Can two wrongs make a right? F508del-CFTR ion channel rescue by second-site mutations in its transmembrane domains

Deletion of phenylalanine 508 (F508del) in the cystic fibrosis transmembrane conductance regulator (CFTR) anion channel is the most common cause of cystic fibrosis. The F508 residue is located on nucleotide-binding domain 1 (NBD1) in contact with the cystic extensions of the transmembrane helices, in particular intracellular loop 4 (ICL4). To investigate how absence of F508 at this interface impacts the CFTR protein, we carried out a mutagenesis scan of ICL4 by introducing second-site mutations at 11 positions in cis with F508del. Using an image-based fluorescence assay, we measured how each mutation affected membrane proximity and ion-channel function. The scan strongly validated the effectiveness of R1070W at rescuing F508del defects. Molecular dynamics simulations highlighted two features characterizing the ICL4/NBD1 interface of F508del/R1070W-CFTR: flexibility, with frequent transient formation of interdomain hydrogen bonds, and loosely stacked aromatic sidechains (F1068, R1070W, and F1074, mimicking F1068, F508, and F1074 in WT CFTR). F508del-CFTR displayed a distorted aromatic stack, with F1068 displaced toward the space vacated by F508, while in F508del/R1070F-CFTR, which largely retained F508del defects, R1070F could not form hydrogen bonds and the interface was less flexible. Other ICL4 second-site mutations which partially rescued F508del-CFTR included F1068M and F1074M. Methionine side chains allow hydrophobic interactions without the steric rigidity of aromatic rings, possibly conferring flexibility to accommodate the absence of F508 and retain a dynamic interface. These studies highlight how both hydrophobic interactions and conformational flexibility might be important at the ICL4/NBD1 interface, suggesting possible structural underpinnings of F508del-induced dysfunction.

The ATP-binding cassette (ABC) transporter family is a large superfamily of proteins (1, 2). Within this superfamily, the cystic fibrosis transmembrane conductance regulator (CFTR) (3) is the only protein known to function as an ion channel (4). Nevertheless, CFTR shares structural features with other ABC transporters. In particular, CFTR’s two asymmetric nucleotide-binding domains (NBDs), NBD1 and NBD2, interact with two transmembrane domains (TMDs) (TMD1 and TMD2) possessing an almost typical Type IV fold (5). Interactions between the TMDs and NBDs are mediated by two pairs of intracellular loops (ICLs). The units formed by a surface depression in each NBD and the two ICLs that contact it have been described as ‘ball-and-socket joints’ (6). While NBD1 forms a relatively shallow socket that interfaces with ICL1 and ICL4, NBD2 forms a deeper socket which contacts ICL2 and ICL3. ICL2 and ICL4 cross over to the NBD linked to the opposite TMD, forming a domain-swapped arrangement like that found in other Type IV ABC transporters, such as bacterial Sav1866 (7–9), MsbA (10), TM287/288 (11), mammalian P-glycoprotein (12), human ABCB10 (13), and McjD (14). In CFTR, interactions at the NBD–TMD interfaces are responsible for coupling ATP binding and hydrolysis to channel gating by transmission of conformational changes from the NBDs to the TMDs, which form the anion-selective permeation pathway (4).

CFTR plays an important physiological role, controlling epithelial secretions in several organs, such as the airways, intestine, pancreas, biliary ducts, sweat glands, and reproductive tracts (15). Cystic fibrosis (CF), caused by loss of function mutations in the CFTR gene, is the most common life-limiting genetic disease in populations of European descent, affecting one in ~2500 to 3000 newborns (16–18). CF-causing mutations are unequally distributed between the two NBD–TMD interfaces (6): 16 are found at the TMD/NBD1 interface, but only five at the TMD/NBD2 interface (list of CFTR2 variants 31 July 2020, https://info.cfr2.org/mutations-history). Possibly the absence of a short helix in NBD1—present in NBD2 and in the NBDs of other ABC transporters (19)—is responsible for a shallower socket and weaker ICL4/NBD1 interactions, rendering the ICL4/NBD1 interface particularly vulnerable to harmful mutations. Moreover, a more dynamic NBD1 structure around the socket might also contribute to causing this mutation hotspot (20).

By far the most common CF-causing variant is F508del, which deletes a phenylalanine in NBD1 that contributes molecular contacts at the ICL4/NBD1 interface (16, 21, 22). Even though X-ray structures of human F508del NBD1 show that
there are only small local changes in the conformation of the loop comprising residues 507 to 511 (23), the deletion has great impact on the biogenesis and function of CFTR. While WT CFTR becomes complex-glycosylated in the Golgi, the F508del mutant does not undergo any detectable complex glycosylation at 37 °C (24–30). Instead, misfolded F508del-CFTR is trapped in the endoplasmic reticulum, ubiquitinated, and then degraded by the proteasome (31).

The minute amount of F508del-CFTR that escapes to the plasma membrane has decreased membrane stability (32–34) and a severe gating impairment. The latter is characterized by a reduction in open probability ($P_\text{o}$) (30, 35–40) caused by a prolonged closed time interval between bursts. The F508del mutation does not strongly affect CFTR pore properties such as single-channel conductance and anion selectivity, although a reduction in current amplitude can occur once instability develops (reviewed in (41)).

Small-molecule modulators that interact with CFTR directly and rescue channel function (potentiators), and/or folding, processing, and trafficking to the plasma membrane (correctors), have been developed for clinical use (42). In the laboratory, one common strategy to promote F508del-CFTR trafficking to the plasma membrane involves incubating cells at low temperature, 26 to 30 °C (30, 34, 43–45). Low temperature provides an energetically favorable F508del-CFTR folding (30) and proteostasis (44) environment, leading to a decrease in misfolding and improved trafficking to the plasma membrane. Moreover, evidence suggests that low temperature promotes trafficking of immature (core-glycosylated) F508del-CFTR, via a nonconventional trafficking pathway that bypasses the Golgi (45, 46).

