4-Phenylbutyric acid treatment rescues trafficking and processing of a mutant surfactant protein C

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Mutations in the *SFTPC* gene, encoding surfactant protein–C (SP-C), are associated with interstitial lung disease (ILD). Knowledge of the intracellular fate of mutant SP-C is essential in the design of therapies to correct trafficking/process of the proprotein, and to prevent the formation of cytotoxic aggregates. We assessed the potential of a chemical chaperone to correct the trafficking and processing of three disease-associated mutant SP-C proteins. HEK293 cells were stably transfected with wild-type (SP-CWT) or mutant (SP-CΔL188Q, SP-CΔ exon4, or SP-CΔC exon7) SP-C, and cell lines with a similar expression of SP-C mRNA were identified. The effects of the chemical chaperone 4-phenylbutyric acid (PBA) and lysosomotropic drugs on intracellular trafficking to the endolysosomal pathway and the subsequent conversion of SP-C proprotein to mature peptide were assessed. Despite comparable SP-C mRNA expression, proprotein concentrations varied greatly: SP-CΔC was more abundant than SP-CWT and was localized to the cell surface, whereas SP-CΔ exon4 was barely detectable. In contrast, SP-CΔL188Q and SP-CWT proprotein concentrations were comparable, and a small amount of SP-CΔL188Q was localized to the endolysosomal pathway. PBA treatment restored the trafficking and processing of SP-CΔL188Q to SP-CWT concentrations, but did not correct the mistrafficking of SP-CΔC or rescue SP-CΔ exon4. PBA treatment also promoted the aggregation of SP-C proproteins, including SP-CΔL188Q. This study provides proof of the principle that a chemical chaperone can correct the mistrafficking and processing of a disease-associated mutant SP-C proprotein.

**Keywords:** pulmonary surfactant protein–C; interstitial lung disease; 4-phenylbutyric acid; chemical chaperone

Intersitial lung disease (ILD) is the result of a multiplicity of pathological processes, from infections to autoimmune diseases (1). In addition to idiopathic forms of the disease, familial ILD has also been reported. Mutations in the gene encoding surfactant protein–C (SP-C) are associated with heritable ILD in both children (2) and adults (1), and are the focus of this study.

**CLINICAL RELEVANCE**

Mutations in the gene encoding surfactant protein–C (*SFTPC*) are associated with interstitial lung disease in both children and adults. The pathogenesis is thought to be related to an accumulation of cytotoxic aggregates of misfolded surfactant protein–C proprotein (proSP-C). No treatment options are currently available. Treatment with 4-phenylbutyric acid (PBA) rescues at least one form of mutant proSP-C from proteasome degradation, and restores trafficking to the endolysosomal pathway as well as the conversion of the proprotein to the mature peptide. This study shows that PBA may be a viable treatment option for some patients with *SFTPC* mutations.

SP-C, encoded by the *SFTPC* gene on chromosome 8 in humans, is a hydrophobic protein produced by alveolar Type II epithelial cells. Human SP-C is synthesized as a 197–amino acid precursor protein (proSP-C) that undergoes multiple proteolytic cleavages as it traffics through the endolysosomal pathway to the lamellar body. The fully processed 35–amino acid mature SP-C peptide is packaged with SP-B and phospholipids into bilayer membranes that are stored in lamellar bodies before secretion into the alveolar airspaces.

Nogee and colleagues described the first disease-associated *SFTPC* mutation in an infant and her mother in 2001 (3). An *SFTPC* mutation that resulted in the skipping of exon 4 (SP-CΔ exon4) was identified in one allele in both subjects, consistent with autosomal dominant inheritance. A number of other autosomal dominant *SFTPC* mutations were subsequently identified with a range of histological phenotypes, even within kindreds. For example, a mutation leading to the substitution of glutamine for leucine at position 188 of proSP-C (SP-CΔ L188Q) resulted in a variable disease onset, with nonspecific interstitial pneumonia (NSIP) in children and usual interstitial pneumonia (UIP) in adults of the same family (4, 5).

