Lipophilicity of the Cystic Fibrosis Drug, Ivacaftor (VX-770), and Its Destabilizing Effect on the Major CF-causing Mutation: F508del

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Received February 14, 2018; accepted June 13, 2018

**ABSTRACT**

Deletion of phenylalanine at position 508 (F508del) in cystic fibrosis transmembrane conductance regulator (CFTR) is the most common cystic fibrosis (CF)-causing mutation. Recently, ORKAMBI, a combination therapy that includes a corrector of the processing defect of F508del-CFTR (lumacaftor or VX-809) and a potentiator of channel activity (ivacaftor or VX-770), was approved for CF patients homozygous for this mutation. However, clinical studies revealed that the effect of ORKAMBI on lung function is modest and it was proposed that this modest effect relates to a negative impact of VX-770 on the stability of F508del-CFTR. In the current studies, we showed that this negative effect of VX-770 at 10 μM correlated with its inhibitory effect on VX-809-mediated correction of the interface between the second membrane spanning domain and the first nucleotide binding domain bearing F508del. Interestingly, we found that VX-770 exerted a similar negative effect on the stability of other membrane localized solute carriers (SLC26A3, SLC26A9, and SLC6A14), suggesting that this negative effect is not specific for F508del-CFTR. We determined that the relative destabilizing effect of a panel of VX-770 derivatives on F508del-CFTR correlated with their predicted lipophilicity. Polarized total internal reflection fluorescence microscopy on a supported lipid bilayer model shows that VX-770, and not its less lipophilic derivative, increased the fluidity of and reorganized the membrane. In summary, our findings show that there is a potential for nonspecific effects of VX-770 on the lipid bilayer and suggest that this effect may account for its destabilizing effect on VX-809 rescued F508del-CFTR.

**Introduction**

Cystic fibrosis (CF) is caused by mutations of a unique ATP-binding cassette anion channel, cystic fibrosis transmembrane conductance regulator (CFTR) (Cutting, 2015). CFTR has a lasso motif, two nucleotide binding domains (NBDs), two membrane spanning domains (MSDs), and a regulatory domain. In addition, the interface between the NBDs and MSDs consists of alpha-helical intracellular loops (ICLs) and coupling helices. The ICL:NBD1 interface is disrupted by the major CF-causing mutation, deletion of phenylalanine at position 508 (F508del), thereby compromising assembly and function of CFTR (Mendoza et al., 2012; Rabeh et al., 2012).

The first US Food and Drug Administration-approved medicine to treat the underlying cause of CF is KALYDECO (ivacaftor or VX-770), a potentiator that enhances channel gating of wild-type (wt) CFTR and CFTR with certain gating mutations (i.e., glycine to aspartic acid at position 551 or G551D) at the cell surface by directly binding to the protein to prolong its open state in an ATP-independent manner (Van Goor et al., 2009; Eckford et al., 2012; Jih and Hwang, 2013). On the other hand, patch clamp studies by Jih and Hwang (2013) suggested that both ATP-dependent and ATP-independent CFTR channel gating are modified by VX-770. Recently, ORKAMBI, a combination of VX-770 and a corrector compound known as lumacaftor (VX-809), was approved for use in patients who are homozygous for the major CF mutation, F508del. The corrector compound, VX-809, acts to partially rescue the defect in protein mistrafficking and misprocessing exhibited by F508del-CFTR.

This work was supported by Cystic Fibrosis Canada; Canadian Institutes of Health Research [Grants MOP-97854, GPG-102171]; and Natural Sciences and Engineering Research Council [Grant RGPIN-2015-043].

https://doi.org/10.1124/mol.118.112177

This article has supplemental material available at molpharm.aspetjournals.org.

**ABBREVIATIONS:** BMOE, bismaleimidodeithane; bSM, egg sphingomyelin; CF, cystic fibrosis; CFTR, cystic fibrosis transmembrane conductance regulator; DMEM, Dulbecco’s modified Eagle’s medium; DOPC, dioleoylphosphatidylcholine; Endo, endoglycosidase; F508del, deletion of phenylalanine at position 508; FBS, fetal bovine serum; Fsk, forskolin; HEK, human embryonic kidney; ICL, intracellular loop; logP, partition coefficient; MSD, membrane spanning domain; NBD, nucleotide binding domain; NEAA, nonessential amino acids; pTIRF, polarized total internal reflection fluorescence; SLC, solute carrier; wt, wild type.
to the cell surface (Van Goor et al., 2011) by repairing the aberrant ICL:NBD1 interfaces (Rabeh et al., 2012; Loo et al., 2013; Okiyone et al., 2013; Ren et al., 2013). VX-770 acts on the VX-809-corrected F508del-CFTR to augment its activity (Van Goor et al., 2009, 2011; Eckford et al., 2012; Jih and Hwang, 2013).