Another laboratory strategy for F508del-CFTR rescue is to introduce second-site (revertant) mutations in cis with F508del. Identified revertant mutations located in NBD1 include V510D/E/A (47–49), I539T (50, 51), G550E (50), R553M/Q (52) and R555K (51, 53). The latter mutations reduce F508del-NBD1 thermodynamic and kinetic instability (54, 55). By contrast, R1070W, located in ICL4 (54, 56, 57), acts by restoring interactions at the ICL4/NBD1 interface (56, 58).

Here, we investigated F508del-CFTR rescue by second-site mutations. We systematically scanned positions 1064 to 1074 at low temperature, 26 to 30 °C (30, 34, 43–45). Low temperature provides an energetically favorable F508del-CFTR folding (30) and proteostasis (44) environment, leading to a decrease in misfolding and improved trafficking to the plasma membrane. Moreover, evidence suggests that low temperature promotes trafficking of immature (core-glycosylated) F508del-CFTR, via a nonconventional trafficking pathway that bypasses the Golgi (45, 46).

Second-site mutations in cis with F508del affect the CFTR anion channel, yellow fluorescent protein (YFP)(H148Q/I152L)-CFTR fluorescence quenching in response to extracellular $\Gamma$ addition was quantified following expression in HEK293 cells. A 24 h incubation at 28 °C was carried out to minimize misfolding and optimize trafficking, and the low temperature was maintained throughout image acquisition. Whole-cell conductance ($G$) and membrane potential ($V_\text{m}$) at steady state, immediately preceding $\Gamma$ addition, were estimated by fitting of a mathematical model to the quenching time course. Moreover, we simultaneously measured CFTR membrane proximity by quantifying the YFP(H148Q/I152L)-CFTR fluorescence located at the border of cells, using mCherry fluorescence as an internal standard for comparison (61). Table S1 summarizes the assay readouts.

Whole-cell conductance

In the control (dimethyl sulfoxide [DMSO]) condition, mutants with the F508del background typically had an average G estimate of around 1 nS ($M = 0.97$ nS, $SD = 1.29$, $N = 64$), consistent with a small anion permeability reflecting endogenous, non-CFTR–mediated conductance and minimal basal phosphorylation of CFTR. Following CFTR activation (through the cAMP pathway) by 10 μM forskolin (Fig. 1A), the anion conductance of WT CFTR increased significantly (forskolin: $Mdn = 117.70$ nS, $N = 17$; DMSO: $Mdn = 2.35$ nS, $N = 20$; see one-tailed Wilcoxon ranking tests in Table S2). For F508del-CFTR too, there was a modest, albeit significant, increase in $G$ after addition of forskolin ($Mdn = 5.7$ nS, $N = 18$) compared to the control condition ($Mdn = 0.86$ nS, $N = 19$). An increased conductance after addition of forskolin compared to DMSO was observed in 24 of 61 mutants with second-site mutations in cis with F508del at the ICL4/NBD1 interface (Fig. 1A and Table S2). Fluorescence quenching timelines are shown in Figure S3.

Table S4 demonstrates that many second-site mutations further impaired F508del-CFTR function—especially substitutions at sites T1064, L1065, R1066, A1067, G1069, and Q1071. By contrast, eight mutations significantly increased the F508del-CFTR–mediated, forskolin-stimulated conductance (Wilcoxon Rank Sum tests, Table S4). Among these, R1070W was particularly effective, increasing F508del-CFTR conductance to 42% of the value measured for WT CFTR ($Mdn = 49.76$ nS, $N = 5$). As a comparison, chronic treatment of...
F508del-CFTR expressing HEK293 cells with the clinically approved CFTR corrector lumacaftor (3 μM VX-809 for 24 h at 28°C) resulted in a G of only 12% of that measured for WT CFTR ($\text{Mdn} = 13.73$ nS, $N = 7$, data not shown). After R1070W, the most successful revertant mutations were F1068M and F1074M, followed by A141S and R1097T. Finally, R1070Q, F1068Q, and R1070F mutations all gave smaller, but still significant, improvements in G (Table S4).

Membrane potential

More than half of the HEK293 cells expressing second-site mutations resulted in a significantly depolarized $V_m$ after steady-state activation with 10 μM forskolin when compared to those expressing F508del-CFTR (Wilcoxon Rank Sum tests, Table S4). However, for most of these cells, a relatively depolarized $V_m$ was also present under control conditions. A significant depolarization of $V_m$ resulting from activation of CFTR with forskolin was seen only for WT CFTR, F508del/R1070W, F508del/R1070Y, and F508del/F1068M (one-tailed Wilcoxon ranking tests (Table S2)).

Membrane proximity

Watershed-based image segmentation on the mCherry images allowed us to approximate the location of the plasma membrane without relying on efficient YFP(H148Q/I152L)-CFTR membrane trafficking. For each cell, the amount of CFTR in proximity to the membrane, denoted as $\rho$, was defined as the ratio of the average normalized YFP(H148Q/I152L)-CFTR fluorescence intensity within the membrane proximal zone (a ~1 μm wide band adjacent to the cell boundary) to the total membrane fluorescence intensity. For each cell, $\rho$ was calculated as the average of all pixels within the membrane proximal zone.

