Reduced PDZ Interactions of Rescued ΔF508CFTR Increases Its Cell Surface Mobility*

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Background: Phenylalanine 508 deletion from CFTR (ΔF508CFTR) is the most common cause of CF.

Results: Diffusional mobility of surface ΔF508CFTR is greater than that of wild type CFTR, irrespective of ΔF508CFTR rescue mechanism.

Conclusion: Defective ΔF508CFTR-PDZ interactions result in greater ΔF508CFTR diffusion and could contribute to its enhanced internalization.

Significance: Restoring ΔF508CFTR-PDZ-scaffolding interaction may have therapeutic benefit.

Deletion of phenylalanine 508 (ΔF508) in the cystic fibrosis transmembrane conductance regulator (CFTR) plasma membrane chloride channel is the most common cause of cystic fibrosis (CF). Though several maneuvers can rescue endoplasmic reticulum-retained ΔF508CFTR and promote its trafficking to the plasma membrane, rescued ΔF508CFTR remains susceptible to quality control mechanisms that lead to accelerated endocytosis, ubiquitination, and lysosomal degradation. To investigate the role of scaffold protein interactions in rescued ΔF508CFTR surface instability, the plasma membrane mobility of ΔF508CFTR was measured in live cells by quantum dot single particle tracking. Following rescue by low temperature, chemical correctors, thapsigargin, or overexpression of GRASP55, ΔF508CFTR diffusion was more rapid than that of wild-type CFTR because of reduced interactions with PDZ domain-containing scaffold proteins. Knock-down of the plasma membrane quality control proteins CHIP and Hsc70 partially restored ΔF508CFTR-scaffold association. Quantitative comparisons of CFTR cell surface diffusion and endocytosis kinetics suggested an association between reduced scaffold binding and CFTR internalization. Our surface diffusion measurements in live cells indicate defective scaffold interactions of rescued ΔF508CFTR at the cell surface, which may contribute to its defective peripheral processing.

The cystic fibrosis transmembrane conductance regulator (CFTR) is a cAMP-regulated chloride/bicarbonate channel predominantly expressed at the apical plasma membrane of epithelial cells in the airways, pancreas, kidney, testes, and gastrointestinal tract (1, 2). Loss-of-function mutations in the CFTR gene cause cystic fibrosis (CF) (1). The most common CF-causing mutation is deletion of phenylalanine at position 508 (ΔF508) of CFTR, which causes its retention in the endoplasmic reticulum (ER) and ubiquitin (Ub)-dependent proteasomal degradation (3, 4).

Cell membrane chloride conductance can be partially restored by maneuvers that correct (or rescue) ΔF508CFTR biosynthetic processing, promoting its exit from the ER and targeting to the cell surface. Experimental rescue maneuvers include permissive temperature (<30 °C), alteration of ER Ca2+, and activation of alternative ER export pathways (5–10). Several classes of drug-like small molecules also restore ΔF508CFTR processing and function (11–13). However, after correction, plasma membrane stability of ΔF508CFTR is reduced compared with that of native, wild-type (wt) CFTR. Incubation of low-temperature rescued (r)ΔF508CFTR at physiological temperature causes conformational destabilization, with ubiquitination, accelerated endocytosis, and lysosomal degradation (14, 15). Similar data have been obtained with rescue by chemical correctors (11, 12, 16).

Residues at the C terminus of CFTR (DTRL) form a class I PDZ (postsynaptic density protein 95/discs large/zebrafish ocludens-1) binding domain that associates with various PDZ domain-containing scaffold proteins including EB505 (ezrin-binding phosphoprotein of 50 kDa), CAL (CFTR-associated ligand), members of the GRASP (Golgi reassembly stacking protein) family and Shank2 (SH3 and multiple ankyrin repeat domains protein 2); however, the role of CFTR-PDZ interactions remains incompletely resolved (17). The CFTR PDZ-binding domain was originally reported to mediate apical polarization (18); however, these findings were challenged (19). Swiatecka-Urban et al. (20) reported that deletion of the CFTR PDZ-binding domain decreased CFTR stability by reducing recycling rates but not endocytosis. In contrast, Lukacs and co-workers (19, 21) reported that short C-terminal deletions from CFTR did not alter protein stability, whereas deletion of...
large regions (>50 amino acids) did increase CFTR turnover. In the Golgi, modulation of the function or abundance of PDZ-domain containing proteins has been shown to correct ΔF508CFTR trafficking (10, 22, 23). Similarly, EBP50 has been implicated in rΔF508CFTR biosynthetic processing. Knockdown of EBP50 in immortalized airway epithelial cells has been reported to reduce the surface stability of both wtCFTR and low-temperature rΔF508CFTR (24). In contrast, overexpression of EBP50 was found to rescue ΔF508CFTR in CBE41o-cells by a mechanism that involved cytoskeletal reorganization through EBP50-RhoA-Rho kinase-ezrin interactions (25, 26).