Unfortunately, ORKAMBI mediates modest and variable clinical responses (Boyle et al., 2014), prompting further discovery efforts for more efficacious corrector compounds. Certain in vitro findings suggest that the modest in vivo effect may reflect an adverse effect of VX-770 on the stability of the VX-809-corrected protein (Cholon et al., 2014; Veit et al., 2014; Matthes et al., 2016). Thus, the aim of the current studies was to interrogate the mechanism underlying this destabilizing effect of VX-770 on the major CF mutant, F508del. We also assessed the specificity of this destabilizing effect of VX-770 by studying its interaction with other membrane proteins (including SLC26A3, SLC26A9, and SLC6A14) as well as the lipid bilayer.

Materials and Methods

Chronic Treatment of F508del-CFTR with VX-770 and Its Analogs. Human embryonic kidney (HEK)-293 cells stably expressing F508del-CFTR (HEK F508del) were plated on 24-well plates (Sarstedt, Nümbrecht, Germany) and maintained in Dulbecco’s Modified Eagle’s Medium (DMEM; Wisent, St-Bruno, Quebec, Canada) supplemented with 10% Fetal Bovine Serum (FBS; Wisent), 1% nonessential amino acids (NEAA; Wisent), 0.6 mg/ml G418 sulfate (Wisent), and 5 μg/ml blasticidin (Wisent) at 37°C with 5% CO₂ overnight. For the initial chronic VX-770 studies, HEK F508del cells were pretreated with VX-770 at a pharmacological concentration (0.1 μM) or at a suprapharmacological concentration (10 μM) with 3 μM VX-809 (Selleck Chemicals, Houston, TX) in the previously described media at 37°C with 5% CO₂ for 48 hours. For the studies with VX-770 and its derivatives, HEK F508del cells were pretreated with VX-770 (Selleck Chemicals) or its various derivatives (SE and KT compounds) synthesized in Dr. Russell Viirre’s laboratory (Toronto, Ontario, Canada) (Krimo Toutah, Maurita Hung, Salma Elmallah, C. Michael Hamilton, Robert N. Young, Christine E. Bear and Russell M) or at a suprapharmacological concentration (10 μM) with 3 μM VX-809 (Selleck Chemicals, Houston, TX) in the previously described media at 37°C with 5% CO₂ for 48 hours. For the studies with VX-770 and its derivatives, HEK F508del cells were pretreated with VX-770 (Selleck Chemicals) or its various derivatives (SE and KT compounds) synthesized in Dr. Russell Viirre’s laboratory (Toronto, Ontario, Canada) (Krimo Toutah, Maurita Hung, Salma Elmallah, C. Michael Hamilton, Robert N. Young, Christine E. Bear and Russell Viirre, manuscript in preparation, refer to Supplemental Fig. 2 for chemical structures of VX-770 derivatives) at a pharmacological concentration (0.1 μM) or at a suprapharmacological concentration (10 μM) in the previously described media at 27°C with 5% CO₂ for 24 hours. The synthesis and characterization of the KT compounds employed the same methodology as (http://digital.library.ryerson.ca/islandora/object/RULA:1478) for the most part but also incorporated additional reaction steps previously published by the Verkman group for modification of the left-hand side of the VX-770 molecule (Pedemonte et al., 2005).

Generation of Mutants and Constructs. The V510C, A1067C, and V510C/A1067C cysteine-less variants of CFTR were kindly provided by Dr. David Clarke (Toronto, Ontario, Canada); and the SLC6A14 construct with a C-terminal FLAG tag was generated by OriGene (Rockville, MD). The F508del mutation was generated on the V510C/A1067C cysteine-less variant of wt-CFTR background with primers (forward: 5’-CATTAAAGAAAAATATTGTTGTTCCTATGATG-3’ and reverse: 5’-CATTAAAGATTTGGTTGTTCCTATGATG-3’) and the KAPA HiFi HotStart PCR Kit (KAPA Biosystems, Woburn, MA). Plasmid DNA was expanded with the QIAprep Spin Miniprep Kit (Qiagen, Hilden, Germany), and the sequence was confirmed with DNA sequencing (TCAG, Toronto, Ontario, Canada). HEK-293 GripTite (GT) cells, kindly provided by Dr. Daniella Rotin’s laboratory (Toronto, Ontario, Canada), were maintained in DMEM (Wisent) supplemented with 10% FBS (Wisent), 1% NEAA (Wisent), and 0.6 mg/ml G418 sulfate (Wisent), transfected with constructs using PolyFect Transfection Reagent (Qiagen) according to the manufacturer’s protocol and temperature-rescued at 27°C with 5% CO₂ for 24 hours.