Figure 1. Effects of second-site mutations in cis with F508del on CFTR conductance, membrane potential, and CFTR membrane proximity. A, CFTR conductance (G in nS, upper panel) and membrane potential ($V_m$ in mV, lower panel), measured in HEK293 cells expressing WT CFTR, F508del-CFTR, or F508del-CFTR with second-site mutations. The dashed reference lines indicate the median conductance (upper panel) and membrane potential (lower panel) after steady-state activation with forskolin of WT CFTR (blue) and F508del-CFTR (red). Asterisks show results of one-tailed Wilcoxon ranking tests used to compare G and $V_m$ after addition of DMSO (control; gray boxes) versus 10 μM forskolin (colored boxes). B, measurements of CFTR membrane proximity ($\rho$) obtained from the same CFTR-expressing HEK293 cells described in A. Dashed reference lines indicate the average CFTR membrane proximity for cells expressing WT CFTR (blue) and F508del-CFTR (red). Mean log$_{10}$ρ values were paired per plate, and paired t-tests were performed on the mean log$_{10}$ρ measurements of cells expressing F508del-CFTR with (colored bars) versus without (gray bars) second-site mutation. CFTR, cystic fibrosis transmembrane conductance regulator; G, whole-cell conductance.
boundary) to the average normalized mCherry fluorescence intensity throughout the entire cell (ρ = f_{YFP membrane}/f_{mCherry cell}). This metric informs about first, trafficking (the fraction of YFP(H148Q/I152L)-CFTR reaching the membrane, f_{YFP membrane}/f_{YFP cell}) and second, the fusion protein’s metabolic stability with respect to mCherry’s (quantified by the ratio f_{YFP cell}/f_{mCherry cell}), a function of overall rates of biosynthesis and degradation. The ρ measurements approximated a lognormal distribution and were log_{10} transformed before determining plate means for each mutation. To evaluate the effects of second-site mutations on F508del-CFTR delivery to the plasma membrane, we compared the mean log_{10} measurements of cells expressing F508del-CFTR in the absence and presence of second-site mutations (Fig. 1B and Table S5). Only R1070W, R1070M, and F1074M significantly increased F508del-CFTR membrane proximity, whereas it was significantly reduced by 25 substitutions (Fig. 1B and Table S5).

**Gating and conduction properties**

CFTR conductance increases approximately linearly with membrane proximity (61), consistent with the ρ metric being proportional to the number of channels at the plasma membrane. Because G is the product of the number of channels at the membrane (N), open probability (P_o), and single-channel conductance (γ), our two assay readouts allow evaluation of the gating and conduction properties (P_o ∙ γ) of CFTR channels located at the plasma membrane (see (61)). The dotted lines in Figure 2 describe conductance as a function of membrane proximity assuming single-channel properties (P_o ∙ γ) of WT CFTR after steady-state activation with forskolin, with the specific experimental conditions used (see Fig. S6). Data points falling above or below the line are suggestive of single-channel activity higher or lower, respectively, than those of WT CFTR.

F1068M (orange five-point star, Fig. 2A) improves the G of F508del-CFTR. The rightward (nonsignificant) shift on the ρ-axis positions the point on the regression line, suggesting that P_o ∙ γ characteristics of F508del/F1068M-CFTR might be close to those of WT CFTR. F508del/R1070M (orange five-point star, Fig. 2B) is significantly shifted to the right compared to F508del-CFTR, suggestive of improved biogenesis, trafficking, and/or membrane stability. However, the mutant falls far below the regression line, suggesting that even though the number of channels at the plasma membrane has increased, P_o ∙ γ is still much lower than that of WT CFTR. By contrast, F508del/R1070W (red circle, Fig. 2B) falls above the regression line, consistent with published single-channel recordings demonstrating rescue of P_o to WT values (while γ is unchanged in both F508del-CFTR and F508del/R1070W, (63)). All substitutions at site F1074, except for methionine (M; orange 5-point star, Fig. 2C), decreased conductance, such that the forskolin-dependent increase in G after activation seen in F508del-CFTR is lost (Table S2). By contrast, the F1074M substitution not only significantly increased conductance but also was one of the three mutants that significantly increased the membrane proximity of F508del-CFTR. Part of the

![Figure 2. Plate matched G-ρ measurements.](image)

G was plotted as a function of membrane proximity (ρ, obtained by back transformation of mean log_10(ρ)). The panels show G-ρ plots for F508del-CFTR in the absence and presence of second-site mutations at residues F1068 (A), R1070 (B), F1074 (C) and identified in the Yorlp screen (A141S and R1097T, D). Markers and error bars represent the mean G ± SEM (y-axis) and back transformed mean ± upper and lower limits of the SEM (x-axis). CFTR, cystic fibrosis transmembrane conductance regulator; G, whole-cell conductance.

**MD simulations**

We used MD simulations to investigate whether efficacy of rescue was correlated with restoration of the interactions between ICL4 and NBD1 found in WT CFTR. For comparison to WT CFTR, we used previously published simulations of the WT zebrafish CFTR (zCFTR) (62), whose sequences at the ICL4/NBD1 interface differ from those of human CFTR (hCFTR) at only six positions in the NBD1 loop including F508 (positions 495–512, human numbering) and eight in ICL4 (positions 1050–1080, Table S7). In subsequent sections, we use the numbering of residues in hCFTR to refer to the positions of residues in both hCFTR and zCFTR (e.g., zR1070 indicates the zCFTR residue corresponding to R1070 in hCFTR, i.e., R1078, see Table S7 and Fig. S8). We selected for analysis two ICL4 second-site mutations, which rescue F508del-CFTR with different efficacy: R1070W that improved markedly both biogenesis and conductance and R1070F, which has only minor effects on conductance.

We constructed a model for the F508del-CFTR system by replacing NBD1 in the ATP-bound structure of hCFTR (64) with the experimental structure of F508del-NBD1 (65). The
resulting system is hereafter termed F508del/R1070. We then generated two additional F508del systems by replacing R1070 with (i) a tryptophan residue (F508del/R1070W) and (ii) a phenylalanine residue (F508del/R1070F). Each system was embedded in a 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC) lipid bilayer and simulated for 2 μs. The ICL4/F508del-NBD1 interfaces were compared with the ICL2/NBD2 interface of the same systems, as well as with previously published, ATP-bound and ATP-free simulations of the WT zCFTR (6, 62, 66).

