TITLE:
SYNERGY-BASED SMALL-MOLECULE SCREEN USING A HUMAN LUNG EPITHELIAL CELL LINE YIELDS ΔF508-CFTR CORRECTORS THAT AUGMENT VX-809 MAXIMAL EFFICACY

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Abbreviations: CFTR, cystic fibrosis transmembrane conductance regulator; ER, endoplasmic reticulum; CFBE, cystic fibrosis bronchial epithelial; MSD, membrane-spanning domains; NBD, nucleotide binding domains; YFP, yellow fluorescence protein; PBS, phosphate-buffered saline; DMSO, dimethyl sulfoxide; HRP, horseradish peroxidase; HA, human influenza hemagglutinin; VX-770, N-(2,4-di-tert-butyl-5-hydroxyphenyl)-4-oxo-1,4-dihydroquinoline-3-carboxamide; VX-809, 3-[6-[[1-(2,2-difluoro-1,3-benzodioxol-5-yl)cyclopropyl]carbonyl]amino]-3-methyl-2-pyridinyl]-benzoic acid.
ABSTRACT

The most prevalent CFTR mutation causing cystic fibrosis, ΔF508, impairs folding of nucleotide binding domain 1 (NBD1) and stability of the interface between NBD1 and the membrane spanning domains (MSDs). The interfacial stability defect can be partially corrected by the investigational drug VX-809 or the R1070W mutation. ‘Second-generation’ ΔF508-CFTR correctors are needed to improve on the modest efficacy of existing CF correctors. We postulated that a second corrector targeting a distinct folding/interfacial defect might act in synergy with VX-809 or the R1070W suppressor mutation.

A biochemical screen for ΔF508-CFTR cell surface expression was developed in a human lung epithelium-derived cell line (CFBE41o-) by expressing chimeric CFTRs with a horseradish peroxidase (HRP) in the fourth exofacial loop either in the presence or absence of R1070W. Using a luminescence read-out of HRP activity, screening of ~110,000 small molecules produced 9 novel corrector scaffolds that increased cell surface ΔF508-CFTR expression by up to 200% in the presence vs. absence of maximal VX-809. Further screening of 1006 analogs of compounds identified from the primary screen produced 15 correctors with EC50 <5 µM. 8 chemical scaffolds showed synergy with VX-809 in restoring chloride permeability in ΔF508-expressing A549 cells. An aminothiazole increased chloride conductance in human bronchial epithelial cells from a ΔF508 homozygous subject beyond that of maximal VX-809. Mechanistic studies suggested that NBD2 is required for the aminothiazole rescue.

Our results provide proof-of-concept for synergy screening to identify second-generation correctors, which, when used in combination, may overcome the ‘therapeutic ceiling’ of first-generation correctors.
INTRODUCTION

CFTR (cystic fibrosis transmembrane conductance regulator) is a cAMP-regulated chloride channel expressed in airway and other epithelia. CFTR is a large membrane glycoprotein containing two membrane-spanning domains (MSD1-2) and three cytoplasmic domains, which include two nucleotide binding domains (NBD1-2) and a regulatory domain (Riordan, 2005). Mutations in CFTR cause the genetic disease cystic fibrosis (CF) in which lung infection and mucus accumulation can lead to life-threatening deterioration of lung function. The most common CFTR mutation, deletion of phenylalanine at residue 508 (∆F508), is present in at least one allele in ~90% of CF patients. The ∆F508 mutation produces CFTR misfolding, with retention at the endoplasmic reticulum (ER), accelerated CFTR degradation at the ER and periphery, and impaired chloride channel gating (Balch et al.; 2011, Du et al., 2009; Gadsby et al., 2006; Riordan, 2008). The reduced plasma membrane chloride permeability in CFTR-expressing cells containing the ∆F508 mutation has been proposed to produce the clinical phenotype by a variety of mechanisms involving abnormal airway surface liquid homeostasis, reduced gland fluid secretion, defective immune cell function, and others (Boucher et al., 2004; Cohen and Prince, 2012; Zielenski et al., 2000).

There has been considerable effort and progress in the development of CFTR-targeted small-molecule therapeutics for CF (Ashlock and Olson, 2011). The CFTR ‘potentiator’ VX-770 (Ivacaftor), which corrects defective channel gating of some CFTR mutants, has been approved for CF therapy caused by the defective channel gating but unimpaired cellular processing and plasma membrane targeting of the G551D-CFTR mutation (Accurso et al., 2010; Van Goor et al., 2009). Several small-molecule ‘correctors’ of ∆F508-CFTR cellular processing have been identified (Pedemonte et al., 2005a, 2005b; Phuan et al., 2011; Robert et al., 2010; Van Goor et al., 2006; Yu et al., 2008). VX-809 is in clinical trials for CF caused by the ∆F508 mutation (Clancy et al., 2012). However, to date VX-809 and other correctors show limited efficacy in primary human bronchial cell cultures from ∆F508-homozygous CF patients, restoring only ~15% of full CFTR activity found in cultures from non-CF patients (Van Goor et al., 2011). Clinical trials with VX-809, alone or together with VX-770, have thus far shown minimal efficacy (NCT01225211). The need for ‘second-generation’ correctors with...
improved efficacy has been widely acknowledged (Hanrahan et al., 2013; Lukacs and Verkman, 2012; Okiyoneda and Lukacs, 2012).

