (see Supplementary Table S6). Underlined residues are further discussed in the Discussion section. (B) ABCB4 transmembrane domains (TM), nucleotide

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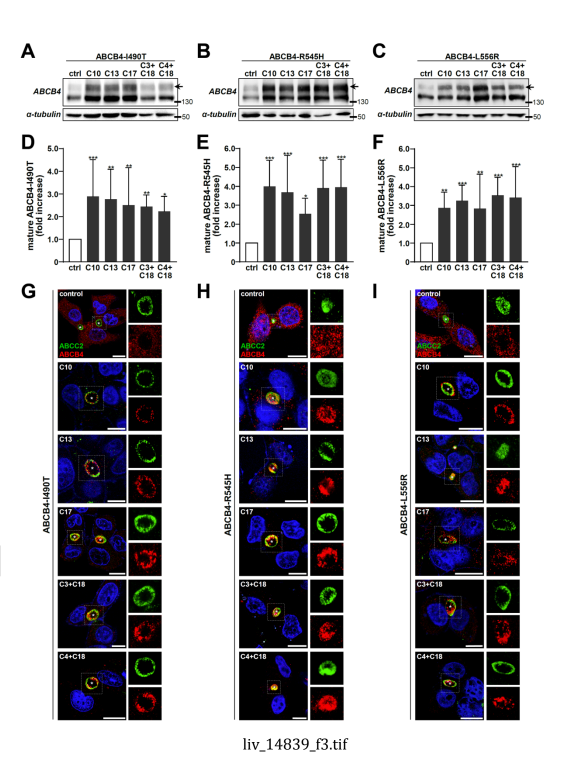
binding domains (NBD) and coupling helices (CH), for which potential interaction with CFTR correctors is suggested in Figures 5B and 6A, are highlighted with the indicated color code.

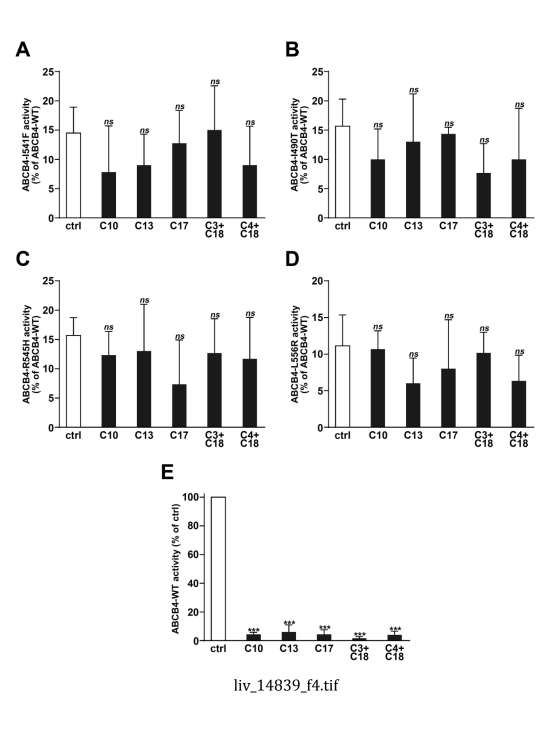


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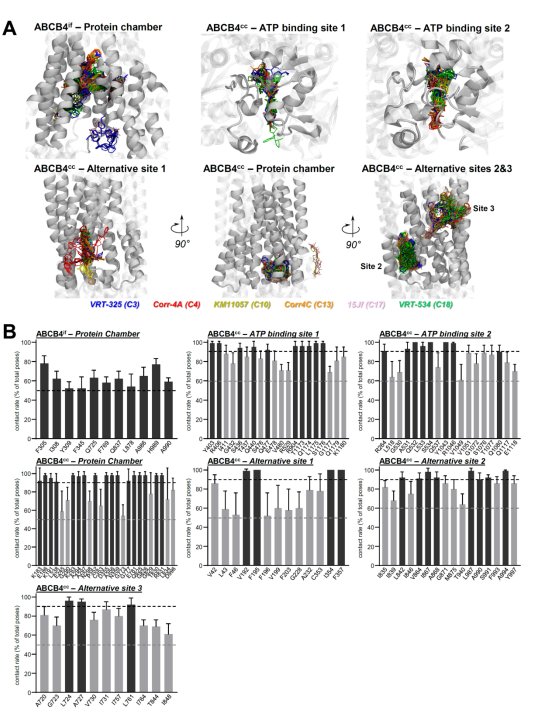