Using single particle tracking (SPT) in live cells, we previously demonstrated that PDZ interactions mediate the formation of a CFTR-containing macromolecular complex at the cell surface (Fig. 1A, top and Refs. 27, 28). SPT offers advantages over ensemble-averaged biophysical approaches used to study protein interactions in live cells. SPT provides information simultaneous on many cell surface proteins at exogenous expression levels with nanometer spatial and millisecond temporal resolution, allowing quantification of complex diffusive behaviors of protein subpopulations (29). In comparison to biochemical approaches involving cell disruption and detergent solubilization, SPT can detect weak or transient interactions in live cells.

In this study, we used SPT to test the hypothesis that ΔF508CFTR-PDZ protein interactions are reduced compared with those of wtCFTR. We found increased cell surface mobility of rΔF508CFTR compared with that of wtCFTR, which were found to involve reduced PDZ-interactions and elements of the peripheral quality control machinery. Disruption of CFTR-PDZ interactions enhanced endocytosis of wild type and ΔF508CFTR, suggesting a possible role of reduced PDZ interactions in the accelerated internalization of ΔF508CFTR.

**Experimental Procedures**

**Antibodies, siRNA, Additional Reagents, and cDNA Constructs**—Antibodies and siGENOME SMARTpool siRNA were used as previously described (15). VX-809 was purchased from Selleckchem. CoPo-22 (13) was provided by the UCSF Cystic Fibrosis Foundation Research and Development Program. Plasmid expression vectors encoding human wt- or ΔF508CFTR with triplet HA tags have been described and characterized previously (11, 15, 27). Additional CFTR constructs containing an external triplet FLAG tag were generated in pcDNA3.1 using engineered human CFTR cDNA (30); constructs were characterized functionally using a halide sensitive YFP assay as previously described (30). The C-terminal domain of CFTR with (residues 1432–1480) and without the PDZ-binding domain (residues 1432–1476) were amplified by PCR and ligated into pmCherry-C1. Plasmids encoding Myc-tagged human CHIP and CHIP-K30A were previously described (15). GFP-GRASP55 encoding plasmid was purchased from OriGene.

**Cell Culture, Transfection, and Rescue of ΔF508-CFTR Processing**—HeLa cell lines stably expressing 3HA-tagged CFTR constructs were previously described (15). An MDCK type II cell line expressing 3HA-tagged wtCFTR was previously described (27), and ΔF508CFTR-expressing cells were generated in an analogous manner. Untransfected COS-7 and MDCK cells were obtained from the UCSF Cell Culture Facility. For all microscopy, cells were cultured on 18-mm coverslips in 12-well cell culture plates for 2–4 days prior to experiments. Cells were transiently transfected using Lipofectamine 2000 (Invitrogen) 2–3 days prior to experiments. Low temperature rescue of ΔF508CFTR was performed by incubating cells at 27 °C for 24–30 h; as necessary, subsequent thermal unfolding was performed by incubation of cells for an additional 1–2 h at 37 °C (15). The following pharmacological maneuvers were used to correct ΔF508CFTR trafficking: 3 μM VX-809 for 24 h (12), 10 μM CoPo-22 for 24 h (13), and 1 μM thapsigargin for 2 h (10).

**Single Particle Tracking and Data Analysis**—Qdot-labeling, SPT data collection, and analysis was performed essentially as described in prior studies (27, 28, 31). Data were collected with a Nikon 100×/N.A.1.3 objective. Datasets were compiled from 110–788 trajectories (average of 215 trajectories per maneuver) obtained from multiple cells on different days. Given that CFTR is endocytosed, experiments were performed within 3–5 min of labeling to reduce errors associated with inclusion of data derived from endocytosed CFTR molecules (which is assumed to be <10% in all datasets). Statistical analysis of SPT data were performed using the Kolmogorov-Smirnov test (31, 32).