Here, we report proof-of-concept for a ‘synergy screening’ approach to identify second-generation ΔF508-CFTR correctors. The idea for synergy screening is that global misfolding and dysfunction of ΔF508-CFTR can be efficiently reversed by stabilizing two major structural deficiencies; the NBD1 stability and the NBD-MSDs interfacial defects (Rabeh et al., 2010), which is supported by data from suppressor mutations and corrector combinations (Okiyoneda et al., 2013). While VX-809 is unable to restore the thermal stability of the isolated ΔF508-NBD1 (Farinha et al., 2013; Okiyoneda et al. 2013; Ren et al., 2013), it is thought to directly stabilize the interface between NBD1 and MSDs (Farinha et al., 2013; Okiyoneda et al. 2013; Ren et al., 2013; Van Goor et al. 2011). Combining genetic modification (e.g. suppressor mutation in the NBD1; 3S) of the ΔF508-NBD1 or chemical chaperones with VX-809 produced robust potentiation of VX-809 effect on the folding and cell surface expression of ΔF508-CFTR (Okiyoneda et al., 2013). Therefore, novel corrector molecules targeting either the NBD1 and/or its interface defects should be preferentially identified by screening in the background of VX-809 or the interface-stabilizing mutation R1070W. To this end, here we established, using human lung epithelium-derived cell lines (CFBE41o-, Ehrhardt et al., 2006; Veit et al., 2012), a novel biochemical screening procedure. Compounds were identified that, when combined with VX-809, increased correction efficacy beyond that of maximal VX-809 alone, supporting the idea of synergy screening and corrector combination therapy for CF caused by the ΔF508 mutation.

MATERIALS AND METHODS

Cell lines

Doxycycline-inducible expression systems were generated by lentivirus transduction using the Lenti-X TetON Advanced Inducible Expression System (Clontech, Mountain View, CA) as described (Veit et al, 2012). For expression of extracellular HRP-tagged CD4TM- ΔF508-NBD1-1S chimeras the extracellular CD4 domain of the previously described CD4T-ΔF508-NBD1-1S (Rabeh et al., 2012) was replaced in-frame with the catalytic domain of horseradish peroxidase (HRP). Madin-Darby canine
kidney (MDCK) type II cells stably expressing HRP-CD4TM-ΔF508-NBD1-1S were maintained in Dulbecco’s modified Eagle’s medium (DMEM) (Invitrogen, Carlsbad, CA) supplemented with 10% fetal bovine serum (FBS) under puromycin (3 μg/ml) and G418 selection (0.2 mg/ml). BHK cells expressing ΔF508 and ΔF508-1218X CFTR-3HA variants have been described (Okiyoneda et al., 2013). The cloning and characterization of 3HA-tagged variants of ΔF508-CFTR, R1070W-ΔF508-CFTR, and 3S-ΔF508-CFTR- (containing the F494N, Q637R and F429S NBD1 suppressor mutations) have been described (Okiyoneda et al., 2013). To replace the 3HA tag in the ΔF508-CFTR variants, the horseradish peroxidise isoenzyme C was introduced into the 4th extracellular loop by using the EcoRV/AvrII restriction sites with a 5’ linker (ctcgaatcaggaggtagtggtggcggaagt) linker, but without a 3’ linker. CFBE41o- cells were grown in minimal essential medium (MEM, Invitrogen) supplemented with 10% FBS, 2 mM L-glutamine and 10 mM HEPES. For propagation, the CFBE41o- cells were cultured in plastic flasks coated with an extracellular matrix consisting of 10 μg/ml human fibronectin, 30 μg/ml collagen from calf skin (Sigma-Aldrich) and 100 μg/ml bovine serum albumin (Sigma-Aldrich) diluted in LHC-8 basal medium (Invitrogen).

For high-throughput screening, CFBE41o- Tet-on cells were plated in black, 96-well microplates (Costar, Corning Inc.) at 15,000 cells/well. ΔF508-CFTR expression was induced 24 h after plating with 0.5 μg/ml doxycycline treatment for 2 d before screening. A549 lung epithelial cells (ATCC CCL-185) stably expressing ΔF508-CFTR (Pedemonte et al., 2010) were provided by Dr. Luis Galietta (Genoa, Italy) and cotransfected with halide-sensitive green fluorescent protein YFP-H148Q/I152L/F46L (Galietta et al, 2001). A549 cells were cultured in DMEM/Ham’s F12 (1:1) containing 10% FBS, 2 mM L-glutamine, 100 U/ml penicillin and 100 μg/ml streptomycin. For functional assay, A549 cells were plated in black, 96-well microplates (Costar, Corning Inc.) at 10,000 cells/well. For short-circuit current measurements, primary cultures of human CF bronchial epithelial cells were isolated and grown at an air-liquid interface for at least 21 days on cell culture inserts (Snapwell; Corning, Lowell, MA) and used when transepithelial resistance was > 1000 μΩ/cm², as described (Yamaya et al., 1992; Fulcher and Randell, 2013).
**Compounds**

A total of 110,000 diverse drug-like synthetic compounds (>90% with molecular size 250-500 Da; ChemDiv Inc.) were used for screening. For optimization, 1006 commercially available analogs from different classes of active compounds from the primary screens were tested.