Very little is known about the pathogenesis of lung disease associated with *SFTPC* mutations. The overexpression of mutant proSP-C can cause cytotoxicity in transfected cells (4) and transgenic mice (6). However, not all individuals who carry a disease-associated mutant *SFTPC* allele are symptomatic (4, 5). This observation led to the hypothesis that misfolded proSP-C is normally rapidly degraded but, under specific conditions (e.g., viral infection), might accumulate to form cytotoxic aggregates that, in turn, could trigger disease onset (7). In this model of disease pathogenesis, the goal of therapy would involve accelerating the turnover of the misfolded proprotein to prevent aggregation. This therapeutic approach comes at the expense of reduced SP-C in the airspaces. An alternative approach would involve the rescue of misfolded proprotein by promoting folding to a stable conformation that will pass quality control in the endoplasmic...
reticulum (ER) and traffic to the endolysosomal pathway for processing to the mature peptide.

Chemical chaperones are small molecules that promote the stabilization and folding of nascent proteins (8). The chemical chaperone 4-phenylbutyric acid (PBA) has been shown to rescue the trafficking of a wide range of misfolded proteins, including the ATP-binding cassette (ABC) transporters ABCB6 (9) and ABCC7/cystic fibrosis transmembrane conductance regulator (10). PBA is one of only two chemical chaperones currently approved by the United States Food and Drug Administration for use in humans. The present study evaluated the potential of PBA to rescue the trafficking and processing of three different mutant SP-C proproteins. The most common SFTPCC allele associated with ILD in both pediatric and adult patients is a substitution of threonine for isoleucine at position 73 of proSP-C (SP-C<sup>I73T</sup>) (11–13). This mutation occurs in a highly conserved region of the linker domain, a 34 amino acid juxtamembrane portion of the C-terminal (luminal) region of the proprotein. In contrast, the SP-C<sup>Δ<exon4</sup></sup> and SP-C<sup>Δ<exon4>Δ<exon11</exon12></sup> mutations are located in the 105–amino acid, C-terminal Brichos domain, a region to which the majority of SFTPCC mutations map. The Brichos domain functions as an intramolecular chaperone for the metastable transmembrane domain of the proprotein (14–16). Disease-associated Brichos mutations are thought to compromise chaperone activity, leading to terminal misfolding and the proteasomal degradation of the proprotein via the ER-associated degradation (ERAD) pathway.

In this study, we used cell lines that stably (constitutively) express nontagged SP-C<sup>WT</sup>, SP-C<sup>I73T</sup>, SP-C<sup>Δ<exon11</exon12></sup>, or SP-C<sup>Δ<exon4></sup> (7, 17) to avoid artifacts associated with the transient overexpression of SP-C proproteins. The stability of each mutant proprotein and its ability to traffic to the endolysosomal pathway for processing to mature peptide were evaluated in the presence or absence of chemical chaperones and lysosomalotropic drugs. The results demonstrate for the first time, to the best of our knowledge, that PBA at clinically achievable and tolerable concentrations can rescue SP-C<sup>Δ<exon4></sup> from ERAD, and restore both trafficking and processing of the proprotein to its mature peptide. A portion of these studies was previously reported in abstract form (18).

**MATERIALS AND METHODS**

**Reagents**

PBA, chloroquine, formic acid, brefeldin A, and glycerol were purchased from Sigma (St. Louis, MO). Bafilomycin A1 was purchased from Calbiochem (San Diego, CA). Antibodies directed against the N-terminus of the SP-C proprotein or the mature SP-C peptide were obtained from Seven Hills Bioreagents (Cincinnati, OH). The generation of the antibody directed against the C-terminal (luminal) domain of SP-C was previously described (16).

**Generation of Stably Transfected Lines**

Full-length human wild-type (SP-C<sup>WT</sup>) or mutant (SP-C<sup>Δ<exon11</exon12></sup>, SP-C<sup>Δ<exon4></sup>Δ<exon4></sup>, or SP-C<sup>Δ<exon7</exon7></sup>) SP-C was subcloned into pTRE2-Hyg (Clontech, Mountain View, CA), and stably transfected cell lines were generated in HEK293 Tet-Off cells, as previously described (7).