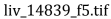
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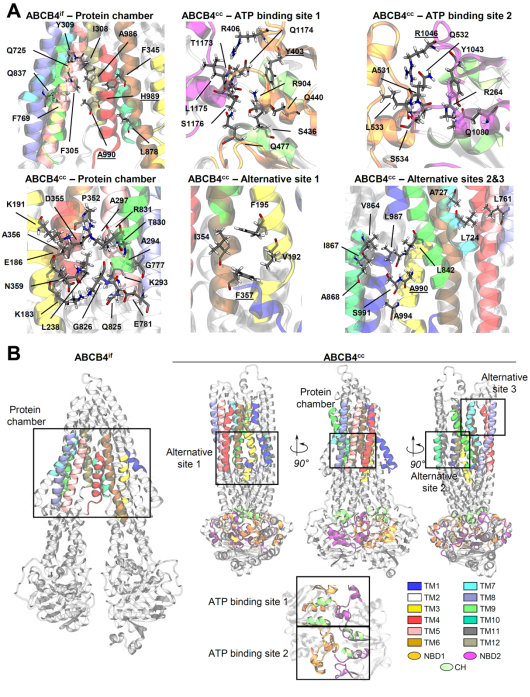




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### **SUPPLEMENTARY INFORMATION**

## Effect of CFTR correctors on the traffic and the function of intracellularly retained ABCB4 variants

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#### SUPPLEMENTARY MATERIALS AND METHODS

**ABCB4 model preparation.** Prior to docking calculations, both models were embedded in water-solvated (3:1) POPC:Cholesterol membrane lipid bilayers using the CHARMM-GUI webserver<sup>1-4</sup>. They were then minimized using the CPU version of PMEMD available in the AMBER18 package, using FF14SB, lipid17 and TIP3P forcefields for protein, lipid and water molecules, respectively<sup>5-8</sup>. Water, ions and lipids were then removed for docking calculations.

**Ligands.** The whole set of 16 CFTR correctors were initially considered for docking calculations, even though special attention was paid to VRT-325 (C3), Corr-4A (C4), KM11057 (C10), Corr-4C (C13), 15Jf (C17) and VRT-534 (C18). Correctors were downloaded from Pubchem (https://pubchem.ncbi.nlm.nih.gov/)<sup>9</sup> except for C7 which was manually built using the Maestro-Schrödinger Suite 2020-1 software (LLC, New York, NY, 2020). Corrector structures were first minimized at the quantum mechanical level using Density Functional Theory (DFT) M06-2X/6-31+G(d,p) level of theory<sup>10</sup>. Frequency analysis was carried out at the same level of theory to at least ensure a local minimum conformation by checking the absence of imaginary frequency. All relevant dihedral angles were allowed to be flexible during docking calculations (see Supplementary Table S1).

Analysis of non-covalent interactions. Considering the whole set of molecular poses, systematic contact (below 4.5 Å), H-bond and  $\pi$ -stacking interaction analyses were performed. H-bond interactions were considered, assuming distance- and angle-cutoffs set up to 3.5 Å and 120°, respectively. Two aromatic rings were considered  $\pi$ -stacked if: i) the distance between both rings' centers of mass is below 5.0 Å; ii) the angle between the two normal vectors to aromatic rings is below 10°. Analyses were carried out and rendered using CPPTRAJ<sup>11</sup>, PyTRAJ19<sup>12</sup> as well as homemade Python3 scripts (available on request) using Numpy20-2<sup>13,14</sup> and Matplotlib22<sup>15</sup> packages. 3D and 2D figures were rendered using the VMD 1.9.3<sup>16</sup> and CS ChemDraw softwares, respectively.