**Screening procedures**

Screening was carried out using a Beckman Coulter (Fullerton, CA) platform (Biomek FX). In one set of assays R1070W-∆F508-CFTR-HRP (R1070W-HRP) expressing CFBE41o- cells were incubated with 100 µL medium containing 25 µM test compounds and 0.5 µg/ml doxycycline, for 24 h at 37 °C. In a second set of assays ∆F508-CFTR-HRP (∆F508-HRP) expressing CFBE41o- cells were incubated with 100 µL medium containing 25 µM test compounds, 2 µM VX-809 and 0.5 µg/ml doxycycline for 24 h at 37 °C. All compound plates contained negative controls (DMSO vehicle) and positive controls (2 µM VX-809). In both assays the cells were washed 4 times with PBS, and HRP activity was assayed by addition of 50 µL/well of HRP-substrate (WesternBright Sirius Kit, Advansta Corp, Menlo Park, CA). After shaking for 5 min, chemiluminescence was measured using a TECAN Infinite M1000 plate reader (TECAN Groups Ltd, Mannedorf, Switzerland) equipped with automated stacker (integration time, 100 ms). Z’ is define as $Z' = 1 - \left[ (3 \times \text{standard deviation of maximum signal control} + 3 \times \text{standard deviation of minimum signal control}) / \text{absolute(mean of maximum signal control-mean of minimum signal control)} \right]$ (Zhang et al., 1999).

**Functional assays**

A549 cells expressing ∆F508-CFTR-YFP were grown at 37 °C / 5% CO₂ for 18-24 h after plating. The cells were then incubated with 100 µL of medium containing test compounds for 18-24 h. At the time of the assay, cells were washed with PBS and then incubated for 10 min with PBS containing forskolin (20 µM) and genistein (50 µM). Each well was assayed individually for I⁻ influx by recording fluorescence continuously (200 ms per point) for 2 s (baseline) and then for 12 s after rapid addition of 165 µL PBS in which 137 mM Cl⁻ was replaced by I⁻. Initial I⁻ influx rate was computed by fitting the final 11.5 seconds
of the data to an exponential for extrapolation of initial slope, which was normalized for background-subtracted initial fluorescence. All compound plates contained negative controls (DMSO vehicle) and positive controls (5 µM VX-809). Fluorescence was measured using a TECAN Infinite M1000 plate reader equipped with a dual syringe pump (excitation/emission 500/535 nm).

**Short-circuit current measurements**

Test compounds (without or with 10 µM VX-809) were incubated with primary human CF bronchial epithelial cells from ΔF508-CFTR–homozygous subjects at the basolateral side for 18-24 h at 37 °C prior to measurements. The apical and basolateral chambers contained identical solutions: 130 mM NaCl, 0.38 mM KH2PO4, 2.1 mM K2HPO4, 1 mM MgCl2, 1 mM CaCl2, 25 mM NaHCO3 and 10 mM glucose. Solutions were bubbled with 5% CO2/95% O2 and maintained at 37 °C. Hemichambers were connected to a DVC-1000 voltage clamp (World Precision Instruments Inc., Sarasota, FL) via Ag/AgCl electrodes and 1 M KCl agar bridges for recording of short-circuit current.

**CFTR plasma-membrane (PM) density measurements**

The PM density of 3HA-tagged CFTR variants was determined by cell surface ELISA (Okiyoneda et al., 2010). HRP-tagged CFTR PM density was measured in a VICTOR Light plate reader (PerkinElmer, Waltham, MA) after addition of 50 µl/well HRP-Substrate (SuperSignal West Pico, Thermo Fisher Scientific, Waltham, MA). PM density measurements were normalized with cell viability determined by Alamar Blue assay (Invitrogen, Carlsbad, CA).

**Differential scanning fluorimetry (DSF)**

Isolation of recombinant human NBD1 containing a single suppressor mutation (1S; F494N) and melting temperature measurement were performed as described (Rabeh et al., 2012). DSF scans of NBD1 (6 µM) were done in 150 mM NaCl, 20 mM MgCl2, 10 mM HEPES and 2.5 mM ATP, pH 7.5 using a Stratagene Mx3005p (Agilent Technologies, La Jolla, CA) qPCR instrument in the presence of 2× Sypro Orange. These studies were performed on recombinant NBD-1S, since we have not seen discernible
differences in the relative thermal stability NBD1 and NBD-1S in the presence of the corrector panel (Rabeh et al., 2012).
RESULTS

Development and validation of synergy-based CFTR screens

Primary screening was done using two stably transfected human lung epithelium-derived (CFBE41o-) cell lines (Ehrhardt et al., 2006). One screen (‘ΔF508 screen’) (Fig. 1A) utilized CFBE41o-cells transfected with humanΔF508-CFTR with a horseradish peroxidase (HRP) inserted in its fourth extracellular loop (ΔF508-HRP CFBE41o-). A second screen (‘R1070W screen’) (Fig. 1B) utilized CFBE41o- cells transfected with ΔF508-CFTR-HRP containing a R1070W mutation (R1070W-HRP CFBE41o-). Cells were cultured on 96-well plates and CFTR synthesis was induced 48 h prior to screening. Cells were grown to confluence prior to addition of test compounds. For the ΔF508 screen cells were incubated with test compounds (at 25 µM) together with 2 µM VX-809; for the R1070W screen cells were incubated with test compounds (at 25 µM) alone. The 25 µM concentration was chosen following initial small-scale screens showing a small percentage of active compounds. After incubation for 18-24 h at 37 ºC, cells were washed and HRP substrate was added for luminescence read-out. Test compounds that were cytotoxic at the screening concentration (25 µM) will result in reduced luminescence signal.