In our MD simulations, we assume that the conformations sampled by the mutant proteins, which reach the plasma membrane are not grossly different from those adopted by WT CFTR. Although mutation effects during biogenesis might result in proteins that fold differently, these effects are minimized by low temperature incubation. Moreover, the significant increase in conductance following stimulation by forskolin (Table S2) for these three mutants suggests that a large proportion of the mutant channels at the plasma membrane retain regulation of gating by cAMP-dependent phosphorylation.

RMSD-based cluster analysis

First, we investigated whether the presence of aromatic residues at the R1070 position alters the structure of the F508del-NBD1 loop. To address this aim, we analyzed the conformation of the F508del-NBD1 loop by means of a RMSD-based cluster analysis of the ICL4/NBD1 interface (ICL4 residues 1050–1080 and F508del-NBD1 loop residues 495–512). With a RMSD cut-off of 0.1 nm, we detected (i) 39 clusters for the F508del/R1070 system, with the first cluster and the top three clusters (Fig. 3, A–C) representing approximately 43% and 69%, respectively, of the total structures; (ii) 35 clusters for the F508del/R1070W system, with the first cluster and top three clusters (Fig. 3, D–F) representing approximately 48% and 73%, respectively, of the total structures; and (iii) nine clusters for the F508del/R1070F system, with the first cluster alone representing approximately 89% of the total structures (Fig. 3G). The same analysis was performed on datasets from simulations on WT zCFTR (Fig. 3, H and I), with cluster 1 representing 93% and 74% of the total structures from the ATP-bound and ATP-free simulations, respectively. Figure 3 demonstrates that the NBD1 loop in the F508del/R1070 and F508del/R1070W mutants explored more extended conformations following the helical part (residues 502–507) of the loop. This can be seen for cluster 1 of the F508del/R1070 system (Fig. 3A) and for the first two clusters of the F508del/R1070W system (Fig. 3, D and E). More compact conformations, similar to the WT hCFTR NBD1 loop conformation (Fig. S8), were retrieved for (i) clusters 2 and 3 of the F508del/R1070 system (Fig. 3, B and C); (ii) cluster 3 of the F508del/R1070W system (Fig. 3F), and (iii) the F508del/R1070F system (Fig. 3G). These folded conformations were also retrieved in the most populated cluster of the ATP-bound and ATP-free zCFTR systems (Fig. 3, H and I).

Aromatic residue side chain orientations

The interface between ICL4 and the NBD1 loop near R1070 is characterized by the presence of several aromatic residues, including F1068 and F1074 from ICL4 and F508 from NBD1. Figure 3, A–G shows the side chain orientation of F1068 (dark gray), R1070X (light gray), and F1074 (light blue) of all the members of a given cluster. We considered the orientation of these interfacing residues by measuring the first rotamer (χ1) of

![Figure 3. Dynamics of the ICL4/F508del-NBD1 interface. Results of the cluster analysis for the following: A–C, F508del/R1070 simulation system; D–F, the F508del/R1070W system; G, the F508del/R1070F system; H, ATP-bound; I, ATP-free zCFTR systems previously simulated (62). For the F508del/R1070 and F508del/R1070W systems, the most populated clusters are shown, while for F508del/R1070F and WT zCFTR, only the first cluster is shown. For each cluster, the left panel is the front view of the interface, with the members of the cluster shown in transparent cartoons. The middle and right panels show the cluster center as cartoon and the side chains of F1068 (black), R1070X (gray), F1074 (light blue), and F508 (yellow) of all the members of the cluster, as lines. CFTR, cystic fibrosis transmembrane conductance regulator; ICL4, intracellular loop 4; NBD1, nucleotide-binding domain 1; zCFTR, zebrafish CFTR.](image-url)
their side chains over the entire simulation time (Fig. 4, A and C). $\chi_1$ for F1068 shows a peak near 300°, corresponding to the side chain orientation of the most populated clusters for F508del/R1070F, F508del/R1070W, and WT zCFTR, (“downwards” side chain orientation shown in Fig. 3, D–I). By contrast, for the F508del/R1070 system, the main peak is at 180°, reflecting the “upwards” movement of the side chain toward the NBD1 loop, as for the most populated clusters in Figure 3, A–C. For the aromatic residues at the 1070 position, R1070W shows a bimodal distribution (middle panel, Fig. 4A), with the major peak near 280 to 300° and a smaller one at approximately 200 to 210°. The major peak corresponds to the side chain orientation shown in Figure 3, D and F, and is shared with R1070F (Fig. 3G), while the smaller peak corresponds primarily to structures that form the second most populated cluster of F508del/R1070W, with the tryptophan side chain flipped upwards (Fig. 3E). No significant differences were retrieved across the three F508del and the two WT zCFTR systems for the $\chi_1$ distribution of F1074. At the ICL2/NBD2 interface, Y275 corresponds to the F1068 position in ICL4 (Table S7), and its $\chi_1$ distribution is centered near 300° (Fig. 4, B and D), similar to F1068 in F508del/R1070W and F508del/R1070F and WT zCFTR. W277 at the ICL2/NBD2 interface corresponds to R1070 in ICL4 (Table S7), and its $\chi_1$ distribution shows a peak near 300° in all simulation systems, as for R1070F and the major peak of R1070W.

In summary, with the exception of F1068 in F508del/R1070, the $\chi_1$ angle of the selected residues at the interface corresponds to the distributions obtained from previous simulations (62) on the WT zCFTR structure (Fig. 4, A and B versus Fig. 4, C and D).