Compound	IUPAC name	Alternative name	Chemical Structure	PubChem CID	MW (Da)	References
C1	6-(1 <i>H</i> -Benzoimidazol-2- ylsulfanylmethyl)-2-(6-methoxy-4- methyl-quinazolin-2-ylamino)- pyrimidin-4-ol	Corr-3a		<u>135486495</u>	445.5	17
C2	2-(1-{4-[(4-Chlorophenyl)sulfonyl]- 1-piperazinyl}ethyl)-4-(1- piperidinyl)quinazoline	VRT-640		<u>23632303</u>	500.1	18
C3	4-(Cyclohexyloxy)-2-(1-(4-[(4- methoxybenzene)sulfonyl]piperazin- 1-yl)ethyl)quinazoline	VRT-325	O O O O O O O O O O O O O O O O O O O	<u>11957831</u>	510.6	19
C4	N-(2-(5-Chloro-2- methoxyphenylamino)-4'-methyl- 4,5'-bithiazol-2'-yl)benzamide	Corr-4a		<u>1144671</u>	457.0	17
C5	4,5,7-trimethyl-N-phenylquinolin-2- amine	Corr-5a	Me Me Me Ne	<u>740918</u>	262.3	17
C6	N-(4-bromophenyl)-4- methylquinolin-2-amine	Corr-5c	Me N N H H H	<u>768745</u>	313.2	17
C7	2-(4-isopropoxypicolinoyl)-N-(4- pentylphenyl)-1,2,3,4- tetrahydroisoquinoline-3- carboxamide	Compound 48		not applicable	485.6	20
C8	N-(2-fluorophenyl)-2-(1 <i>H</i> -indol-3- yl)-2-oxoacetamide	not applicable	HN-C	<u>2986288</u>	282.3	18
С9	7-Chloro-4-[4-(4- chlorophenyl)sulfonylpiperazin-1- yl]quinoline	KM11060		<u>1241327</u>	422.3	21
C10	7-Chloro-4-[4- (phenylsulfonyl)piperazin-1- yl]quinoline	KM11057		<u>1241326</u>	387.9	21
C12	N-(4-Fluorophenyl)-4-(4- methylphenyl)-1,3-thiazol-2-amine	Corr-2i	F HS HS Me	<u>625111</u>	284.4	17
C13	N-[2-(3-Acetyl-phenylamino)-4'- methyl-[4,5']bithiazolyl-2'-yl]- benzamide	Corr-4c	Me N H Me	<u>2230879</u>	434.5	17
C14	N-[2-(2-Methoxy-phenylamino)-4'- methyl-[4,5']bithiazolyl-2'-yl]- benzamide	Corr-4d	N S N OMe	<u>1117191</u>	422.5	17
C15	Phenyl-[4-(4-vinyl-phenyl)-thiazol- 2-yl]-amine	Corr-2b	S NH	<u>683025</u>	278.4	17
C17	N-[5-[2-(5-Chloro-2- methoxyanilino)-1,3-thiazol-4-yl]-4- methyl-1,3-thiazol-2-yl]-2,2- dimethylpropanamide	Compound 15Jf		<u>11958611</u>	437.0	22
C18	1-(1,3-Benzodioxol-5-yl)-N-[5-[(S)- (2-chlorophenyl)-[(2R)-2- hydroxypyrrolidin-1-yl]methyl]-1,3- thiazol-2-yl]cyclopropane-1- carboxamide	VRT-534		<u>86621212</u>	498.0	18

# Supplementary Table S1. Chemical structures of CFTR correctors used in this study

bold green bounds indicate allowed rotations during molecular docking calculations

Supplementary Table S2. Technical parameters of blind and refined molecular docking calculations.

Storestand	Madallad a service a	Decking colordation	Valence and an a	E	Box cente	er		Box size				
Structure	Structure Modelled sequence	Docking calculation	Volume search space	x	у	Z	x	у	Z			
ABCB4 <sup>if</sup>	42-629	Blind	Overall	61.6	60.6	59.4	80.0	65.0	120.0			
	692-1251	Refined	Protein chamber A	60.0	62.7	94.4	29.5	29.5	29.5			
			Protein chamber B	69.0	53.0	83.0	26.0	20.0	30.0			
ABCB4 <sup>cc</sup>	42-629	Blind	Overall	60.5	62.1	67.2	80.0	65.0	120.0			
	692-1251	Refined	Protein chamber	68.0	73.0	79.0	20.0	20.0	20.0			
			Alternative site 1	51.1	72.9	80.0	23.0	23.0	23.0			
			Alternative site 2	66.0	52.0	84.0	20.0	20.0	20.0			
			Alternative site 3	68.0	69.0	100.0	26.0	28.0	20.0			
			ATP-binding site 1	52.7	70.3	31.5	23.0	28.0	23.0			
			ATP-binding site 2	65.2	49.3	30.7	23.0	28.0	23.0			