**Determination of CFTR Surface Density, Stability, and Endocytosis Rates**—Epitope-tagged CFTR surface density and endocytic rates were determined essentially as described in a prior study (15). Cell surface density of epitope-tagged CFTR variants was measured by labeling surface CFTR at 4 °C with primary antibody (anti-HA or anti-FLAG) followed by: (i) a secondary antibody conjugated to fluorophore for image-based detection (after fixation), or (ii) horseradish peroxidase conjugated secondary antibody and Amplex Red (Invitrogen) for quantification in an ELISA format. Relative ΔF508CFTR surface density presented in Fig. 4 is the normalized difference in surface density before and after incubation at 37 °C for 2 h. To determine endocytic rates for CFTR variants, cells were labeled with primary antibody and incubated at 37 °C for 0 or 10 min prior to labeling with secondary antibody to determine CFTR surface density. Nonspecific antibody labeling was assessed using non-transfected cells. Control experiments in HeLa cells expressing 3HA-tagged CFTR verified that imaging and Amplex Red-based analysis gave equivalent results. Datasets are presented as mean ± S.E. from 3–8 independent experiments. For Amplex Red, individual measurements were performed using 6–8 parallel samples and normalized to protein content; for image-based analysis, cell surface CFTR density measurements were performed on >30 cells per data set. Statistical analysis was by t test.

**Results**

**Increased Cell Surface Diffusion of rΔF508CFTR Compared with wtCFTR**—Initial studies were done using HeLa cells expressing CFTR constructs containing triplet hemagglutinin tags (CFTR3HA) engineered into the fourth external loop region; these cells were used previously to characterize and identify cellular components involved in rΔF508CFTR processing (15). For surface mobility measurements using SPT, cell
surface CFTR was selectively labeled by sequential incubation with low concentrations of anti-HA antibody, secondary biotin-conjugated Fab fragment, and streptavidin-conjugated Qdots (emission 655 nm). As in prior studies (27, 31), Qdot labeling specificity was confirmed by absence of Qdot labeling when primary antibody or secondary Fab fragment were omitted, or when ΔF508CFTR was not rescued (data not shown). Diffusion of Qdot-labeled CFTR was measured at 37 °C by analysis of image sequences acquired at 30 Hz over 10 s (Fig. 1B). Mobility of wtCFTR was highly confined (Fig. 1B, top), as found previously for wtCFTR expressed at endogenous levels in multiple cell types, including human airway epithelial cells (27, 28).

Immediately following low temperature rescue, where ΔF508CFTR plasma membrane stability, endocytosis and ubiquitination are similar to that of wtCFTR (15), the diffusion (measured at 37 °C) of ΔF508CFTR (ΔF508CFTR37) was also similar to that of wtCFTR (Fig. 1B, middle). However, thermal unfolding of ΔF508CFTR by 2 h of incubation at 37 °C (ΔF508CFTR37), which causes accelerated endocytosis (~2-fold), enhanced ubiquitination and an overall reduction in cells surface ΔF508CFTR levels (15), produced an increase in mobility (Fig. 1B, bottom). Trajectories generated from analysis of multiple image sequences (Fig. 1, B, right and D) were summarized as cumulative probability distributions of particle range at 3 s. Fig. 1C shows significantly greater range for ΔF508CFTR37 diffusion (blue) in comparison to wtCFTR (black) or ΔF508CFTR27 (red, see Fig. 1D for representative trajectories). Diffusion of wtCFTR was not altered under the same culture conditions used to rescue and thermally unfold ΔF508CFTR (supplemental Fig. S1A).