Cysteine Cross-linking. HEK-293 GT cells were grown on 24-well plates (Sarstedt) at 37°C with 5% CO₂ overnight and then transfected with cys-less variants of CFTR. Cells were treated with DMSO (0.1%), VX-809 (3 μM), VX-770 (0.1 or 10 μM), or VX-809 plus VX-770 (0.1 or 10 μM) at 27°C with 5% CO₂ for 24 hours. The next day, cells were treated with 50 μM bismaleimidodithioene (BMOE; Life Technologies, Rockford, IL) in DMEM (Wisent) supplemented with 10% FBS (Wisent), 1% NEAA (Wisent), and 0.6 mg/ml G418 sulfate (Wisent) at 27°C with 5% CO₂ for 1 hour.

Deglycosylation Studies. Cys-less variants of CFTR were lysed from HEK-293 GT cells into radioimmunoprecipitation assay (50 mM Tris-Base, 150 mM sodium chloride, 1 mM EDTA, 1% (v/v) Triton X-100, 0.1% (v/v) SDS, 1× protease inhibitor cocktail (AMRESCO, Cleveland, OH), pH 7.4) buffer. Protein samples were treated with Endoglycosidase H and PNGase F according to the manufacturer’s protocol (New England Biolabs, Ipswich, MA).

Chronic Treatment of SLC Proteins with VX-770. HEK-293 GT cells were grown on 24-well plates (Sarstedt) at 37°C with 5% CO₂ overnight and transfected with the SLC constructs. Cells were then treated with various concentrations of VX-770 (0, 0.1, 1, 2, 10 μM) at 27°C with 5% CO₂ for 24 hours.

Immunoblotting. HEK F508del and HEK-293 GT cells were lysed into radioimmunoprecipitation buffer and protein samples were subjected to SDS-PAGE. CFTR protein samples were run on 6% Tris-glycine sodium dodecyl sulfate (TG SDS) gels (Life Technologies, Carlsbad, CA) and SLC protein samples were run on 8% TG SDS gels (Life Technologies). Protein samples were transferred to nitrocellulose paper and probed with the following primary antibodies overnight at 4°C shaking: IgG2b mAb596 antibody (University of North Carolina at Chapel Hill, Chapel Hill, NC, code: A4; Cystic Fibrosis Foundation Therapeutics Inc., Bethesda, MD) for CFTR protein (1.5000 for F508del-CFTR, 1.0000 for V510C/A1067C cys-less variant of wt-CFTR, 1.5000 for V510C/A1067C cys-less variant of F508del-CFTR), calnexin-specific rabbit pAb (1:10,000, cat. no.: C4731; Sigma-Aldrich, St. Louis, MO) for calnexin loading control, anti-mouse FLAG antibody (1:2500; Sigma-Aldrich) for SLC protein, and anti-β-actin (1:10,000, ab8229; Abcam, Cambridge, UK) for actin loading control. The samples were then probed with secondary antibodies for 1 hour at room temperature shaking at concentrations twice as diluted as the primary antibodies as follows: horseradish peroxidase-conjugated goat anti-mouse IgG secondary antibody (Pierce, Rockford, IL) for CFTR protein and actin loading control, horseradish peroxidase-conjugated goat anti-rabbit IgG secondary antibody (Pierce, Rockford, IL) for CFTR protein and actin loading control, horseradish peroxidase-conjugated goat anti-rabbit IgG secondary antibody (Pierce, Rockford, IL) for calnexin loading control, and anti-mouse HA (MMS-101R; Covance, Princeton, NJ) for SLC protein. Immunoblots were exposed using Amersham enhanced chemiluminescent reagent (GE Healthcare Life Sciences, Mississauga, Ontario, Canada) with the Li-Cor Odyssey Fc (LI-COR Biosciences, Lincoln, NE) at a linear range of exposure. Densitometry of immunoblot bands were analyzed with Image Studio Lite software (Version 5.2.5) from Li-Cor Odyssey Fc (LI-COR Biosciences).