## (A) Box parameter coordinates (center and size)

reported coordinates are defined according to PDB orientations available on request

# (B) Flexible residues in each volume search space.

Structure	Docking calculation	Volume search space	Flexible residues
ABCB4 <sup>if</sup>	Blind	Overall	None
	Refined	Protein chamber A	234, 305, 309, 311, 312, 337, 338, 339, 345, 346, 349, 725, 728, 729, 841, 870, 874, 944, 952, 978, 982, 989
		Protein chamber B	191, 195, 196, 201, 224, 236, 239, 242, 243, 345, 346, 349, 355, 357, 359
ABCB4 <sup>cc</sup>	Blind	Overall	None
	Refined	Protein chamber	234, 296, 298, 300, 349, 355, 766, 767, 768, 769, 770, 772, 774, 778, 830, 837
		Alternative site 1	44, 46, 195, 196, 239, 357
		Alternative site 2	838, 870, 874, 937, 940, 941, 944, 945, 991, 992, 993
		Alternative site 3	309, 311, 312, 316, 338, 345, 725, 728, 768, 769, 770, 772, 774, 778, 830, 837
		ATP-binding site 1	166, 402, 403, 405, 435, 436, 437, 438, 440, 476, 477, 478, 557, 558, 902, 905, 908, 1156, 1166, 1173, 1174, 1176, 1179, 1180
		ATP-binding site 2	513, 514, 517, 524, 532, 534, 537, 538, 1042, 1043, 1045, 1053, 1071, 1076, 1077, 1080, 1116, 1117, 1118, 1199, 1200

CFTR ABCB4 <sup>if</sup>				ABCB4 <sup>cc</sup>						
Correctors	Protein chamber A	Out of zone	Alternative site 1	ATP-binding site 1	ATP-binding site 2	Out of zone				
C1	0.98	0.02	0.02	0.28	0.46	0.24				
C2	0.89	0.11	0.56	0.05	0.16	0.23				
C3	0.90	0.10	0.52	0.07	0.18	0.24				
C4	1.00	0.01	0.29	0.16	0.50	0.06				
C5	0.94	0.07	0.25	0.18	0.40	0.17				
C6	0.91	0.10	0.33	0.18	0.32	0.17				
C7	1.00	0.00	0.43	0.06	0.11	0.40				
C8	0.95	0.05	0.04	0.20	0.34	0.42				
С9	0.92	0.08	0.26	0.12	0.44	0.18				
C10	0.90	0.10	0.35	0.12	0.41	0.13				
C12	0.95	0.05	0.32	0.21	0.41	0.07				
C13	0.99	0.01	0.24	0.24	0.52	0.01				
C14	0.96	0.04	0.26	0.16	0.53	0.05				
C15	0.97	0.03	0.36	0.17	0.37	0.11				
C17	0.98	0.02	0.17	0.16	0.46	0.22				
C18	0.92	0.08	0.44	0.04	0.16	0.36				
Average (Overall)	$0.95\pm0.04$	$\begin{array}{c} 0.05 \pm \\ 0.04 \end{array}$	$0.30\pm0.15$	$0.15\pm0.07$	$0.36\pm0.14$	$0.19\pm0.12$				
Average (Selected)	$0.95\pm0.05$	$\begin{array}{c} 0.05 \pm \\ 0.04 \end{array}$	$0.34\pm0.13$	$0.13\pm0.07$	$0.37\pm0.16$	$0.17\pm0.13$				

**Supplementary Table S3.** Spatial distributions of the whole set of CFTR correctors over defined binding sites and out of these zones originally obtained from blind molecular docking calculations.

averages  $\pm$  SD are indicated for all correctors as well as for selected correctors reported in bold

### Ben Saad et al., Supplementary information

**Supplementary Table S4.** Minimum and maximum affinity energies ( $\Delta G_{min}$  and  $\Delta G_{max}$ , respectively; in kcal.mol<sup>-1</sup>) obtained from refined molecular docking calculations on ABCB4<sup>if</sup> (A) and ABCB4<sup>cc</sup> (B), as well as minimum docking affinity mean signed deviation (MSD, kcal.mol<sup>-1</sup>) with respect to Protein chamber A and ATP-binding site 1 for ABCB4<sup>if</sup> and ABCB4<sup>cc</sup>, respectively.