**Analyses of SP-C mRNA and Protein Expression**

SDS-soluble and insoluble proteins were isolated, normalized to the total cell lysate protein concentration, and analyzed by SDS-PAGE/ Western blotting, as previously described (6, 7, 15). Pulse-chase experiments were performed as previously described (17). The isolation of RNA for quantitative PCR and microarray analyses is described in the online supplement. Unfolded protein response element luciferase reporter assays and confocal microscopy were performed exactly as previously described (7, 19). Immunogold labeling and ultrastructural analyses of HEK293 cells are described in the online supplement.

**Statistical Analysis**

All experiments were performed at least three times, except where noted. Data were analyzed using GraphPad Prism, version 5.0 (GraphPad Software, Inc., La Jolla, CA). The data are representative of at least three independent transfection experiments, with each group performed in triplicate. Data are plotted as means ± SD, and were analyzed by the Tukey multiple comparisons test. Statistical significance was defined as P < 0.05.

**RESULTS**

**SFTPCC Mutations Result in Widely Variable Proprotein Stability**

Except where noted, all experiments in this study were performed in stably transfected HEK293 cells to model the constitutive expression of SP-C. Multiple cell lines stably expressing nontagged SP-C<sup>WT</sup> or mutant SP-C (SP-C<sup>I73T</sup>, SP-C<sup>Δ<exon4></sup>Δ<exon4></sup> or SP-C<sup>Δ<exon4>Δ<exon11</exon12></sup>) were grown in the absence of doxycycline to induce the expression of the proprotein, and screened to identify four cell lines that expressed similar concentrations of SP-C mRNA (Figure 1A). Cell lysates (30 μg) of each line were subsequently immunoblotted with antibodies directed against the N-terminus of the SP-C proprotein or the mature SP-C peptide, to assess relative protein expression (Figure 1B). Despite comparable SP-C mRNA concentrations, proSP-C expression varied greatly among the four cell lines (Figure 1B). SP-C<sup>WT</sup> proprotein was considerably more abundant than SP-C<sup>Δ<exon4></sup> proprotein, which was previously shown to be rapidly degraded by the proteasome (7, 17). SP-C<sup>Δ<exon11</exon12></sup> mutant proprotein was also more stable than SP-C<sup>Δ<exon4></sup> proprotein and similar to SP-C<sup>WT</sup> proprotein. Notably, SP-C<sup>Δ<exon7</exon7></sup> proprotein accumulated to a much greater extent than either SP-C<sup>WT</sup> or the other two mutant proproteins. The abundance of the SP-C<sup>Δ<exon7</exon7></sup> mutant was particularly striking when all forms of the proprotein were considered, including two unique bands recently identified by Beers and colleagues (20), composed of a prominent processing intermediate (Figure 1B, white arrowhead) and a slower-migrating form of the proprotein (Figure 1B, black arrowhead, and Figure 2C). Thus the stability of proSP-C varies widely with the mutation: SP-C<sup>Δ<exon4></sup> accumulates, SP-C<sup>Δ<exon4>Δ<exon4></sup> is rapidly degraded, and SP-C<sup>Δ<exon11</exon12></sup> exhibits an intermediate stability similar to that of SP-C<sup>WT</sup>.

The expression of SP-C<sup>WT</sup> in cells lacking lamellar bodies, such as HEK293 cells, results in the anterograde trafficking of the proprotein to the lysosome, where it is processed to the mature peptide (19) (Figure 1B, center). In contrast, the SP-C<sup>Δ<exon11</exon12></sup> and SP-C<sup>Δ<exon4>Δ<exon4></sup> mutants are ERAD substrates that are retained in the ER and degraded by the proteasome (17). The Western blotting of cell lysates detected a small amount of mature peptide in cells stably expressing SP-C<sup>Δ<exon11</exon12></sup>, suggesting that some of the proprotein escaped ER quality control when constitutively expressed in HEK293 cells (Figure 1B, center). No mature peptide was identified in cells expressing SP-C<sup>Δ<exon4>Δ<exon4></sup> or SP-C<sup>Δ<exon7</exon7></sup>, although longer exposure of the blot detected some mature peptide in cells expressing SP-C<sup>Δ<exon7</exon7></sup> (not shown).