VX-809 produced a concentration-dependent increase in HRP luminescence signal after incubation with cells at 37 ºC or 27 ºC (Fig. 1A and 1B) in both cell lines, with similar EC₅₀ of ~0.3 µM. In ΔF508-HRP CFBE41o- cells at 37 ºC VX-809 increased signal maximally to ~250 luminescence arbitrary units (a.u.) over DMSO-control baseline of ~60 a.u., representing a ~4-fold signal increase. Similarly, with the R1070W-HRP CFBE41o- cells, VX-809 increased signal maximally to ~220 a.u. over DMSO-control baseline of ~85 a.u., representing a ~2.5-fold signal increase (bar graphs in Fig. 1A and 1B). Therefore, both cell lines produced robust signals with a good dynamic range for high-throughput screening.

Low-temperature rescue (27 ºC) of ΔF508-CFTR increased HRP luminescence signal by ~2-fold (compared with 37 ºC) in ΔF508-HRP CFBE41o- cells and ~3-fold in R1070W-HRP CFBE41o- cells. VX-809 and low-temperature together further increased HRP luminescence. EC₅₀ values were 30 nM and 78 nM in the low-temperature rescued ΔF508-HRP and R1070W-HRP CFBE41o- cells.
respectively.

Preferential correction of ΔF508-CFTR-3HA with the NBD1 stabilizing 3S mutations (F494N, Q637R and F429S) compared to CFTR carrying the R1070W interface stabilizing mutation has been taken as evidence that VX-809 preferentially stabilizes the interface between NBD1 and MSDs but not the NBD1 folding defect CFTR (Okiyoneda et al., 2013). This was preserved in the CFTR-HRP context, indicating that the HRP fusion preserved behavior of the ΔF508 variants (Supplemental Fig. 1). The relative insensitivity of R1070W-HRP to VX-809 was utilized to identify correctors that act in synergy with VX-809.

**Identification of ΔF508-CFTR correctors by synergy screens**

A total of 110,240 drug-like small synthetic molecules were tested in the ΔF508 and R1070W screens. As summarized in Fig. 2A, in the ΔF508 screen 164 active compounds were identified based on >50% increase in luminescence signal over that of 2 μM VX-809 alone. After re-testing, five compounds, grouped into three classes, were confirmed from the ΔF508 screen. Fig. 2B shows the structures of the three most active compounds, H-01, J-01 and K-01. For the R1070W screen 25 active compounds were identified based on >50% increase in luminescence signal over that of DMSO. After retesting, nine compounds, grouped into six classes, were confirmed from the R1070W screen. Fig. 2D shows structures of the six most active compounds, A-01, B-01, C-01, D-01, E-01 and F-01.

Because different small molecule collections were used for the ΔF508 and R1070W screens, we cross-tested all active correctors in both the ΔF508-HRP and R1070W-HRP CFBE41o- cell lines (Supplemental Fig. 2). Five compounds, A-01, B-01, C-01, H-01 and K-01, were active in both cell lines. However, compounds D-01, E-01 and F-01, discovered from the R1070W screen, were not active in ΔF508-HRP CFBE41o- cells. J-01, discovered from the ΔF508 screen, was not active in R1070W-HRP CFBE41o- cells. We further tested the concentration-dependent activities of A-01, B-01, H-01 and K-01 (in the presence of 2 μM VX-809) in ΔF508-HRP CFBE41o- cells (Fig. 3A). H-01 was the most potent corrector with EC₅₀ ~1.5 μM and maximal signal >300% over that produced by 2 μM VX-809. However, H-01, when tested alone in ΔF508-HRP CFBE41o- cells, had little activity (Supplemental Table 1). We
also measured the concentration-dependent activities of A-01, B-01, D-01, H-01 and K-01 (Fig. 3B) in R1070W-CFBE41o- cells and found that D-01 is the most potent corrector with EC\textsubscript{50} \approx 1.2 \ \mu M and maximal signal \approx 65\% of that produced by 2 \ \mu M VX-809. Supplemental Table 1 and Supplemental Fig. 2 summarize EC\textsubscript{50} of all compounds in both cell lines, with and without 2 \ \mu M VX-809. We conclude that H-01 and D-01 are the most active class of compounds discovered from the screens.

To confirm that the HRP luminescence assay reports the apical plasma membrane CFTR in the CFBE41o- cells, the relative correction determined in the HRP assay was compared to that detected using an extracellular 3xHA tagged ΔF508-CFTR (ΔF508-CFTR-3HA) expressed in CFBE41o- cells by cell-surface ELISA, as described (Veit et al., 2012). A linear correlation was found for a panel of correctors (Fig. 3C), confirming the results obtained from the CFTR-HRP luminescence assay.

**Structure-activity analysis**

1006 commercially available analogs of active compounds were tested to establish structure-activity relationships. Fig. 4A shows concentration-dependence data of H analogs (in the presence of 2 \ \mu M VX-809) in ΔF508-HRP CFBE41o- cells. Several class H analogs increased HRP luminescence with low micromolar EC\textsubscript{50}. Similar compound potency and efficacy was found for class D analogs in R1070W-HRP CFBE41o- cells (Fig. 4C). Supplemental Table 2 and Supplemental Fig. 3 summarize EC\textsubscript{50} for the most active class D and H analogs. We found that the most active analogs have similar activities as the original compounds identified in the primary screen. Structural determinants of activity for class D and H compounds are summarized in Figs. 4B and 4D. Class D correctors are 2-aminothiazoles, with best activities found for analogs with R\textsuperscript{1} substituents phenyl, thiophene and furan. Electron-withdrawing aromatic groups at the R\textsuperscript{1} position, such as nitrophenyl, biphenyl, pyridine and naphthalene, reduced activity. Active groups on the thiazole include methyl (for example, D-02, D-03), naphthalenes (D-04, D-05) and 6-methyl-cyclohexyl (D-01) rings. Rings such as hindered t-butyl-cyclohexyl and cyclohexanone reduced activity. Class H analogs contained a unique dihydrossoiro-indene scaffold. Analogs with different substituents on the phenyl ring (R\textsuperscript{1}) and on the nitrogen (R\textsuperscript{2}) were examined. For R\textsuperscript{1}, class H analogs with the phenyl ring substituted at the 4-position...
with halides and electron-neutral alkyl groups were active. Substitution at the 2-position on the phenyl ring reduced activity. For the R\(^2\) position, analogs with naphthalene and di-substituted phenyl rings had greatest activity, whereas electron-withdrawing group such as benzyl, nitro-substituted phenyl and cyclohexyl rings reduced activity.