	P	rotein chamber A		Protein chamber B				
	$\Delta G_{min}$	$\Delta G_{max}$	#	$\Delta G_{min}$	$\Delta G_{max}$	#		
C1	-10.7	-9.1	399	-10.0	-6.5	263		
C2	-10.3	-8.9	400	-10.5	1.4	135		
С3	-10.3	-8.9	400	-10.4	-5.2	177		
C4	-9.6	-8.2	398	-9.6	-6.7	177		
C5	-9.6	-7.3	379	-8.7	-0.1	330		
C6	-9.3	-6.9	398	-8.0	-5.7	343		
C7	-12.2	-8.5	278	-9.3	-1.2	254		
C8	-9.5	-7.2	383	-8.0	-6.1	400		
С9	-9.9	-7.8	397	-9.7	-6.1	131		
C10	-9.7	-7.6	400	-9.3	-5.3	219		
C12	-8.9	-7.3	400	-8.1	-3.8	362		
C13	-10.3	2.3	399	-10.2	3.5	158		
C14	-9.8	-7.9	398	-9.5	1.7	180		
C15	-8.8	-7.1	397	-8.0	-5.6	375		
C17	-8.9	-7.3	399	-8.1	-5.9	377		
C18	-10.5	-8.8	400	-10.1	-4.8	300		
MSD		-			$0.7\pm0.8$			

(A)

# indicates the selected number of poses for each molecule, with a maximum 2 kcal.mol<sup>-1</sup> range with respect to the top-ranked pose for each ABCB4 state/CFTR corrector/region triad. Selected CFTR correctors are shown in bold.

	P	rotein chamb	er	Alter	native site	1	Alter	native site	2	Alter	native site	3	ATP	binding si	te 1	ATP	binding si	te 2
	$\Delta G_{min}$	$\Delta G_{max}$	#	$\Delta G_{min}$	$\Delta G_{\text{max}}$	#	$\Delta G_{min}$	$\Delta G_{max}$	#									
C1	-9.2	5.9	69	-9.7	-8.1	399	-9.8	-0.5	223	-7.1	-5.9	400	-11.6	-9.0	365	-11.1	-8.8	385
C2	-5.2	8.1	61	-10.2	-8.9	393	-9.6	3.9	321	-7.1	-6.0	400	-11.4	-7.5	175	-12.4	-8.7	138
С3	-7.4	8.8	46	-10.4	-8.5	400	-9.3	0.7	386	-7.2	-6.4	400	-12.1	-7.8	111	-11.9	-8.8	219
C4	-6.7	7.4	112	-8.8	-5.5	396	-8.3	6.6	334	-6.9	-0.4	398	-9.9	-7.4	372	-10.3	-7.4	327
C5	-8.7	6.7	111	-10.0	-7.1	207	-7.5	7.1	363	-6.3	-5.0	400	-9.3	-6.2	358	-9.5	-6.1	259
C6	-7.5	7.7	91	-9.2	-6.6	280	-7.2	5.3	329	-6.1	-4.1	399	-9.0	-6.3	307	-9.0	-5.5	342
C7	-8.1	3.6	57	-9.7	-7.8	397	-8.1	5.7	389	-7.1	-5.9	400	-9.9	-3.1	111	-11.6	-5.8	249
C8	-8.7	5.9	213	-8.5	-6.7	398	-7.4	2.2	388	-6.0	-2.8	398	-9.3	-7.5	397	-9.6	-7.0	327
С9	-7.0	8.1	36	-9.0	-7.3	400	-8.1	4.3	258	-6.4	-5.6	400	-10.2	-5.9	248	-10.7	-7.6	289
C10	-7.7	8.7	33	-9.3	-7.0	352	-7.9	4.3	255	-6.5	-5.2	399	-9.8	-6.8	267	-10.5	-7.5	310
C12	-7.6	7.8	96	-9.1	-7.2	398	-8.0	6.6	193	-6.1	-4.3	399	-8.7	-6.7	388	-9.2	-6.3	315
C13	-7.9	7.0	104	-10.1	-8.6	400	-8.7	2.6	327	-7.3	-6.2	400	-10.2	-7.4	359	-10.7	-3.5	304
C14	-7.9	5.6	132	-9.1	-4.2	397	-8.3	1.5	277	-6.6	-5.6	399	-9.9	-7.1	325	-10.0	-6.6	324
C15	-7.4	8.2	106	-8.6	-6.6	391	-7.6	7.1	252	-6.0	-4.3	400	-8.4	-6.3	386	-9.3	-5.6	255
C17	-5.4	8.8	127	-8.5	-6.5	392	-7.4	3.8	385	-6.5	-5.2	400	-9.2	-6.6	327	-9.4	-6.8	368
C18	-7.5	6.7	119	-9.8	-7.3	264	-9.3	1.2	380	-6.4	-5.6	400	-11.0	-2.0	336	-10.1	-7.4	192
MSD		$2.5\pm1.5$		0	$.6 \pm 0.8$		1	$.7 \pm 0.5$	•	3	$0.4\pm0.8$			-			$-0.3 \pm 0.6$	