We also analyzed two other residues, W496 of the NBD1 loop and Y1073 in ICL4, that contribute to the aromatic nature of the interface and are located in proximity to the amino acids discussed above (Fig. S9). In the three F508del systems, the W496 side chain is oriented upwards and not toward the residues at positions 1068, 1070, and 1074 (Fig. S9, A–C), with its $\chi_1$ distribution centered at 180° (Fig. S9F). This orientation is shared with the WT zCFTR systems (Fig. S9, D–E and G). In the WT zCFTR systems, the W496 side chain is more dynamic and can adopt an alternative conformation, corresponding to the additional peak centered at approximately 75°, especially in the ATP-bound system (Fig. S9G). This orientation corresponds to a side chain pointing toward zF508. However, this alternative conformation of W496 does not affect the orientation of the residues at positions zF1068 and zF1074, as shown by their $\chi_1$ distribution (Fig. 4C). Y1073 is located next to F1074 and above the residue at position 1070, and its side chain adopts similar conformations in the F508del and the WT zCFTR systems, pointing toward the water environment (Fig. S9, A–G).

Spatial overlap between F508 in WT CFTR and side chains at position 1070

To test for overlap of R1070F and R1070W with the region occupied by F508 in WT hCFTR, the ICL4/NBD1 interface from the hCFTR structure was superimposed on the center of cluster 1 for the F508del/R1070F system (Fig. S10A) and those of the three most populated clusters of the F508del/R1070W system (Fig. S10, B–D). While R1070F projected toward the space vacated by F508 (Fig. S10A), the overlap was greater for the F508del/R1070W system, particularly cluster 1 (Fig. S10B). We also compared the orientation of R1070F and R1070W with that of W277 at the ICL2/NBD2 interface (Fig. S10, E–H). Similar results were obtained with this comparison; a higher

![Figure 4. Aromatic side chain orientations at TMD/NBD interfaces.](Image 46x95 to 547x322)
degree of overlap was achieved with the R1070W mutant (Fig. S10F versus Fig. S10E).

**Hydrogen-bond interactions at the ICL4/F508del-NBD1 interface**

In our previous WT zCFTR simulations (62), we observed that zR1070 was highly dynamic, oriented both toward and away from the interface, and forming transient interactions with zF508 and other residues of the NBD1 loop (Fig. 5A and Table S7). We therefore investigated whether equivalent interactions were present in the F508del systems. We found that R1070 and R1070W formed hydrogen bonds with residues E504, I507, and G509 of the NBD1 loop. Interactions also occurred between Y1073 and E504 (Fig. 5B). Overall, hydrogen-bond interactions in the F508del systems, while not lasting for the entire simulation time, were more frequent and persistent than those in the WT zCFTR system.

**Discussion**

Our systematic, empirical scan of ICL4, seeking mutations capable of repairing F508del defects, highlighted a number of second-site revertant mutations. The replacement of R1070 with a tryptophan (54, 56, 57) was found to be most effective at increasing plasma membrane levels and function of F508del-CFTR. This confirms patch-clamp recordings, which show how the R1070W mutation restores F508del-CFTR plasma membrane stability and gating kinetics to levels measured for WT CFTR (63). Using MD simulations, the structural basis of this revertant mutation’s action was investigated. Our systematic scan also identified a small number of new revertant mutations, which mitigated the defects of F508del to some degree. Taken together, our results underscore the importance, for CFTR biogenesis and function, of a dynamic ICL4/NBD1 interface characterized by multiple transient hydrophobic and hydrogen-bond contacts.

**F670del-Yor1p revertants**

We introduced CFTR equivalents of the F670del-Yor1p revertant mutations F270S (67) and R1116T (60) into F508del-CFTR. In hCFTR, the corresponding revertant mutations are A141S (in transmembrane helix 2, TM2) and R1097T (in TM11), respectively. Both revertant mutations, situated on adjacent TMs at the same horizontal plane within the membrane-embedded portion of the protein, significantly rescued F508del-CFTR activity. Structurally, the two residues are coupled to the TMD/NBD1 ball-and-socket joint, through helical portions of TM2 (linking to ICL1) and TM11 (linking to ICL4) (68). F270S only rescued the folding and trafficking of F670del-Yor1p in combination with another revertant mutation, R1168M, in TM12 (67). By contrast, A141S, by itself, significantly rescued the conductance of F508del-CFTR, albeit it was without effect on its membrane proximity, suggesting an effect on $P_o$ and/or $\gamma$. Unlike F270S, R1116T, by itself, increased F670del-Yor1p function. Our data show that R1097T too increased F508del-CFTR conductance by improving its gating and/or conduction properties. While we cannot completely rule out an effect on conduction, residues in TM2 and TM11 do not play a major role in lining the inner vestibule of the pore (69) (but note (70)), suggesting effects on $P_o$ are more likely. It is possible that in F670del-Yor1p, as in...
**F508del-CFTR ion channel rescue by revertant mutations**

F508del-CFTR, the enzymatic cycle time is greatly prolonged by a slow transition from the inward-facing to outward-facing conformation (corresponding to opening in CFTR, (71–73)). R1116 and R1097 are linked, via TM11, to the TMD/NBD1 interface altered by the phenylalanine deletion. The threonine substitutions might facilitate coupling of NBD dimerization to the TMD rearrangement resulting in an outward-facing conformation. Further studies are required to better understand the mechanistic details of this rescue.

**Why is R1070W such an effective revertant mutation?**

Introducing the R1070W mutation into the F508del-CFTR background greatly increased the CFTR-mediated anion conductance. In an attempt to identify the structural basis of this remarkably efficient rescue, we compared three different MD simulations of F508del systems (F508del/R1070, F508del/R1070W, and F508del/R1070F) with simulations previously performed on the WT zCFTR systems (62). While other MD simulations have analyzed the NBD dimer interface (74), we focused on the ICL4/NBD1 interface. Two potentially important features of protein dynamics emerged, suggesting that both the indole nitrogen and the aromatic ring in R1070W play important roles (Fig. 6).