FLiPR Membrane Potential Assay. The FLiPR membrane potential assay was conducted as previously described (Molinski et al., 2015). Briefly, HEK F508del cells were grown on black 96-well plates with clear bottoms (Corning, Corning, NY) at 37°C with 5% CO₂ overnight. Cells were then treated with 5 μM VX-809 at 37°C with 5% CO₂ for 48 hours. The next day, cells were loaded with 0.5 mg/ml FLiPR membrane potential dye (Molecular Devices, Sunnyvale, CA) in sodium gluconate buffer as previously described (Molinski et al., 2015) for 45 minutes with 5% CO₂ at 37°C. Fluorescence was read at excitation of 530 nm and emission of 560 nm on the SpectraMax i3X.
fluorescence plate reader at 37°C (Molecular Devices). Baseline fluorescence was read for 5 minutes. CFTR was stimulated by vehicle, 1 μM forskolin (Fsk; Sigma-Aldrich), 1 μM Fsk with VX-770 (10 μM), or 1 μM Fsk with the derivative SE-03 (10 μM) for 10 minutes. CFTR was then inhibited with 10 μM CFTR inhibitor-172 (CFTRinh-172; Cystic Fibrosis Foundation Therapeutics Inc. and Rosalind Franklin University of Medicine and Science, Chicago, IL) for 12 minutes. Curves were normalized to last point of baseline and the maximal points of stimulation were plotted with GraphPad Prism v6.01 (San Diego, CA).

Polarized Total Internal Reflection Fluorescence Microscopy. The formation of supported lipid bilayers of 1:1:1 dioleoylphosphatidylcholine/egg sphingomyelin/cholesterol (DOPC/SphCH/SM) and 1:1:1 DOPC/distearylphosphatidylcholine/cholesterol with DiLC2(3) (1 mol %; Life Technologies) in buffer (10 mM HEPES, 150 mM NaCl, pH 7.4) were prepared as previously described (Walsh et al., 2014). All images were acquired using a home-built pTIRF system incorporating four excitation laser lines built on an Olympus IX70 inverted microscope. All images were captured at 15-second intervals for 1 hour with a water-cooled Evolve 512 EMCCD camera (Photometrics, Tucson, AZ) controlled by µ-Manager (http://www.micro-manager.org). Fluorescent probes were excited by substrate parallel (Fp, s-polarized) and perpendicular (Fp, p-polarized) polarized excitation at 532 nm through a half-wave liquid crystal variable retarder (LCC25 1111A; Thorlabs, Newton, NJ). Two bilayer studies were performed with multiple fields of view examined for each bilayer (three to six fields). Order parameter S2 values were determined for each region of interest and calculated on a per pixel basis using an in-house macro. Refer to previous publications for more details (Oreopoulos and Yip, 2009; Oreopoulos et al., 2010).

Statistical Analyses. Results were plotted and analyzed on GraphPad Prism v6.01. Results were presented as mean ± S.D. with each replicate representing one biological replicate from an individual experiment. For all statistical analyses, one-way ANOVA and Dunnett’s post hoc test were conducted on GraphPad Prism v6.01 for multiple comparisons of conditions to the control. The P values were automatically adjusted by GraphPad Prism v6.01 for the multiple comparisons and P < 0.05 was considered significantly different from the control.

Results

Destabilizing Effect of Long-Term Treatment with Micromolar Concentration of VX-770 on the Abundance of VX-809-corrected F508del-CFTR Protein. As previously reported (Cholon et al., 2014; Veit et al., 2014; Matthes et al., 2016), we observed a destabilizing effect of long-term (48 hours) treatment of VX-770 at a high concentration (10 μM) at 37°C on the relative abundance of the mature band C (Fig. 1B) and immature band B (Fig. 1C) of the VX-809-rescued F508del-CFTR protein in the HEK-293 expression system. As previously reported (Cholon et al., 2014), this was not related to a significant decrease in cell viability, with comparable abundance of calnexin expression across the treatments (Fig. 1A).

Micromolar Concentration of VX-770 Reverses the Effect of VX-809 in Stabilizing the ICL4:NBD1 Interface of Cys-Less Variant of F508del-CFTR. We were prompted to determine if this destabilizing effect of VX-770 was due, in part, to a reversal of the positive effect of VX-809 on F508del-CFTR assembly. The corrector, VX-809, is thought to act by stabilizing the aberrant interaction between F508del in NBD1 and the intracellular loop ICL4 (conferring by MSD2) (Rabeh et al., 2012; Okiyoneda et al., 2013). As in previous publications (He et al., 2008; Loo et al., 2008; Serohijos et al., 2008; Aleksandrov et al., 2010; Loo and Clarke, 2011; Chin et al., 2017), we assessed this interaction by studying the propensity for chemical cross-linking between cysteine residues engineered at V510 (NBD1) and A1067 on ICL4 in the context of a full-length cysteine-less (cys-less) variant of CFTR protein for both wt and F508del. The following studies were performed at 27°C as the cys-less variant of CFTR protein is misprocessed at 37°C and requires temperature rescue at 27°C to promote its maturation (Loo and Clarke, 2006; Wang et al., 2007; He et al., 2008, 2013).