# indicates the selected number of poses for each molecule, with a maximum 2 kcal.mol<sup>-1</sup> range with respect to the top-ranked pose for each ABCB4 state/CFTR corrector/region triad. Selected CFTR correctors are shown in bold

Conformations	Binding sites	H-bondi	ng interactions	van der W	aals interactions
		Residue	Involvement rate	Residue	Involvement rate
ABCB4 <sup>if</sup>	Protein chamber A	Gln725	0.21	Phe338	0.24
		Trp234	0.08	Phe776	0.21
		Tyr312	0.06	Phe982	0.18
		Tyr309	0.05	Phe993	0.14
		Gln772	0.05		
	Protein Chamber B	Ser242	0.06	Phe345	0.81
		Lys191	0.05	Phe357	0.19
ABCB4 <sup>cc</sup>	Protein chamber	Thr830	0.15		
	Alternative site 1	Arg361	0.04	Phe357	0.54
				Phe195	0.46
	Alternative site 2	Ser991	0.06		
	Alternative site 3			Phe760	0.46
	ATP-binding site 1	Arg904	0.27	Tyr403	1.00
		Arg529	0.1		
		Gln440	0.09		
		Tyr403	0.09		
		Gln1179	0.07		
		Arg406	0.06		
		Ser476	0.05		
	ATP-binding site 2	Arg1046	0.21	Tyr1043	1.00
		Gln1080	0.21		
		Gln537	0.13		
		Arg264	0.06		

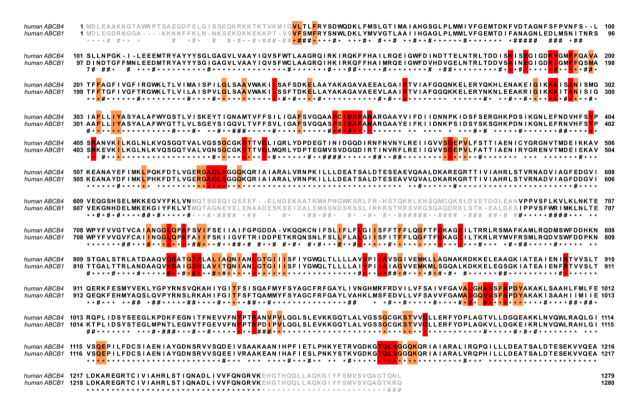
**Supplementary Table S5.** Description of non-covalent interactions of CFTR correctors with ABCB4<sup>if</sup> and ABCB4<sup>cc</sup>.

**Supplementary Table S6.** Sequence-based definitions of ABCB4<sup>if</sup> (A) and ABCB4<sup>cc</sup> (B) binding sites from molecular docking calculations. Locations according to transmembrane helices (TM), nucleotide binding domains (NBD) and coupling helices (CH) are reported.