_F508del/R1070W restores ICL4/NBD1 aromatic interactions_

In WT CFTR, the interface between ICL4 and NBD1 is characterized by the presence of several aromatic residues. In MD simulations of WT zCFTR, we found that the side chains of zF1068, zF508, and zF1074 loosely stack in an alternating fashion (Fig. 3, H and I). In the MD simulation of F508del/R1070, the arrangement of aromatic side chains at the ICL4/NBD1 interface deviated noticeably from this pattern, due to the upward movement of F1068 toward the region vacated by the deletion of F508. Our MD simulations of F508del/R1070W and F508del/R1070F showed that both R1070W and R1070F side chains occupy the region of space sampled by F508 in WT zCFTR (Fig. 3, D–G). As a result, the loose stacking of aromatic residues in WT zCFTR and hCFTR is restored. The bulkier side chain of R1070W permitted better overlap with the region occupied by F508, particularly for the most populated cluster of F508del/R1070W, comprising 48% of the total conformations (Fig. S10). It is interesting to note that at the homologous ICL2/NBD2 interface, a tryptophan, W277 (ICL2), corresponds to R1070 and projects its side chain toward the NBD2 loop (positions 1303–1309), where a proline (P1306) is present at a position equivalent to F508. A tryptophan–proline pair is completely conserved at these positions at the ICL2/NBD2 interface among a set of asymmetric ABC transporter sequences (75). Similarly, an aromatic side chain is completely conserved at a position equivalent to F508 at the ICL4/NBD1 interface. However, a basic side chain at a position equivalent to R1070 is much less conserved (75).

_F508del/R1070W restores ICL4/NBD1 hydrogen-bond interactions, allowing flexibility of the NBD1 loop_

In WT zCFTR, the NBD1 loop mostly adopts a compact, folded structure, but several residues in ICL4, including zR1070, are highly dynamic and form transient hydrogen bonds with residues in the NBD1 loop, including zF508 (Fig. 5A, see also (22, 76, 77)). Our MD simulations of the three F508del systems highlighted a clear dichotomy. F508del/R1070 and F508del/R1070W had a more flexible NBD1 loop, sampling more extended conformations and forming transient hydrogen bonds with residues of ICL4. By contrast, F508del/R1070F adopted a more rigid interface, with no groups at the 1070 position capable of establishing ICL4/NBD1 hydrogen bonds (Fig. 5B). R1070W oriented toward the interface occupying the region sampled by zR1070 in WT zCFTR, formed similar hydrogen bond contacts, albeit more frequently (Fig. 5A versus Fig. 5B). R1070F adopted a similar orientation. However, because hydrophilic contacts are not possible for the phenylalanine side chain, the ICL4/NBD1 interface in this mutant is fixed and cannot sample alternative conformations with similar stability. This is demonstrated by the absence of significant changes in the NBD1-loop conformation throughout the simulation (the top cluster, Fig. 3G, includes 89% of all structures). Functionally, a dynamic ICL4/NBD1 interface is crucial for CFTR gating, as demonstrated by the rapid and reversible interruption of channel activity that occurs upon formation of covalent cross-links between F508C and F1068C in a cyst-less CFTR background (21). In addition, recent studies suggest that a frequent “uncoupling” of NBD1 is an integral feature of WT CFTR gating (20, 78). A tryptophan side chain at position R1070, capable of both hydrogen bonding and hydrophobic contacts, would allow the protein to adopt conformations in which the interface is alternatively buried or more solvent exposed.

_Methionine substitutions at positions F1068 and F1074_

Following R1070W, the most effective F508del-CFTR revertant mutations are F1068M and F1074M. These results are consistent with a crucial role played by the aromatic cluster at
the core of the ICL4/NBD1 interface (including F1068, F508, and F1074) highlighted by our MD simulations and analysis (Figs. 3 and 4). Methionine-aromatic interactions can occur at longer distances and are potentially more robust compared to purely hydrophobic contacts or salt bridges (79). The linear (nonbranched) aliphatic chain that includes a sulfur atom gives more flexibility than any other hydrophobic side chain (80), possibly allowing strong interactions with the aromatic cluster, despite the distortions caused by the absence of F508.

In addition, F1068M and F1074M might allow transient polar interactions (81, 82). Strikingly, one of the critical phenylalanines coordinating permeating anions in the Fluc-Ec2 channel can only be functionally replaced by methionine (83). We speculate that rescue by the introduced methionines in CFTR might rely on a dynamic switching between relatively strong hydrophobic interactions and interactions with polar residues of the NBD1 loop or with the solvent.

Potential translational impact of this study

The results of the mutagenic scan presented here advance our understanding of the defect caused by absence of phenylalanine 508, the variant carried by most people with CF of European descent. As our understanding of the structure and MD of the CFTR protein improves, information on the detailed impact of the deletion will inform efforts to design future therapies capable of optimizing the rescue of CFTR biogenesis and function.

The strong validation our data provides of the effectiveness of the revertant mutation R1070W might invigorate efforts to design novel gene therapy treatments for CF. Partial restoration of function might be achieved by cytosine base editing (transforming the arginine-encoding codon CGG to the tryptophan-encoding TGG). Base editing is more efficient and less prone to error than homology-directed repair ((84, 85) and PT Harrison, personal communication). In addition, R1070W rescue of F508del-CFTR channels is likely to have synergistic effects with Class I and Class III correctors (58, 86) potentially allowing a simplification of modulator therapy. Thus, base-editing gene therapy might lead to benefits for F508del homozygous patients, in the interim, while the homology-directed repair gene therapy technology is advanced.

Experimental procedures

Plasmid and site-directed mutagenesis

Mutations were introduced in the pIRES2-mCherry-YFPCFTR plasmid (see (61)) with help of complementary primers containing mutations (Eurofins MWG Operon) using the QuikChange protocol for site-directed mutagenesis (Stratagene). Sanger sequencing either outsourced to SourceBio-science or by the UCL Sequencing Facility with a 3100-Avant Genetic Analyzer (Applied Biosystems) was used to confirm the introduced mutations.