The V510C/A1067C cys-less variant of wt-CFTR runs as two bands: the mature, complex glycosylated band C and the immature, core glycosylated band B (Fig. 2A). As previously reported, band C is insensitive to Endoglycosidase (Endo) H, whereas band B is deglycosylated, and this form is labeled as band A. Both band B and C are deglycosylated by PNGase F as shown on the left panels of Fig. 2A. We confirmed that two cysteine residues (V510C on NBD1 and A1067C on ICL4) could be cross-linked by a cell-permeable maleimide reagent, bismaleimidoethane (BMOE), in a V510C/A1067C cys-less variant of wt-CFTR protein with the appearance of a cross-linked species (✝) that migrates as a 250-kDa protein on a SDS-PAGE gel (Fig. 2A) (Chin et al., 2017). We confirmed that this band (✝) was formed intramolecularly rather than intermolecular cross-linking, because it was not produced when single cysteine-containing constructs (V510C or A1067C) were expressed alone or together (Supplemental Fig. 1) (Chin et al., 2017). We showed that the band (✝) is Endo H resistant and, hence, complex glycosylated. Interestingly, an N-glycanase resistant band (∆) was apparent in the cross-linker-treated cells but not in the absence of cross-linker (Fig. 2A; Supplemental Fig. 1). We suggest on the basis of its glycosidase resistance that this unique band is the cross-linked form of band A.

A V510C/A1067C cys-less variant of F508del-CFTR is misprocessed with no appearance of band C on the immunoblot, with just the misprocessed band B expressed (Fig. 2A) (Loo et al., 2008; Loo and Clarke, 2011). Treatment with the cell-permeable cross-linker (BMOE) led to the appearance of a unique band (✝). This band (✝) shifted slightly following Endo H treatment to band (∆), confirming that it is not the C band, but rather the cross-linked band B. The band (∆) of this mutant protein is N-glycanase resistant, suggesting that it represents the cross-linked band A.

Long-term (24 hours) treatment of V510C/A1067C cys-less variant of F508del-CFTR with VX-809 (3 μM) did not lead to its maturation and the appearance of band C, likely due to the number of mutations introduced to eliminate endogenous cysteines (Fig. 2B). However, this treatment did lead to a significant increase in the cross-linking between V510C and A1067C with an increased ratio of the cross-linked band B (✝) relative to band B (Fig. 2, B and C). These findings are consistent with previous studies, showing its positive effect in improving interaction between NBD1 and ICL4 (Van Goor et al., 2011; Loo et al., 2013; Ren et al., 2013). This positive effect of VX-809 was reversed when VX-770 (10 μM) was included in this long-term treatment (Fig. 2, B and C). This abolition of NBD1:ICL4 assembly following the combination treatment could explain the effect observed on processing as documented in Fig. 1. As in the case of the protein processing studies shown in Fig. 1, the destabilizing effect of VX-770 on domain:domain assembly is dose dependent, and no destabilizing effect on cross-linking was observed following cotreatment with VX-770 at 0.1 μM (Fig. 2, B and C).
Micromolar Concentrations of VX-770 Also Exert a Destabilizing Impact on Other Membrane Proteins.

We were interested to determine if the destabilizing effect of 10 μM VX-770 was specific to F508del-CFTR. To maintain similar conditions to our cysteine cross-linking studies, we tested its specificity by applying this concentration of VX-770 at 27°C on HEK-293 cells expressing other membrane proteins, including solute carriers (SLCs): SLC26A3, SLC26A9, and SLC6A14. These membrane proteins are particularly important, because they have been shown to interact with and/or modify CFTR channel activity in various epithelial tissues (Ko et al., 2004; Xu et al., 2005; Gupta et al., 2006; Bertrand et al., 2009; Chang et al., 2009; Avella et al., 2011; Karunakaran et al., 2011; Anagnostopoulou et al., 2012). Furthermore, these SLC genes have been shown to modify CF disease phenotypes (Sun et al., 2012; Li et al., 2014b; Corvol et al., 2015; Strug et al., 2016). Thus we reason that it would be disadvantageous for CFTR modulatory drugs to also affect the expression of these membrane proteins.

Firstly, analysis of the SLC proteins by SDS-PAGE revealed multiple glycosylation states as previously reported (Li et al., 2014a). Interestingly, the steady-state abundance of all the SLCs in this study was significantly decreased with chronic treatment with micromolar concentrations of VX-770 at 27°C (Fig. 3). SLC26A3 appeared to be the most susceptible to VX-770, because its protein expression significantly decreased at concentrations of VX-770 as low as 0.1 μM (Fig. 3A). On the other hand, the abundance of SLC26A9 and SLC6A14 was significantly decreased starting at higher concentrations of 2 and 10 μM VX-770 respectively (Fig. 3, B and C). Therefore, the destabilizing effect of suprapharmacological concentrations of VX-770 is not specific to F508del-CFTR.