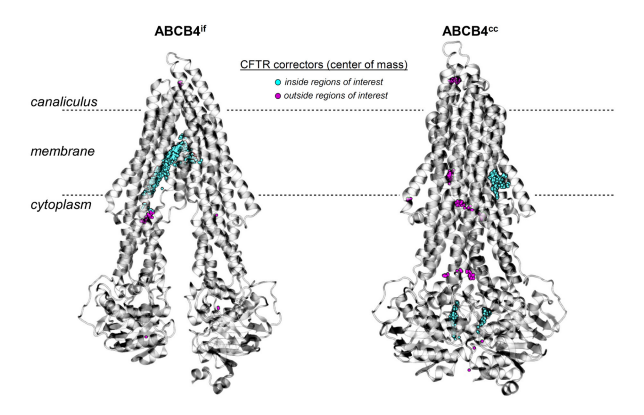
(A)

	Protein Chamber A	Protein Chamber B	Color code in Figure 6A-B
TM1	-	63-70	
TM2	-	137-143	
TM3	-	183-203	
TM4	227-240	223-243	
TM5	291-314	307-309	
TM6	336-353	341-362	
NBD1	-	-	
TM7	717-731	-	
TM8	756-780	-	
TM9	828-846	-	
TM10	867-878	877-880	
TM11	-	-	
TM12	975-993	-	
NBD2	-	-	
CH2-3	-	-	
CH4-5	-	-	
CH8-9	-	-	
CH10-11	-	-	

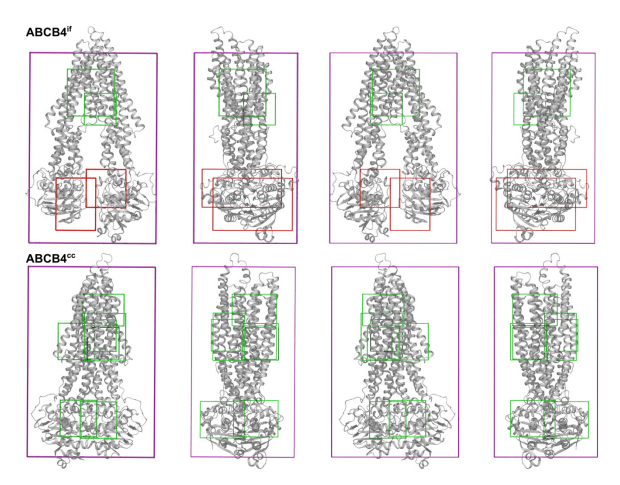
	Protein	Alternative	Alternative	Alternative	ATP-binding	ATP-binding	Color code
	Chamber	site 1	site 2	site 3	site 1	site 2	in Figure 6A-B
TM1	-	42-65	-	71-79	-	-	
TM2	-	-	-	-	-	-	
TM3	-	185-200	-	-	-	-	
TM4	230-239	225-241	-	216-225	-	-	
TM5	291-305		-	305-320	-	-	
TM6	347-360	346-361	-	334-347	-	-	
NBD1	-	-	-	-	401-415, 431-446 474-481, 555-561	479-485, 510-539, 561-563	
TM7	714-719	-	-	720-734	-	-	
TM8	766-781	-	-	752-766	-	-	
TM9	828-838	-	835-844	840-853	-	-	
TM10	-	-	864-877	-	-	-	
TM11	-	-	937-947	-	-	-	
TM12	-	-	984-996	973-986	-	-	
NBD2	-	-	-	-	1118-1123, 1155-1181, 1203-1204	1041-1053, 1071-1085, 1115-1118, 1197-1203, 1247-1248	
CH2-3	-	-	-	-	161-167	-	
CH4-5	-	-	-	-	-	262-268	
CH8-9	-	-	-	-	-	799-804	
CH10-11	-	-	-	-	901-909		