**Functional measurements of halide transport in human A549 lung epithelial cells**

A cell-based fluorescence assay of iodide influx was used for functional studies. Human lung epithelium-derived A549 cells stably expressing ΔF508-CFTR and an iodide-sensitive YFP were incubated with the test compounds, without or with VX-809, at 37 °C for 24 h (Fig. 5A). Iodide influx was measured by addition of extracellular iodide in the presence of maximal concentration of a potentiator (50 µM genistein) and a cAMP agonist (20 µM forskolin). Representative iodide influx data for fluorescence plate reader assays of D-01 and H-01, alone or with 2 µM VX-809, are shown in Fig. 5B. Increased ΔF508-CFTR conductance is seen as a greater negative slope. Concentration-dependence data for A-01, B-01, D-01, H-01, K-01, without and with VX-809, are shown in Fig. 5C. Supplementary Table 3 summarizes EC\(_{50}\) and V\(_{\text{max}}\) values. The compounds showed moderate functional activity in A549 cells. For example, the EC\(_{50}\) of aminothiazole D-01 is 2.9 µM with V\(_{\text{max}}\) 43% of that produced by 2 µM VX-809. When added together with VX-809, most correctors increased ΔF508-CFTR function by ~20% over maximal VX-809 (2 µM). D-01 and H-01 were most active, with similar EC\(_{50}\) ~0.6 µM and V\(_{\text{max}}\) ~120%.

**Mechanistic studies of corrector action**

To assess whether the correctors identified here can restore ΔF508-NBD1 stability in vivo, cell surface expression of the HRP-CD4-ΔF508-NBD1-1S chimera was measured in MDCK-II cells. This approach probes the in vivo conformational stability of the isolated NBD1 tethered to a reporter molecule, as based on our observation that the CD4-ΔF508-NBD1 chimera cell surface density is proportional to NBD1 thermal stability (Rabeh et al., 2012). The plasma membrane density of the HRP-CD4-ΔF508-NBD1-1S chimera was increased in the presence of 30 µM C-01 to the level detected
in the presence of the chemical chaperone glycerol (5%) (Fig. 6A) (Sato et al., 1996). The C-01 induced in vivo rescue could be accounted for by the direct thermo-stabilization of the NBD1-1S, based on the melting temperature ($T_m$) shift of the domain as monitored by differential scanning fluorimetry (Rabeh et al., 2012). 30 µM C-01 was able to increase the NBD1-1S $T_m$ to that found in the presence of 5% glycerol (Fig. 6B and data not shown). C-01 is one of few compounds so far described to act directly on NBD1 (Okiyoneda et al., 2013). Even though this compound acted only at 30 µM and showed limited efficacy on full-length CFTR, its scaffold might be used for future SAR studies.

It has been demonstrated that corrector 4 (C4)-mediated rescue of the ∆F508 CFTR folding defect requires the presence of the NBD2 in CFTR (Okiyoneda et al., 2013). To investigate whether any of the correctors identified here may target the NBD2 interface, their synergy with C4 was evaluated in R1070W-HRP CFBE41o- cells using the HRP luminescence assay. While the relative rescue efficiency of A-01, B-01, C-01 and K-01 did not change in the presence of C4, the D-01 rescue effect was prevented by C4 (Fig. 6C). This observation suggests that D-01 and C4 may target an overlapping site on NBD2. In agreement, truncated ∆F508-CFTR-3HA lacking the NBD2 (∆F508-1218X-CFTR) prevented rescue by D-01 regardless of the presence of VX-809 (Fig. 6D), supporting the conclusion that NBD2 may represent one of the targets of D-01. These results provide a mechanistic basis of the synergistic action of D-01 with VX-809 as seen in the functional measurements.

**Short-circuit measurements in primary cultures of ∆F508-homozygous human bronchial CF epithelial cells**

Functional assays were also done in well-differentiated primary cultures of human bronchial epithelial cells from a ∆F508-CFTR–homozygous subject. Short-circuit current was measured across electrically tight polarized monolayers cultured on porous supports at an air-liquid interface in which test compounds were added to the bath solution for 18-24 h prior to measurements. Sodium current was blocked with amiloride. Cells incubated with 10 µM VX-809 alone (Blanchard et al., 2014) showed increased short-circuit current in response to forskolin and genistein, which was inhibited by CFTR$_{inh}$-172 (Fig. 7A). Little current was seen in the absence of VX-809. D-01 at 30 µM together with
10 μM VX-809 increased chloride conductance more than that of VX-809 alone. D-01 alone did not increase current in the bronchial epithelial cells. The other correctors, including H-01, when tested in human bronchial epithelial cells of the same ΔF508-CFTR–homozygous subject, showed small increases in chloride current (Supplemental Fig. 4). Fig. 7B summarizes chloride current responses following forskolin and forskolin + genistein. Chloride current increased by 2.3 μA/cm² (forskolin) and 3.4 μA/cm² (forskolin + genistein) when cells were treated with D-01 and VX-809 versus VX-809 alone. The fractional stimulation of short-circuit current by forskolin alone compared to the maximum current measured in the presence of forskolin and genistein was increased two-fold by D-01, suggesting that the corrector combination favors the ΔF508-CFTR native-like conformation, which is more susceptible to activation by forskolin and less dependent on genistein.
DISCUSSION