Lipophilicity of VX-770 Appears To Be Correlated with the Destabilizing Effect on F508del-CFTR. We propose that the nonspecific, destabilizing effects of suprapharmacological concentrations of VX-770 may be related to the known effects of this compound on the lipid bilayer (Baroni et al., 2014). To test this idea, VX-770 derivatives (Supplemental Fig. 2) with varying degrees of lipophilicity, as assessed by their partition coefficient (logP) values, were incubated with HEK-293 cells expressing temperature-rescued F508del-CFTR and their effects on the processing (band C/C1B) of F508del-CFTR determined.

In this study, the HEK-293 cells expressing F508del-CFTR were treated with VX-770 or a derivative for 24 hours at 0.1 or 10 μM at 27°C. We observed a negative correlation between the lipophilicity or logP values and the relative band C/C1B ratio (Fig. 4B). Derivatives with low logP scores (i.e., SE-02, SE-03, KT-226, and KT-227) were associated with a higher
lipophilicity of VX-770 affects F508del-CFTR stability.

**Fig. 2.** Suprapharmacological concentration of VX-770 abolishes VX-809 rescue effect at the ICL4:NBD1 interface of cys-less variant of F508del-CFTR. (A) Immunoblot of deglycosylation studies of V510C/A1067C cys-less variants of wt- and F508del-CFTR without (−) and with (+) BMOE treatment. V510C/A1067C cys-less variant of wt-CFTR (−) BMOE: No glycanase (−) treatment resulted in bands C and B. Endoglycosidase H (H) treatment resulted in band C and band A. PNGase F (F) treatment resulted in band A. V510C/A1067C cys-less variant of F508del-CFTR (−) BMOE: No glycanase (−) treatment resulted in cross-linked band C (✝), population containing band C and cross-linked band B (C*), and band B. Endoglycosidase H (H) treatment resulted in cross-linked band C (✝), population containing band C and cross-linked band B (C*), and band B. PNGase F (F) treatment resulted in cross-linked band A (n), and band A.

**Fig. 3.** VX-770 exerts destabilizing effect on membrane proteins of the solute carrier (SLC) family. (i) Representative immunoblots of (A) SLC26A3, (B) SLC26A9, and (C) SLC6A14 show different glycosylation forms of the proteins as previously reported (Li et al., 2014a). SLC proteins were treated with 0, 0.1, 1, 2, and 10 μM of VX-770 at 27°C for 24 hours with actin as loading control. (ii) Densitometry analyses of total protein relative to DMSO treatment and actin loading control of (A) SLC26A3 (n = 4 for all conditions), (B) SLC26A9 (n = 5 for all conditions), and (C) SLC6A14 (n = 4 for all conditions). SLC26A3 expression significantly decreased after 0.1 μM VX-770, SLC26A9 expression significantly decreased after 2 μM VX-770, and SLC6A14 expression significantly decreased after 10 μM VX-770. Results are presented as mean ± S.D. and analyzed by one-way ANOVA and Dunnett’s post hoc test (*P < 0.05; **P < 0.01; ***P < 0.001 compared with vehicle).

**Micromolar Concentration of VX-770 Increase the Membrane Fluidity in a Lipid Raft Model.**

Lipophilicity of VX-770 affects F508del-CFTR stability. Given the potential role for lipophilicity in VX-770-dependent effects on membrane protein stability, we were prompted to determine if the nonspecific destabilizing effects of micromolar concentrations of VX-770 were mediated by disrupting lipid rafts. We tested this hypothesis using a synthetic lipid raft model and tested the effect of micromolar VX-770 on membrane fluidity and bilayer integrity. Polarized total internal reflection fluorescence (pTIRF) microscopy was employed to study the effect of 10 μM VX-770 on supported lipid bilayers that contained DOPC/cholesterol/distearoylphosphatidylcholine band C/C+B ratio than the derivatives with high logP scores (i.e., KT-51, KT-126, KT-127, and KT-128) (Fig. 4, A and B). For example, the derivative SE-03, while exhibiting potentiator activity (Fig. 4, D and E), did not reduce steady-state protein abundance at 10 μM (Fig. 4C).
and DOPC/cholesterol/sphingomyelin (bSM) bathed in a HEPEs buffer (Simons and Toomre, 2000; Calder and Yaqoob, 2007). Two distinct bilayers were tested for the effect of VX-770 with three to six fields of view analyzed per bilayer (Fig. 5, A and B). Order parameter, $P_2$, measurements and the histogram movies demonstrated that 10 $\mu$M VX-770 immediately increased the bilayer fluidity as well as reorganized subdomains within the DOPC/bSM/cholesterol supported lipid bilayers. The histograms were determined from the selected field of view and that same field of view was used to track the dynamic changes in the bilayer upon VX-770 addition. As such, our interpretations as to the effect of these agents on membrane stability and structure are drawn from relative changes in the individual histogram characteristics for the chosen field of view. From that perspective, we can clearly see that there is a clear shift to a higher mean $P_2$ value and a tighter distribution upon VX-770, which is strongly suggestive of increased membrane fluidity. This interpretation is certainly bolstered by the before- and after-VX-770 addition, which shows significant changes in the bilayer domain structures. Conversely, the relative decrease and broadening of the distribution of $P_2$ values (i.e., emergence of a second lower $P_2$ peak) upon SE-03 addition can only be interpreted at this point as evidence of a greater extent of membrane reorganization. Intriguingly, the extent of subdomain restructuring due to SE-03 addition appears to be significantly less than in the VX-770 case, as is evidenced by the relatively minor changes in bilayer morphology.
Discussion