We previously established that confined diffusion of wtCFTR in live cells is dependent on its C terminus PDZ binding motif, PDZ domain containing scaffold proteins, and coupling to the actin cytoskeleton via ezrin (27). This finding is consistent with in vitro protein-protein interaction data suggesting the involvement of wtCFTR in a cell surface macromolecular complex. Here, to investigate the involvement of PDZ interactions in ΔF508CFTR confinement in live cells, a chimeric protein consisting of the C terminus of CFTR (residues 1432–1480, which does not contain the endocytic motifs Y1433FLVI, Y1434YDSI, and Y1439LL) fused to mCherry (mCh-CFTR-C) was expressed to compete with CFTR for attachment to PDZ domain proteins. As expected from our previous data (27), mCh-CFTR-C expression greatly increased wtCFTR diffusion (Fig. 1, C and D, gray). Low-temperature corrected ΔF508CFTR has been demonstrated to interact with EB50 (24) and mCh-CFTR-C also increased the diffusion of ΔF508CFTR37 (Fig. 1, C and D, orange) and ΔF508CFTR27 (Fig. 1, C and D, green) to the same degree as wtCFTR, suggesting that PDZ interactions contribute to the confinement of cell surface ΔF508CFTR. As previously
demonstrated for wtCFTR (27), expression of a CFTR C-terminal fusion protein that lacked the PDZ-binding motif (mCh-CFTR-C-ΔPDZ) did not alter the diffusion of wtCFTR, ΔF508CFTR27, and ΔF508CFTR37 in HeLa cells (supplemental Fig. S1B). These findings show in live cells that interactions with PDZ-domain-containing proteins contribute to the tethering of wt and low-temperature rescued ΔF508CFTR, but that interactions between rΔF508CFTR27 and PDZ domain-containing proteins are reduced compared with wtCFTR or rΔF508CFTR37.

CFTR diffusion was measured in additional cell types and with transiently transfected CFTR constructs to verify the robustness of observations made in HeLa cell lines. In MDCK cell lines stably expressing CFTR3HA constructs, incubation of ΔF508CFTR-expressing cells at 27 °C increased ΔF508CFTR cell surface density to a level ~6-fold reduced compared with wtCFTR (supplemental Fig. S1C). As predicted from prior studies in different cell types (14, 15), incubation at 37 °C for 1 h to thermally denature rΔF508CFTR produced an ~50% decrease in plasma membrane rΔF508CFTR density (Fig. 2A, left) and an ~2-fold increase in endocytosis compared with wtCFTR; however, ΔF508CFTR internalization was similar to wtCFTR immediately after rescue (Fig. 2A, right).

SPT analysis of CFTR diffusion in MDCK cell lines showed highly confined diffusion of wtCFTR (black) and rΔF508CFTR27 (red, Fig. 2B); however, incubation for 1 h at 37 °C increased rΔF508CFTR diffusion as seen in individual trajectories (blue) and cumulative probability distributions (Fig. 2C, black and blue). The diffusion of wtCFTR and rΔF508CFTR27 were significantly increased by expression of mCh-CFTR-C (Fig. 2C, gray and green). A comparable increase in channel diffusion was seen with deletion of the C-terminal PDZ-binding motif of CFTR (CFTR-ΔPDZ, Fig. 2C, pink), suggesting that expression of mCh-CFTR-C mobilizes wt and rΔF508CFTR to the same degree as deletion of the PDZ-binding motif. In control experiments, mCh-CFTR-C-ΔPDZ did not alter the diffusion of wtCFTR (Fig. 5C), rΔF508CFTR27, or rΔF508CFTR37 (supplemental Fig. S1D), verifying the involvement of PDZ-domain interactions in confinement of wtCFTR, rΔF508CFTR27, and rΔF508CFTR37. The smooth shape of cumulative probability distributions suggested the absence of distinct CFTR subpopulations with different mobilities. As such, data were summarized in bar graph format (Fig. 2D), showing median (blue) and inter-quartile (25th to 75th percentiles, red) range values. As described above, data from MDCK cell lines (Fig. 2D) shows significantly different diffusion of rΔF508CFTR37 versus wtCFTR or rΔF508CFTR27 (bars 1–3), and increased CFTR diffusion upon disruption of PDZ interactions (bars 3–6).