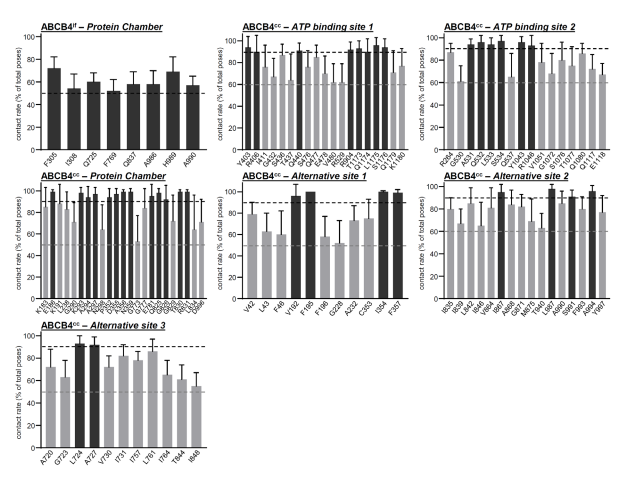
**Supplementary Figure S1. Sequence alignment of human ABCB4 and human ABCB1.** The protein sequences of human ABCB4 (NP\_000434.1) and ABCB1 (NP\_000918.2) were aligned using the Uniprot alignment tool (https://www.uniprot.org/align/) and represented using Jalview software (https://www.jalview.org/). Conservation of residues are represented as follows: \*, identical; # similar. Residues shown in Figure 5B are highlighted either in red for contact rates above 90% or in orange otherwise. Residues in grey (N- and C-terminal domains and L<sub>1</sub>-linker) were not built in the 3D models.



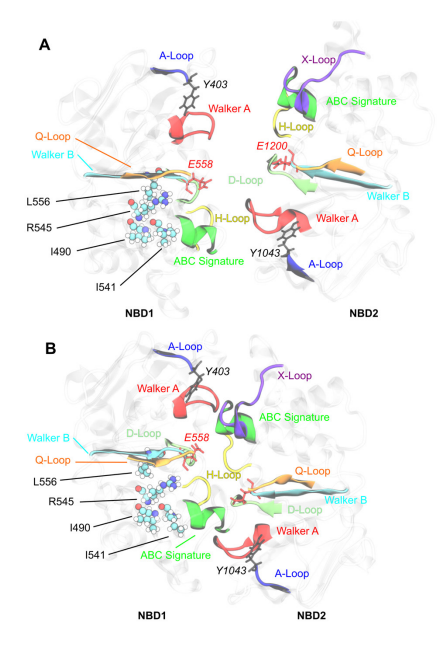
**Supplementary Figure S2. Blind** *in silico* molecular docking of CFTR correctors into ABCB4<sup>if</sup> and ABCB4<sup>cc</sup>. Blind molecular docking simulations with CFTR correctors shown in Supplementary Table S1 were performed using the two indicated conformations of ABCB4 (see *Materials and Methods* for details). Four hundred poses from 20 independent simulations are displayed for each corrector. For the sake of clarity, only the corrector centers of mass are represented and color-coded according to their localization inside or outside the regions defined in Supplementary Table S2. Results from these analyses are summarized in Supplementary Table S3.



**Supplementary Figure S3. Refined** *in silico* molecular docking of CFTR correctors into ABCB4<sup>if</sup> and ABCB4<sup>cc</sup>. Refined molecular docking simulations with CFTR correctors shown in Supplementary Table S1 were performed using the two indicated conformations of ABCB4 (see *Materials and Methods* for details), for which regions were originally obtained from blind molecular docking calculations (boxes colored in purple). Regions obtained from ABCB4<sup>if</sup> were also sampled in ABCB4<sup>cc</sup> and *vice versa*. Regions colored in red are regions which were sampled without exhibiting poses from refined molecular docking calculations while regions colored in green were further considered.



Supplementary Figure S4. Contact rates of ABCB4 residues with all CFTR correctors. The contact rates of ABCB4 residues identified in Figure 5B with all correctors shown in Supplementary Table S1 are represented. Dash lines indicate the cut-offs chosen for considering the involvement of the residues. Means ( $\pm$  SD) of 20 independent simulations per CFTR corrector are shown.



Supplementary Figure S5. Structural mapping of the I490, I541, R545 and L556 on ABCB4<sup>if</sup> (A) and ABCB4<sup>cc</sup> (B). Location of the four ABCB4 variants investigated in the present study (van der Waals representation, atom-type coloring), focusing on the NBDs. Key structural features of NBD are colored, as well as the catalytic glutamates (Glu558 and Glu1200, licorice representation, red) and adenosine-binding tyrosines (Tyr403 and Tyr1043, licorice representation, black).

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