Cell culture, transfection, and incubations

HEK293 cells were maintained at 37 °C in Dulbecco’s modified Eagle’s medium, supplemented with 2 mM L-glutamine, 100 units/ml penicillin, 100 μg/ml streptomycin, and 10% fetal bovine serum (all from Life Technologies, Inc). Cells were seeded in black-walled 96-well plates (Costar, Fisher Scientific) coated with poly-D-lysine and transiently transfected with the pIRES2-mCherry-YFPCFTR plasmid using Lipofectamine 2000 (Life Technologies), following the manufacturer’s instructions. After transfection, cell plates were incubated at 37 °C for 24 h, then at 28 °C for a further 24 h to minimize misfolding (30). Before imaging, cells were washed twice in 100 μl of standard buffer (140 mM NaCl, 4.7 mM KCl, 1.2 mM MgCl2, 5 mM Hepes, 2.5 mM CaCl2, and 1 mM glucose, pH 7.4).

Image acquisition

The ImageXpress Micro XLS (Molecular Devices), an automated inverted wide-field fluorescence microscope with a temperature-controlled chamber (set to 28 °C), was used for image acquisition. Protocols for automated imaging and fluid additions were created using MetaXpress software (Molecular Devices). A 20 × objective was used to take 16 bit images of both mCherry (excitation/emission filters at 531 ± 20/592 ± 20 nm) and YFP-CFTR (excitation/emission filters at 472 ± 30/520 ± 35 nm). To evaluate CFTR activity at steady state, images of mCherry and YFP fluorescence were taken every 2 s. After following the baseline for 20 s, CFTR was activated by the addition of 50 μl of standard buffer containing forskolin (10 μM final concentration) or DMSO (control, 0.05 %). After a further 230 s, when CFTR is expected to be gating at steady state, 50 μl of Γ- buffer (as standard buffer with 400 mM NaI instead of 140 mM NaCl) were added to achieve the extracellular Γ- concentration of 100 mM. Further, forskolin/DMSO were added so that their concentration was not altered by the second fluid addition. After this, image acquisition continued for another 40 s.

Image analysis

Images were analyzed using MATLAB mathematical computing software (MathWorks) as described (61). In brief, to estimate CFTR membrane proximity, a watershed transform-based segmentation was performed on binarized mCherry images after noise was removed. Cells were removed from analysis if they had (i) an area of <108 μm2 or >5400 μm2, (ii) a major axis length of >32.4 μm, (iii) if the area over the perimeter was <25 or >300 μm, or (iv) if they were touching the image border. The membrane-proximal zone was defined as a 1.08 μm band within the border of each cell. After background correction, YFP and mCherry fluorescence intensity were normalized to the median YFP and mCherry fluorescence intensities of cells expressing WT CFTR on the same plate. Cells were removed from analysis if their average normalized fluorescence intensity fell below 0. For each cell, the CFTR membrane proximity (ρ) was defined as the average normalized YFP fluorescence intensity within the membrane-proximal zone over the average normalized mCherry fluorescence within the entire cell.
**F508del-CFTR ion channel rescue by revertant mutations**

A separate image analysis protocol was used to assess CFTR activity. CFTR was activated with 10 μM forskolin and allowed to reach steady state, after which 100 mM extracellular Γ was added. Images were corrected for background noise with help of a cell selection mask created using binarized mCherry images taken at the first and last timepoints. A mathematical model was then fitted to the cell YFP fluorescence time course (see below).

**mCherry fluorescence quality control**

The mCherry fluorescence is related to the amount of bicistronic mRNA transcribed from the plasmid. The normalized mCherry fluorescence was quantified using images collected with the steady-state activity and membrane proximity protocols (see Fig. S11 and Table S12). In both datasets, mCherry fluorescence intensity of cells expressing F508del/Q1071F-CFTR was significantly lower compared to other variants. This difference persisted using a new DNA preparation of the plasmid. For this reason, the F508del/Q1071F mutant was not studied further and removed from the dataset.

**Mathematical model**

The steady-state membrane potential (V_m in mV) and the CFTR-mediated whole-cell Cl− conductance (G in nS) in the presence of 140 mM symmetrical [Cl−] can be estimated by fitting a mathematical model, with the experimental details defining initial conditions in the modeled system, to the fluorescence quenching time course (see (87)). A description of the model is provided in Text S13 and Fig. S14 shows the YFP(H148Q/I152L) quenching traces (solid red circles), together with the time course of several modeled variables, including the proportion of anion-free YFP(H148Q/I152L). Because, in our system, YFP(H148Q/I152L) fluorescence is completely quenched by anion binding (88), fitting the observed fluorescence time course to the predicted time course of the anion-free fluorophore allows parameter estimation. To account for variations in transfection efficiency, the mean mCherry fluorescence intensity within the cells was then used to normalize the G obtained by fitting.

At 28 °C, the estimated transient endogenous anion conductance (G_{trans}) under control (DMSO) conditions decayed more slowly than previous estimates at 37 °C (87). We first ran the model with four free parameters (V_m, G, G_{trans}, and τ_{trans}) see Text S13). However, at 28 °C, there was more overlap between the transient and CFTR-mediated currents making it harder to reliably estimate the values of parameters describing the transient current. For this reason, we constrained G_{trans} and τ_{trans} to the average values obtained from the negative (DMSO) controls (9 nS and 11.4 s, respectively) and ran all the fits estimating only V_m and G.

**MD simulations**

**System setup**

To build a starting structure for F508del-CFTR, we aligned one F508del-NBD1 from the hCFTR F508del NBD1 dimer (PDB ID 2PZF; (65)) to the NBD1 of full-length ATP-bound, hCFTR (PDB ID 6MSM; (64)). This alignment was made based on the α-carbon atoms and gave a RMSD of 0.8 Å. A chimera model was then built by replacing the WT NBD1 with the F508del-NBD1 from PDB ID 2PZF. This system is referred to as F508del/R1070. ATP molecules and Mg^{2+} ions were included at the binding sites. We used this structure to prepare two additional simulation systems, namely F508del/R1070F and F508del/R1070W, by replacing the arginine at position 1070 with a phenylalanine and a tryptophan, respectively. This in silico mutagenesis was performed using PyMOL (PyMOL Molecular Graphics System, Version 2.5, Schrödinger, LLC.), and for each mutation, the rotamer with the least overlap with nearby residues was chosen.