CFTR diffusion was further studied in COS-7 fibroblasts that were transiently transfected to express CFTR constructs containing external FLAG epitopes (CFTR3FLAG). As expected, surface density decreased and endocytosis increased after 37 °C incubation of low temperature corrected ΔF508CFTR (Fig. 2E). The diffusion of wtCFTR and rΔF508CFTR27 were similar in COS-7 cells, whereas incubation of ΔF508CFTR-expressing cells at 37 °C increased diffusion by ~30% (Fig. 2F, bars 1–3). Deletion of the PDZ binding motif from wtCFTR increased diffusion (Fig. 2F, bar 4), as did deletion from ΔF508CFTR following low temperature rescue (ΔFΔPDZ27) and subsequently incubated at 37 °C (ΔFΔPDZ27) (Fig. 2F, bars 5 and 6). The magnitude of the effect on diffusion of PDZ-domain deletion varied for wtCFTR and rΔF508CFTR in different cell types; however, this difference likely reflects differences in plasma membrane properties as MDCK cells have been shown to reduce the diffusion of inert probe molecules as compared with other cell types (33). Further, the diffusion of wt and ΔF508CFTR in the absence of PDZ interactions in both MDCK and COS-7 cells was comparable to that of AQP1, a membrane transport protein that has largely unhindered diffusion (33). These studies confirm that in different cell types ΔF508CFTR rescued by low-temperature incubation interacts with PDZ...
domain proteins in a similar manner as wtCFTR, but that thermal unfolding at 37 °C increases ΔF508CFTR diffusion by partially disrupting the interactions between rΔF508CFTR and PDZ domain-containing scaffold proteins.

ΔF508CFTR Diffusion Is Increased with Alternative Correction Methods—SPT experiments were done to investigate cell surface ΔF508CFTR-PDZ interactions following different correction maneuvers. Treatment of MDCK cell lines expressing CFTR3HA with 3 μM VX-809, a corrector drug in Phase II clinical trials (12, 34), for 24 h gave ΔF508CFTR cell surface levels similar to those produced by low-temperature rescue (supplemental Fig. S1C). SPT trajectories of ΔF508CFTR diffusion following VX-809 correction appeared less confined than those of wtCFTR under control conditions (Fig. 3A, red and black trajectories) and range values differed significantly (Fig. 3B, bars 1 and 3). Expression of mCh-CFTR-C increased the diffusion of VX-809-corrected ΔF508CFTR (Fig. 3A, blue) to that found for wtCFTR with mCh-CFTR-C or for CFTRΔPDZ (Fig. 3B, bars 4–6), confirming the involvement of PDZ interactions in restricting rΔF508CFTR mobility after VX-809 treatment. Control experiments verified that mCh-CFTR-C-ΔPDZ did not alter the diffusion of rΔF508CFTR corrected with VX-809 (data not shown).

Additional experiments were performed in transiently transfected MDCK cells using VX-809 to correct 3FLAG-tagged ΔF508CFTR to verify results generated in cell lines (Fig. 3, C and D). In transiently transfected MDCK cells, wtCFTR mobility was reduced when compared with VX-809 rescued ΔF508CFTR (Fig. 3C, black and red; and Fig. 3D, bars 1 and 2). The diffusion of cell surface ΔF508CFTR was increased by expression of mCh-CFTR-C (but not mCh-CFTR-C-ΔPDZ) or deletion of the ΔF508CFTR C-terminal PDZ-binding motif (rΔFAPDZ) (Fig. 3, C and D and data not shown). VX-809 treatment did not alter the diffusion of wtCFTR or CFTRΔPDZ, suggesting that VX-809 does not perturb plasma membrane properties (for example see Fig. 3B, bars 1 and 2; Fig. 3D, bars 5 and 6).

The diffusion of cell surface ΔF508CFTR in MDCK cell lines was also increased compared with wtCFTR (Fig. 3E) following correction by thapsigargin (black), the small molecule corrector CoPo-22 (red), or transient expression of GFP-GRASP55 (blue; for all datasets p < 0.001 compared with wtCFTR, Fig. 3F, top). These treatments did not alter the diffusion of wtCFTR (Fig. 3E, left and Fig. 3F, bottom). For GRASP55-mediated ΔF508CFTR rescue, immunocytochemical confirmed GFP-GRASP55 co-localization with the Golgi marker giantin (Fig. 3G). Photobleaching experiments using laser scanning confocal microscopy indicated that the diffusion of the inert membrane probe glycophosphatidylinositol-anchored GFP was not altered by each of the maneuvers used to correct ΔF508CFTR trafficking (data not shown). These studies thus demonstrate that, as found for low temperature correction, rescue by a variety of unrelated maneuvers produces cell surface ΔF508CFTR with abnormal PDZ-domain interactions relative to wtCFTR.