This study was done to investigate the idea that a synergy screen might identify ΔF508-CFTR correctors, which, when used in combination, would have greater maximal efficacy than individually used correctors. The underlying hypothesis is that distinct structural defects in ΔF508-CFTR each require correction, such that simultaneous correction of distinct defects would achieve greater efficacy than correction of a single defect. One screen was done using cells expressing ΔF508-CFTR in which test compound was added together with VX-809, an established corrector that has been extensively characterized and is in clinical trials. Though the precise correction mechanism of VX-809 has not been resolved, current data suggests that VX-809 may target multiple sites at the NBD1-MSDs interface and interact with the N-terminal fragment of CFTR, represented by MSD1 or MSD1-NBD1 (Farinha et al., 2013; He et al., 2013; Loo et al., 2013; Okiyoneda et al., 2013; Ren et al., 2013). Mutagenesis studies and thermo-stabilization of ΔF508-CFTR suggest that VX-809 interacts directly with the channel (Okiyoneda et al., 2013), although indirect effects cannot be precluded. A second screen was done in cells expressing R1070W-ΔF508-CFTR (in the absence of VX-809), since in the background of genetically stabilized ΔF508-NBD1 the R1070W mutation was necessary and sufficient to restore robust CFTR domain assembly and cell surface expression (Mendoza et al., 2012; Rabeh et al., 2012; Thibadeau et al., 2010).

Screening was done using a human lung epithelium-derived cell line, CFBE41o-, that was stably transfected with HRP-tagged ΔF508-CFTR or R1070W-ΔF508-CFTR. The CFBE41o- cell line was selected as a readily transfectable cell line that is predicted to recapitulate the human bronchial epithelium (Ehrhardt et al., 2006). However, it is recognized that quality control mechanisms for ΔF508-CFTR processing are cell type-dependent (Pedemonte et al., 2010), so that there are potential concerns for screens done with any cell line. We note that VX-809 is an analog of a corrector identified in a ΔF508-CFTR mouse fibroblast cell line (Van Goor et al., 2010). The CFTR constructs used here for screening were engineered with an HRP in their fourth extracellular loop for robust plate reader-based luminescence measurements of cell surface CFTR expression. The constructs were transfected using a tetracycline-inducible promoter in order to prevent phenotypic drift of the CFBE41o- cells during
passages. The luminescence HRP assay significantly simplified the CFTR detection as compared to HA-tagged variants (Okiyoneda et al., 2010), and increased signal-to-noise ratio, reproducibility and dynamic range.

The screens identified small molecules that functioned as correctors when used individually, and, when used together with VX-809, had greater efficacy than maximal VX-809 alone. Several classes of compounds were identified that produced a greater than 140% increase in HRP luminescence when added with VX-809 in transfected CFBE41o- cells. These compounds were verified by independent biochemical assays, either from accumulation of complex-glycosylated ΔF508-CFTR-3HA by immunoblot (Supplemental Fig. 5) or cell surface ELISA in transfected CFBE41o- cells, thus confirming the use of HRP-tagged CFTR as valid screening tool for modulators of ΔF508-CFTR biogenesis. Most of the compounds were also active in a secondary functional assay done in ΔF508-CFTR-transfected A549 cells, albeit with relative lower activity. The most active correctors in A549 cells were class D and H correctors, having low micromolar potency. In primary human bronchial cell cultures from a ΔF508 homozygous CF patient, most compounds, including the dihydrospiro-indene H-01, showed little activity. Cell-specific corrector activity has been described (Pedemonte et al., 2010), although the mechanisms responsible are not known. Of note, the 2-aminothiazole D-01 acted in synergy with VX-809 in the ΔF508 human bronchial epithelia, increasing chloride current greater than VX-809 alone. The substantial increase seen with forskolin alone in the D-01-treated cells suggests that the D-01/VX-809 corrector combination is able to partially correct the ΔF508-CFTR folding defect. Mechanistic studies suggested that D-01 is unable to stabilize the NBD1 thermodynamically, but likely targets the NBD2 or its interface with NBD1 or MSDs, as the presence of the NBD2 is required for the D-01 rescue effect and the domain deletion prevented the rescue of ΔF508-1218X-CFTR.

There are several prior reports on biological properties of the corrector scaffolds identified in this study. We previously reported 2-aminoarylthiazole ΔF508-CFTR correctors that are structurally similar to the 2-aminothiazoles (class D) identified here (Pedemonte et al., 2005a). Mechanism of action studies suggested that 2-aminoarylthiazoles improved ΔF508-CFTR folding at the ER and stability at the cell surface (Loo et al., 2008). Recent reported biological activities of aminothiazoles includes inhibition of
prion replication (Gallardo-Godoy et al., 2012), antimicrobial activity against methicillin-resistant
Staphylococcus aureus (Annadurai et al., 2012), and γ-secretase modulators for treatment of Alzheimer’s
disease (Lübbers et al., 2011). Similar dihydrospiro-indenes (class H) have been reported as inhibitors of
Enterococcus faecalis and Staphylococcus aureus phenylalanyl-tRNA synthetases with low nanomolar
potency (Yu et al., 2004). Dihyrospiro-indenes have also been reported to be inhibitors of human
papillomavirus type 11 (HPV11) E1-E2 protein-protein interaction (Goudreau et al., 2007). To our
knowledge, the channel-modulating effects of dihydrospiro-indene have not been reported.