System preparation for MD simulations was performed using CHARMM-GUI (89–91) as follows: Each F508del system was inserted in a POPC lipid bilayer using a simulation box of approximately 13, 13, and 18 nm in x, y, and z directions, respectively. This resulted in a total of 444 POPC lipids per system. Potassium and chloride ions were added at a concentration of 0.15 M. The simulations were performed with GROMACS 2019.3 (92, 93), using the CHARMM36 force field with the TIP3P water model, as implemented in GROMACS (94). Each system was minimized with position restraints on the backbone and side chain atoms using a force constant of 50 kJ mol\(^{-1}\) nm\(^{-2}\) and from 2000 to 0 kJ mol\(^{-1}\) nm\(^{-2}\) for the side chain atoms, as recommended by the CHARMM-GUI protocol. The last stage of the equilibration was 20 ns long, with position restraints only on the backbone atoms using a force constant of 50 kJ mol\(^{-1}\) nm\(^{-2}\). Production runs, with no position restraints applied, lasted 2 μs with a time step of 2 fs. The temperature was maintained at 303.15 K using the Nosé–Hoover algorithm (95, 96) and a relaxation time of 1 ps. Semi-isotropic pressure coupling was applied with the Parrinello–Rahman barostat (97) with a relaxation time of 12 ps to maintain the pressure at 1 bar. The LINCS algorithm (98) was employed to constrain H-atom bonds. Neighbor searching was made with the Verlet cutoff scheme, with van der Waals interactions switched to 0 between 1 and 1.2 nm. The long-range electrostatic interactions were treated using the Particle Mesh Ewald (PME) algorithm (99, 100).

**Analyses**

The cluster analysis was performed using the tool gmx cluster in GROMACS 2019.6. For this analysis, for each simulation system, we extracted a trajectory containing only the atoms of the interface between ICL4 (residues 1050–1080 in hCFTR) and the F508del-NBD1 loop (residues 495–512 in hCFTR), with frames saved every 500 ps. The clustering was made with the Daura algorithm (101) using a cutoff of 0.1 nm. For the RMSD calculation, the alignment was performed on the backbone (N, αC, C) atoms of all the residues at the interface. The distribution of the χ₁ dihedral angle (N-Cα-Cβ-Cy) for selected aromatic residues at the interface
was calculated using the *gnx angle* tool in GROMACS2019.6 and plotted using Matplotlib libraries (102). The probability density was calculated over 2 µs-long trajectories for the F508del systems and over 1 µs-long trajectories for the previously published zCFTR systems (62). For clarity, although the ATP-bound protein is a Walker B glutamate E1372Q (corresponding to E1371Q in hCFTR) mutant, we refer to it here as “WT” to contrast it to the F508del mutants (note that the 6MSM hCFTR used to generate the F508del systems carries the same glutamate to glutamine mutation). The number of hydrogen bonds at the interface was estimated as a function of time based on a hydrogen–donor–acceptor angle cutoff of 30° and on a donor–acceptor distance cutoff of 0.35 nm, as implemented in the *gnx hbond* tool in GROMACS2019.6. Plots were made using Matplotlib libraries (102). Figures with snapshots from simulations or experimental structures were generated with either VMD1.9.3 (103) or PyMOL 2.0.6 (The PyMOL Molecular Graphics System, Schrödinger, LLC).

**Data and statistical analysis**

Statistical analysis was carried out in MATLAB (MathWorks) using the MATLAB Statistics Toolbox. Before statistical tests were performed, distributions were examined to assess whether they approximated normal distributions and whether there was homogeneity of variances. If the normality and homogeneity assumptions for parametric testing were not met, data were either transformed to meet the assumptions or analyzed using nonparametric tests. When *post hoc* tests consisted of all possible pairwise comparisons between groups, the Tukey–Kramer procedure was applied (using the *multcompare* function in Matlab) to prevent inflation of the type I error rate. By contrast, when comparisons were planned, the Benjamini–Hochberg procedure with a false discovery rate of 10% was applied to control the family-wise error rate. To describe the data, the following common statistical abbreviations were used: M (mean), Mdn (median), SD (standard deviation), SEM (standard error of the mean), N (number of measurements in the sample), and df (degrees of freedom). Statistical tests were performed two-sided unless otherwise specified. Data in the boxplots represent median, quartiles, range, and outliers, and data in other graphs represent mean ± SEM. The significance level was prespecified as α = 0.05. *p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001.

**Data availability**

Most data are presented in the figures of the article. In addition, supporting information is available, which includes a description of the mathematical model used for fitting the quenching traces, tables detailing the statistical analyses performed, and additional figures illustrating the scan and MD simulations.

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**Supporting information**—This article contains supporting information (62, 64, 87, 88, 104–109).

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**Abbreviations**—The abbreviations used are: ABC, ATP-binding cassette; CFTR, cystic fibrosis transmembrane conductance regulator; DMSO, dimethyl sulfoxide; G, whole-cell conductance; hCFTR, human CFTR; IRES, internal ribosome entry site; ICL, intracellular loop; MD, molecular dynamics; NBD, nucleotide-binding domain; Popen, open probability; PDB, Protein Data Bank; POPC, 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine; TM1-12, transmembrane helix 1–12; TMD, transmembrane domain; Vm, membrane potential, after steady-state activation of CFTR; YFP, yellow fluorescent protein; zCTFR, zebrafish CFTR.

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**References**

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between the sulfur atom of methionine/cysteine and backbone amides in proteins. J. Phys. Chem. Lett. 6, 1385–1389