Components of the Peripheral Protein Quality Control System Modulate ΔF508CFTR Mobility—Experiments were performed to identify cellular components that are responsible for reduced ΔF508CFTR-PDZ interactions. Using a small interfering (si)-RNA screen, Okiyoneda et al. (15) previously identified constituents of the peripheral protein quality control (PPQC) system that recognize and eliminate low-temperature corrected and thermally denatured ΔF508CFTR from the cell surface. This approach identified the E3 Ub ligase CHIP (C terminus of Hsc70-interacting protein) and the chaperone Hsc70 (heat shock cognate of 70 kDa) as key determinants of rΔF508CFTR turnover such that knockdown of either protein increased cell surface stability. Here, we tested whether altering the abundance or activity of CHIP and Hsc70 would also affect ΔF508CFTR mobility. rΔF508CFTR37 diffusion was...
The peripheral protein quality control (PPQC) system alters rΔF508CFTR diffusion. A, effects of siRNA-mediated knockdown of CHIP or Hsc70 (left, top) on rΔF508CFTR<sub>37</sub> surface levels (left, bottom, values >1 represent stabilization of rΔF508CFTR<sub>37</sub>, relative to control conditions) and diffusive range (right) in low temperature corrected HeLa cells. For reference, median and interquartile values for wtCFTR diffusive range are shown by dashed lines (bar 4, right). B, transient co-expression of rΔF508CFTR<sub>37</sub> with CHIP but not CHIP-K30A alters rΔF508CFTR<sub>37</sub> diffusive range (top) and surface levels (bottom) in low temperature corrected COS-7 cells. C, siRNA-mediated CHIP knockdown in HeLa cells (left) and CHIP expression in MDCK cells (right) alter rΔF508CFTR diffusion (top) and surface levels (bottom) in VX-809-treated cells. For reference, median and interquartile values for wtCFTR diffusive range in HeLa cells are shown by dashed lines (bar 4, left). Statistical difference is denoted (*); for SPT data p < 0.001 and for surface stability data p < 0.05. SPT data were derived from 276–316 trajectories for low temperature corrected HeLa cells, from 124–255 trajectories for COS-7 cells, from 160 and 226 trajectories for VX-809-treated HeLa cells and from 162–298 trajectories for MDCK cells. Data for CFTR surface stability are mean ± S.E. for 3–8 independent experiments.

To further investigate the role of CHIP in rΔF508CFTR surface diffusion, wild type, and mutant CHIP constructs were transiently expressed in COS-7 cells. As expected based on earlier observations (15), overexpression of CHIP in COS-7 cells reduced rΔF508CFTR<sub>37</sub> surface stability (Fig. 4B, bottom) and also increased rΔF508CFTR<sub>37</sub> surface diffusion (Fig. 4B, top). CHIP binds to Hsc70 and Hsp90 (heat shock protein of 90 kDa) via an N-terminal tetratricopeptide repeat domain, and these associations are blocked in the CHIP-K30A mutant (35). Expression of CHIP-K30A in COS-7 cells did not alter rΔF508CFTR<sub>37</sub> diffusion or stability (Fig. 4B), indicating that CHIP recognition of thermally destabilized rΔF508CFTR is dependent upon Hsc70 and Hsp90. In VX-809-treated HeLa cells rΔF508CFTR diffusion was similar to that of rΔF508CFTR<sub>37</sub>, as would be expected from prior studies in MDCK cells (Fig. 4C, top left). Further, siRNA knockdown of CHIP reduced rΔF508CFTR mobility and increased surface stability (Fig. 4C, left). CHIP expression also reduced confinement and cell surface stability of rΔF508CFTR in transiently transfected MDCK cells treated with VX-809; however, expression of CHIP-K30A did not alter diffusion or stability of VX-809-corrected rΔF508CFTR, confirming that CHIP recognition of rΔF508CFTR required Hsc70 and Hsp90 (Fig. 4C, right). In control experiments, wtCFTR mobility was not altered by CHIP knockdown in HeLa cells or CHIP overexpression in COS-7 cells; non-targeted siRNA did not alter wtCFTR or rΔF508CFTR diffusion in HeLa cells (supplemental Fig. 5E). These studies indicate that components of the PPQC machinery including CHIP, Hsc70 and Hsp90, contribute to the increased diffusion of cell surface ΔF508CFTR rescued by low-temperature or VX-809.