The current studies provide a potential explanation for the concentration-dependent, destabilizing effects of chronic exposure to VX-770 on VX-809-rescued F508del-CFTR protein (Cholon et al., 2014; Veit et al., 2014; Matthes et al., 2016). At the concentrations of VX-770 that reduce F508del-CFTR protein stability, we also observed reduction of the abundance of other membrane proteins (SLC26A3, SLC26A9, and SLC6A14), pointing to its lack of specificity. From our studies of model lipid rafts, we deduce that this negative effect of high concentrations of VX-770 is related to its lipophilicity and disruption of these membrane structures.

Our biochemical studies showed that the concentration of VX-770 (10 μM) that reduced VX-809-mediated augmentation in F508del-CFTR processing also reversed VX-809-mediated enhancement of the interaction between the intracellular loop conferred by MSD2 (ICL4) and NBD1. As in previous studies (He et al., 2008; Loo et al., 2008; Serohijos et al., 2008; Aleksandrov et al., 2010; Loo and Clarke, 2011; Chin et al., 2017), we probed the interaction at this interface in the context of a cysteine-less variant of F508del-CFTR protein harboring cysteine residues engineered in ICL4 (A1067C) and NBD1 (V510C). In our experimental system, we could not detect VX-809-induced maturation and processing of the V510C/A1067C cysteineless variant of F508del-CFTR protein in immunoblots. However, VX-809-induced interface assembly in the immature, core glycosylated band B of the protein was detectable as a unique cross-linked band (*) and the relative abundance of this band was reduced in the presence of 10 μM VX-770 but not in the presence of 100 nM VX-770. Hence, we interpret the destabilizing effects of VX-770 at this concentration to be due to inhibition of the positive effects of VX-809 at this interface. However, as discussed in the subsequent paragraphs, this effect is likely not specific to this interface or even the mutant CFTR protein.

The same concentration of VX-770 that induced destabilizing effects on F508del-CFTR assembly also reduced the steady-state abundance of the three SLC proteins tested. We chose these proteins because they are known to be localized in the same tissues and membranes as CFTR and modify epithelial transport function (Ko et al., 2004; Mount and Romero, 2004; Sun et al., 2012). Hence, any nonspecific effect of VX-770 on these membrane proteins may have a potential to modify disease and/or therapeutic response to ORKAMBI. Of these proteins, SLC26A3 was the most sensitive to the destabilizing effect of VX-770, with a significant decrease in protein abundance apparent (at least in the HEK-293 expression system) at 1 μM, the serum concentration reported by Vertex Pharmaceuticals (https://pi.vrtx.com/files/uspi_ivacaftor.pdf). Previous studies have shown that loss of function mutations of SLC26A3 caused a rare disease known as congenital chloride-losing diarrhea via impaired chloride absorption and fluid/bicarbonate secretion (Höglund et al., 1996). Intestinal inflammation, which is common in diarrhea, has been found to further

(Fig. 5, C and D), supporting the claim the VX-770 effects are mediated at least in part by its relative lipophilicity.
reduce SLC26A3 expression through direct binding to NF-κB (Kumar et al., 2017). Hence, it is possible that the destabilizing effect of VX-770 that we observed on SLC26A3 may modify the ORKAMBI response in the intestinal tissues.