In summary, the results here provide proof-of-concept for the paradigm of synergy-based screening
to identify corrector combinations with greater efficacy than individually used correctors, and support
the idea that the ΔF508 mutation confers multiple structural defects in the CFTR chloride channel.
While the compounds identified here produced only a modest increase in maximal correction efficacy,
their activity and synergy with VX-809 in different human lung epithelial cell lines, including primary
human bronchial cell cultures, supports further synergy screens to identify efficacious corrector
combinations.
ACKNOWLEDGMENTS

We thank Dr. Luis Galietta (Genoa, Italy) for providing transfected A549 cells and Dr. Dieter Gruenert (UCSF, San Francisco) for the parental CFBE41o- cell line.
AUTHORSHIP CONTRIBUTIONS

Participated in research design: Phuan, Veit, Lukacs, and Verkman.

Conducted experiments: Phuan, Veit, Tan, and Roldan.

Wrote or contributed to the writing of the manuscript: Phuan, Veit, Roldan, Finkbeiner, Lukacs, and Verkman.
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FIGURE LEGENDS

Figure 1. **High-throughput synergy screens for identification of ΔF508-CFTR correctors.**
A. Screening assay for ΔF508-CFTR-HRP CFBE41o cells (top) showing incubation with 25 μM test compounds and 2 μM VX-809 for 24 h at 37 ºC. Cell surface ΔF508-CFTR was assayed by a luminescence readout of HRP activity. Concentration-dependence data (bottom left) and bar graph for luminescence readout of VX-809 effect (S.E., n = 4) at 27 and 37 ºC. B. Screening assay for R1070W-ΔF508-CFTR-HRP CFBE41o cells (top) showing incubation with 25 μM test compounds for 24 h at 37 ºC. Concentration-dependence data (bottom left) and bar graph for luminescence readout of VX-809 effect (S.E., n = 4) at 27 and 37 ºC.

Figure 2. **Screening results.** A. Summary of primary findings of the ΔF508-CFTR screen. B. Chemical structures of three classes of correctors identified from the screen. C. Summary of primary findings of the R1070W-ΔF508-CFTR screen. D. Chemical structures of six classes of correctors identified from the screen.

Figure 3. **Corrector concentration-dependence studies.** A. Concentration-dependence data of A-01, B-01, H-01 and K-01, with 2 μM VX-809, in ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3). B. Concentration- dependence data of A-01, B-01, D-01, H-01 and K-01 in R1070W-ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3). C. Correlation between PM density of ΔF508-HRP and ΔF508-3HA in CFBE41o- cells treated with correctors (5 and 25 μM, 24 h, 37 ºC) alone or in combination with 3 μM VX-809. Data were fitted by linear regression analysis with R² = 0.94 (S.E., n = 3).

Figure 4. **Structure-activity relationships of class D and H compounds.** A. Concentration-dependence data of class D analogs in R1070W-ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3). B. Structural determinants of corrector activity of class D compounds. C. Concentration-dependence data of class H analogs, together with 2 μM VX-809, in ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3). D. Structural determinants of corrector activity of class H compounds. Control-(DMSO) background corrected
luminescence signals.

Figure 5. **Functional assay in A549 cells expressing ΔF508-CFTR and a halide-sensitive YFP.**
A. Assay schematic. A549 cells were incubated with test compounds, with or without 2 μM VX-809, at 37 °C for 24 h. ΔF508-CFTR function was assayed in a plate reader from the kinetics of YFP fluorescence quenching in response to extracellular iodide addition in the presence of forskolin (20 μM) and genistein (50 μM). B. Representative data curves showing iodide influx at different [D-01] and [H-01], without and with 2 μM VX-809. C. Concentration-dependence data of A-01, B-01, D-01, H-01 and K-01, without and with 2 μM VX-809 (S.E., n = 3). Fitted curves for single-site activation model.

Figure 6. **Corrector mechanism of action.** A. Plasma membrane expression of temperature-rescued (26 °C, 48 h) HRP-CD4TM-ΔF508-NBD1-1S in MDCK cells. Cells were treated with the indicated correctors for 24 h at 26 °C followed by 1 h chase at 37 °C. B. Representative melting curves (left panel) and melting temperature (Tm, right panel) of human ΔF508-NBD1-1S determined by differential scanning fluorimetry. C-01, D-01 or H-01, or the chemical chaperone glycerol, were present during thermal unfolding at the indicated concentrations. C. Relative effect of corrector 4 (10 μM, 24 h, 37 °C) on the PM density of R1070W-HRP CFBE41o- cells treated with A-01 – D-01, H-01, J-01, K-01 or VX-809 at the indicated concentrations. D. PM density of ΔF508-CFTR-3HA and ΔF508-1218X-CFTR-3HA treated with D-01 (5 and 25 μM, 24 h, 37 °C) alone (left panel) or in combination with VX-809 (right panel) in BHK cells. Errors represent means ± SEM of three independent experiments (B, D) or means ± SD of 8 measurements in two independent experiments (A, C). *, p < 0.05; **, p < 0.01 by unpaired t-test.