PDZ Interactions Stabilize wtCFTR at the Cell Surface—The present study indicates that reduced rΔF508CFTR surface stability is associated with increased diffusive mobility. Given that endocytosis is the initial step in reducing rΔF508CFTR surface density, an experimental strategy was developed to quantitatively investigate whether the observed increase in diffusion associated with reduced PDZ interactions of ΔF508CFTR could alter CFTR internalization rates. Our approach relied upon exogenous expression of the CFTR C terminus (mCh-CFTR-C) to increase wtCFTR mobility and thus mimic rΔF508CFTR surface dynamics (Fig. 5A). CFTR3HA-expressing MDCK cells were enriched for mCh-CFTR-C content by fluorescence-activated cell sorting (FACS) to generate a cell population with ~80% mCh-CFTR-C-positive cells that were readily identified in image-based assays of CFTR trafficking and surface stability (Fig. 5B, left). Expression of mCh-CFTR-C in FACS-enriched cells was relatively homogeneous (Fig. 5B, right), albeit ~40-fold less than in prior experiments that used transient transfection of the same construct. Surface expression of wtCFTR was similar in control and mCh-CFTR-C-enriched cells, suggesting that expression of exogenous CFTR C-terminal domains does not alter CFTR stability (supplemental Fig. S1C). This result is consistent with data from CFTR with short C-terminal deletions that show similar stability to wtCFTR (19). SPT confirmed that wtCFTR diffusion in mCh-CFTR-C-enriched cells was greater (by ~25%) than that in control cells (Fig. 5C, bars 1 and 3), and similar to that of ΔF508CFTR rescued by low-temperature or VX-809 (Fig. 5C, bars 4 and 5).

Endocytosis of wtCFTR in mCh-CFTR-C-enriched cells was increased by ~30% as compared with wtCFTR (Fig. 5D, bar 1 and 3, p < 0.005, ~16 versus 21%), suggesting that PDZ interactions stabilize CFTR at the cell surface over time scales of minutes. In support of this conclusion, deletion of the CFTR C-terminal PDZ binding domain (CFTRΔPDZ), which is expected to fully account for any PDZ-dependent component...
of CFTR internalization (and increased diffusive by ~50%, Fig. 2), increased endosyndy by ~60% as compared with wtCFTR (Fig. 5D, bar 4, p < 0.001, ~16 versus 26%). Therefore, compared with wtCFTR, the percentage increase in surface diffusion and in endocytosis rate after deletion of the CFTR PDZ-binding domain or upon partial disruption of PDZ interactions to model ΔF508CFTR diffusion is in reasonable agreement. Control experiments confirmed that CFTR diffusion (Fig. 5C, bar 2) and internalization (Fig. 5D, bar 2) were not altered by enrichment of cells for expression of the PDZ-binding-domain deleted CFTR C terminus (mCh-CFTR-C-ΔPDZ). Finally, in MDCK cells transiently expressing VX-809-rescued ΔF508CFTR3FLAG, co-expression of mCh-CFTR-C at high levels enhanced ΔF508CFTR internalization, confirming that PDZ interactions can also influence rΔF508CFTR endocytosis (Fig. 5E). These studies indicate that (i) PDZ interactions stabilize CFTR at the cell surface over time scales of minutes, and (ii) increasing the diffusion of wtCFTR to that of rΔF508CFTR is sufficient to alter internalization kinetics.

**DISCUSSION**

SPT provides definitive information about the motions of individual channels, receptors or lipids in the plasma membrane of live cells, which can reveal complex diffusive behaviors over extended time and distance scales (29, 36). Our prior SPT studies on wtCFTR showed that it integrates into an actin-associated, macromolecular complex that includes PDZ domain-containing proteins and ezrin (27, 28). Here, SPT analysis of the dynamics and interactions of rescued ΔF508CFTR revealed novel properties relative to wtCFTR that were related to the peripheral quality control mechanism and, potentially, ΔF508CFTR internalization dynamics.

In live cells at physiological temperature, we found consistently greater cell surface mobility of rΔF508CFTR (by ~30%) compared with wtCFTR, regardless of correction mechanism, which was due to distinct association with PDZ scaffold proteins. Incubation of low temperature-corrected ΔF508CFTR at 37 °C causes increased ubiquitination and endocytosis, and consequent reduced plasma membrane stability (15, 37). Though rΔF508CFTR-PDZ interactions were similar to those of wtCFTR just after low-temperature correction, incubation at 37 °C increased rΔF508CFTR diffusion and reduced PDZ binding. Unfolded ΔF508CFTR is eliminated from the cell surface by the PPQC system. We found that modulation of the PPQC system also altered the diffusion of rΔF508CFTR, suggesting an interdependence of rΔF508CFTR surface stability and mobility.