SLC26A9 confers an apical anion channel, and this function augments the CAMP-regulated channel activity of CFTR in the non-CF airway epithelium and in the epithelium of patients bearing gating mutations such as G551D (Bertrand et al., 2009, 2017). Strug et al. (2016) showed that polymorphisms in the SLC26A9 gene associate with the response of patients bearing G551D to VX-770 and interpreted these results to suggest that functional expression of SLC26A9 enhances the therapeutic efficacy of VX-770 in this cohort of patients. Therefore, VX-770 in patient tissues is not impairing the positive effect of SLC26A9 on the potentiation of the G551D channel. On the other hand, the misfolded F508del-CFTR protein interacts with and prevents SLC26A9 trafficking to the apical surface (Bertrand et al., 2009, 2017). Correction of F508del-CFTR by VX-809 is thought to promote SLC26A9 trafficking to the cell surface indirectly (Bertrand et al., 2009, 2017). In vitro studies of bronchial explant tissue from F508del homozygotes suggest that the magnitude of pharmacological correction by VX-809 is also augmented by SLC26A9 expression (Strug et al., 2016). Hence, the results of the current study suggest that the possible deleterious effect of chronic VX-770 on SLC26A9 protein stability should be further studied given its importance in modifying the pharmacological rescue conferred by this therapy.

SLC6A14 is an apical amino acid transporter, and we have shown that it functions to remove nutrient amino acids from the surface of respiratory epithelial cells (Di Paola et al., 2017). This function is associated with reduced adherence of P. aeruginosa to the surface of respiratory epithelial cells (Di Paola et al., 2017). We speculate that reduced surface stability of SLC6A14 induced by high concentrations of VX-770 would potentially enhance P. aeruginosa attachment and biofilm formation and we are testing this hypothesis.

We propose that the destabilizing effect of VX-770 is related to its lipophilic property. Interestingly, treatment of F508del-CFTR with a panel of VX-770 derivatives showed that the degree of VX-770 lipophilicity is correlated to its destabilizing effect. SE-03 lacks two tert-butyl groups on the phenol ring (Supplemental Fig. 2) and is less lipophilic than VX-770 (the logP value of SE-03 is 2.67, whereas it is 5.76 for VX-770), yet it retains activity as a potentiator. Interestingly, SE-03 does not reduce F508del-CFTR protein stability even at the high concentration of 10 μM. Biophysical studies of a lipid raft model showed that unlike VX-770, SE-03 addition did not disperse lipid rafts comprised of cholesterol and sphingomyelin. Together, these findings support the hypothesis that the nonspecific effects of VX-770 on membrane protein stability are mediated through VX-770 interaction with and disruption of lipid rafts, a membrane organizing structure known to facilitate protein interactions and signaling (Bi et al., 2001; Patel et al., 2008; Steinberg, 2008; Abu-Arish et al., 2015).

The nonspecific effect of chronic VX-770 on the steady-state abundance of F508del-CFTR, other membrane proteins, and lipid rafts was observed at the suprapharmacological concentration of 10 μM VX-770. According to Matthes et al. (2016), a peak free plasma concentration of VX-770 is 1.5–8.5 nM, considerably lower than other estimates and the concentration at which we observed the nonspecific effects of VX-770 (Hanrahan et al., 2017). According to the drug data sheet provided by Vertex, after a single recommended dose of 150 mg administered after a meal, the peak plasma concentration reaches 768 ng/ml or 1 to 2 μM (https://pi.vrtx.com/files/uspi_ivacaftors.pdf). This value is an order of magnitude less than the concentration (10 μM) that caused significant loss of CFTR and other membrane proteins (including SLC26A9) in the in vitro studies of this paper. However, according to a previous study (Trillert and Hug, 2014), there was a large range in serum concentrations of VX-770 in different patients, ranging from 400 to 3000 ng/ml, with the upper limit approaching 10 μM. Importantly, the concentration of VX-770 in the tissues of CF patients treated with VX-770 is uncertain. According to the European Medicine Agency (EMA/473279/2012), VX-770 accumulates in lung and tracheal epithelial tissues relative to plasma in rat studies (http://www.ema.europa.eu/docs/en_GB/document_library/EPAR_-_Public_assessment_report/human/002494/WC500130766.pdf). Hence, it remains unknown if tissue concentrations could approach the higher concentration of 10 μM shown to exert nonspecific, destabilizing effects.

It is important to acknowledge that VX-770 exerts a beneficial effect overall when given as a monotherapy to individuals bearing gating mutations in CFTR. However, our studies suggest that future clinical studies are required to monitor off-target effects in this population taking this monotherapy continuously over their lifetime. Finally, it remains possible that the variable clinical efficacy of ORKAMBI in the F508del patient population is partially associated with variation in its tissue accumulation among patients and/or off-target effects of VX-770 on other membrane proteins variably expressed in different patients.

Acknowledgments
We thank Dr. David Clarke for providing the V510C, A1067C and V510C/A1067C cys-less variants of wt-CFTR. We thank Dr. Reinhart Reithmeier and OriGene for providing the SLC constructs. We thank Dr. Steven Molinski for the helpful discussions on these studies. We thank Dr. Jacqueline McCormack for help in editing the manuscript.

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