Figure 7. **Functional assays in primary cultures of human bronchial epithelial cells from a homozygous ΔF508 CF patient.** A. Representative short-circuit current recordings. Cells were incubated at 37 °C for 24 h with DMSO vehicle, 10 μM VX-809, 30 μM D-01 or 10 μM VX-809 + 30 μM D-01. Concentrations were: amiloride, 10 μM; forskolin, 20 μM; genistein, 50 μM; CFTRinh-172, 10...
μM. **B.** Summary of changes in short-circuit current (ΔI_{sc}) produced by forskolin alone and forskolin + genistein from experiments as in A (S.E.; n = 3 cultures each). *, p < 0.05; by unpaired t-test.
Figure 1

A

ΔF508-CFTR + test compound + VX-809
37 °C, 24 h

measure luminescence

B

R1070W-ΔF508-CFTR + test compound
37 °C, 24 h

measure luminescence
ΔF508-CFTR primary screen
50,000 compounds
164 hits (>50% luminescence signal over 2 µM VX-809)
5 confirmed hits
3 compound classes (H, J, K)

R1070W-ΔF508-CFTR primary screen
60,240 compounds
25 hits (>50% luminescence signal over DMSO control)
9 confirmed hits
6 compound classes (A, B, C, D, E, F)

Figure 2
Figure 3
Figure 4

Class D - aminothiazoles

Class H - dihydrospiro-indene

**Active** ring = 

**Inactive** ring = 

**Best** $R^1$ =

**Inactive** $R^1$ =

**Best** $R^2$ =

**Inactive** $R^2$ =
Figure 5
Figure 6
Figure 7
MOLECULAR PHARMACOLOGY

Supplemental Data

Synergy-based small-molecule screen using a human epithelial cell line ΔF508-CFTR correctors that augment VX-809 maximal efficacy

Puay-Wah Phuan, Guido Veit, Joseph Tan, Ariel Roldan, Walter E. Finkbeiner, Gergely Lukacs, and A.S. Verkman

Supplemental Table 1. Corrector activities in ΔF508-HRP and R1070W-HRP CFBE41o- cells measured with and without VX-809.

Supplemental Table 2. Corrector activities of selected class A, D and H analogs, with and without VX-809, in ΔF508-HRP and R1070W-HRP CFBE41o- cells.

Supplemental Table 3. Functional activities of correctors in A549 cells expressing ΔF508-CFTR and halide-sensitive YFP.

Supplemental Figure 1. Plasma membrane (PM) density of HRP-tagged ΔF508-CFTR.

Supplemental Figure 2. Correctors activities in R1070W and DF508 cell lines.

Supplemental Figure 3. Dose-dependent activities D and H analogs.

Supplemental Figure 4. Functional assays in primary cultures of human bronchial epithelial cells from homozygous ΔF508 CF patient.

Supplemental Figure 5. Immunoblot of ΔF508-CFTR.
**Supplemental Table 1.** Corrector activities in ΔF508-HRP and R1070W-HRP CFBE41o- cells measured with and without VX-809

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<th>Compound</th>
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* as percentage of V<sub>max</sub> for VX-809
**Supplemental Table 2.** Corrector activities of selected class A, D and H analogs, with and without VX-809, in ΔF508-HRP and R1070W-HRP CFBE41o- cells

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Supplemental Table 3. Functional activities of correctors in A549 cells expressing ΔF508-CFTR and halide-sensitive YFP.

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* as percentage of Vₘₐₓ for VX-809
Supplemental Figure 1. Plasma membrane (PM) density of HRP-tagged ΔF508-CFTR with the suppressor mutations 3S or R1070W in presence or absence of VX-809 (3 µM, 24 h, 37 ºC) was measured by luminescence in CFBE41o- cells. Data shown as percent of wild-type CFTR PM density (S.E., n = 3).
Supplemental Figure 2. Correctors activities in R1070W and ΔF508 cell lines. A. Activities of correctors in cell lines. B. Dose-response data of class A-01, B-01, H-01 and K-01 in ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3). C. Dose-response data of class A-01, B-01, D-01, H-01 and K-01, synergized with 2 μM VX-809, in R1070W-ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3).
Supplemental Figure 3. Dose-dependent activities D and H analogs. A. Dose-response data of class H analogs without 2 µM VX-809 in ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3). B. Dose-response data of class D analogs, with 2 µM VX-809, in R1070W-ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3). C. Dose-response data of class H analogs, with (left) and without (right) 2 µM VX809, in R1070W-ΔF508-CFTR-HRP CFBE41o- cells (S.E., n = 3).
**Supplemental Figure 4.** Functional assays in primary cultures of human bronchial epithelial cells from a homozygous ΔF508 CF patient. Representative short-circuit current recordings. Cells were incubated at 37 °C for 24 h with DMSO vehicle, 10 µM VX-809, 10 µM VX-809 + 30 µM compounds. Untreated wild-type CFTR from non-CF human bronchial epithelial cells were shown as further control. Concentrations were: amiloride, 10 µM; forskolin, 20 µM; genistein, 50 µM; CFTRinh-172, 10 µM.
Supplemental Figure 5. Immunoblot of ΔF508-CFTR. Effect of the indicated correctors (A-01 - D-01, J-01, K-01 25 µM, H-01 5 µM, 24 h, 37 °C) alone or in combination with 3 µM VX-809 on the expression pattern of ΔF508-CFTR-3HA and R1070W-ΔF508-CFTR-3HA in CFBE41o- cells. CFTR was visualized using anti-HA antibody, anti-Na+/K+-ATPase antibody was used as loading control.