The role of CFTR-PDZ interactions is incompletely understood. PDZ interactions were initially reported to mediate CFTR polarization, but subsequent contradictory evidence was reported (18, 19). Inconsistent data has also been reported on the role of PDZ interactions in CFTR turnover (19–21). Several studies have concluded that multivalent PDZ scaffolds facilitate compartmentalization of proteins to regulate CFTR activity (17, 38). Our experimental approach provided evidence in live cells for involvement of PDZ interactions in CFTR complexation and stabilization at the cell surface over time scales of minutes. This conclusion is based on experimental approaches that either deleted the PDZ-binding domain from CFTR or used expression of an exogenous peptide (mCh-CFTR-C) to com-
pete with CFTR for PDZ domain-containing scaffold proteins. Previously, knockdown of EBP50 was shown to reduce surface stability of wtCFTR and low-temperature rΔF508CFTR over hours (24). However, interpretation of this study is difficult because EBP50 is a multi-domain protein that interacts with distinct PDZ-binding domain proteins and the actin cytoskeleton. Further, altering EBP50 expression profoundly alters the actin cytoskeleton (26).

In experiments designed to quantitatively relate CFTR diffusion to internalization dynamics in MDCK cells, deletion of the CFTR PDZ-binding domain (~50% increase in diffusion compared with wtCFTR) enhanced endocytosis by ~60% whereas partial disruption of PDZ interactions to mimic rΔF508CFTR diffusion (~30% increase in diffusion) increased internalization by ~30%. Endocytosis of ΔF508CFTR in MDCK cells after low-temperature correction and thermal unfolding, or VX-809 treatment, were ~1.8-fold greater than that of wtCFTR. Therefore, the proportion of rΔF508CFTR internalization that relates to increased rΔF508CFTR diffusion (Fig. 5D, blue dashed line) could be ~50%. Although reduced scaffold binding and consequent accelerated endocytosis would predispose rΔF508CFTR to lysosomal trafficking and degradation, the overall elimination of rΔF508CFTR is ultimately dependent on channel ubiquitination (14, 15). Indeed, there are several disease-causing CFTRs with C terminus deletions yet relatively mild disease, which has suggested that severe disease, as found for the ΔF508 mutation, requires CFTR structural destabilization and consequent ubiquitination in addition to enhanced endocytosis (14, 15, 19, 21).

A limited number of prior studies have demonstrated that PDZ interactions stabilize proteins at the cell surface by reducing endocytic uptake. For instance, stimulated G-protein coupled receptors that contain PDZ-binding domains segregate into specific clathrin-coated pits to greatly extend their surface lifetime through actin association (39). Similar to the present study, a correlative analysis of protein diffusion and internalization has been performed for the neuronal α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor (AMPAR) (40). The AMPAR forms a complex with Stargazin and the PDZ scaffold PSD-95, which result in receptor immobilization at the neuronal postsynaptic density. Disruption of Stargazin-PSD-95 interactions with a cell permeable peptide mimetic of the Stargazin PDZ binding domain increased AMPAR diffusive range by ~1.3-fold and endocytosis by ~4-fold (40). For CFTR, Thelin et al. (41) reported that the S13F mutation increased CFTR diffusion coefficient by only ~20%, but increased internalization by ~2-fold. However, in that study CFTR was reported as mobile under control conditions, presumably due to CFTR expression at super-physiological levels (27), making interpretation of these findings difficult.

Components of the PPQC network, including CHIP, Hsc70 and Hsp90, were found here to alter the plasma membrane dynamics of rΔF508CFTR after low temperature or chemical correction. Although the mechanism responsible for PPQC modulation of rΔF508CFTR diffusion was not established, we speculate that rΔF508CFTR-PDZ interactions might be reduced by ubiquitination, or perhaps sterically hindered by rΔF508CFTR-CHIP/Hsp90/Hsc70 complexation. Further work is required to address these hypotheses. CHIP and associated quality control components have been shown to mediate ubiquination of ΔF508CFTR at the cell surface and the ER, and inhibition of CHIP causes ΔF508CFTR accumulation in the ER (15, 42, 43). Functional inhibition of CHIP or a CHIP-binding partner thus represents a potential therapeutic approach to stabilize rΔF508CFTR at the cell surface.

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