The block in processing of the SP-C<sup>Δ<exon7</exon7></sup> proprotein to the mature peptide, coupled with the accumulation of proprotein, suggested that the I73T mutation may impart increased stability to the proprotein. To test this hypothesis, pulse-chase experiments were performed in HEK293 cells stably expressing WT or mutant SP-C. SP-C<sup>WT</sup> was synthesized as a proprotein (Figure 2A, band 1) that was posttranslationally modified to a larger proprotein (Figure 2A, band 2), with an estimated t½ of 145 minutes. The conversion of band 1 to band 2 was incomplete and blocked by brefeldin A (Figure E1A in the online supplement),
consistent with proprotein modification in a compartment distal to the ER. In contrast, very little SP-C L188Q was converted to band 2 and the proprotein was rapidly degraded, with a t 1/2 of 43 minutes (Figure 2B). Band 1 of SP-C I73T was rapidly converted to band 2 and a higher molecular weight protein (Figure 2C, band 3). The formation of band 3 was inhibited by brefeldin A (Figure E1B) or by the mutation of isoleucine at position 73 to serine (I73S) or lysine (I73K) (Figure E1C). Both bands 2 and 3 of SP-C I73T were very stable and did not decrease over the 2-hour chase period (Figure 2C). Thus the I73T mutation is associated with decreased proprotein processing/degradation that leads to accumulation.

PBA Treatment Stabilizes SP-C Proprotein and Promotes Conversion to Mature Peptide

To determine if chemical chaperones would promote processing of the mutant proproteins to the mature peptide, cell lines were treated with PBA or glycerol. PBA was added at up to 2.5 mM and glycerol at up to 2.5% (higher doses of either reagent result in toxicity in all cell lines), and the cell lysates were probed for both proSP-C and mature SP-C. Because glycerol treatment did not alter SP-C concentrations, only the PBA data are reported here. PBA treatment did not significantly alter concentrations of SP-CWT proprotein or mature peptide (Figure 3A). In contrast, PBA treatment promoted a significant accumulation of SP-C L188Q and SP-CΔexon4 proproteins (Figure 3A, top). Proprotein stabilization was accompanied by a corresponding increase in mature peptide in cells expressing SP-C L188Q but not SP-C Δexon4 (Figure 3A, center). Although PBA did not alter SP-C I73T proprotein concentrations, treatment was associated with a small but significant increase in mature peptide. However, the proprotein/mature peptide ratio remained much higher for SP-C I73T than for SP-C L188Q or SP-C WT, consistent with the hypothesis that generation of the mature peptide is partly blocked in cells expressing SP-C I73T. Overall, PBA treatment resulted in the stabilization of the SP-C L188Q proprotein and conversion to mature peptide at concentrations approximating those of SP-C WT. Importantly, the effect of PBA was reproduced in three different HEK293 cell lines stably expressing SP-C L188Q (Figure E2A), as well as in mouse lung epithelial (MLE-12) cells transiently transfected with SP-CWT (Figure E2B).

The accumulation of mutant protein is frequently associated with the induction of ER stress. In this study, no evidence of ER stress was detected in any of the untreated SP-C cell lines (Figure 3B), consistent with our previous finding that the constitutive/stable expression of SP-C Δexon4 in HEK293 cells is not accompanied by ER stress (7). The absence of ER stress in cells expressing SP-C I73T suggested that the proprotein might accumulate in a compartment outside the ER (to be described later). PBA-induced ER stress (Figure 3B) may be related to the stabilization and retention of the proprotein in the ER that, in turn, allows more time for folding.

PBA Treatment Induces Detergent-Insoluble Aggregates of SP-C Proprotein

Mutant proSP-C has a high propensity to form intracellular SDS-insoluble aggregates. To assess aggregate formation, lysates of cells stably expressing SP-C WT or mutant SP-C were extracted.
with 1% SDS, and the detergent-insoluble fractions were analyzed by Western blotting. High molecular weight (> 250 kD) SP-C aggregates were consistently detected in SDS-insoluble fractions of all four cell lines, indicating that both mutant and WT SP-C can aggregate (Figures 4A and 4B). In the absence of PBA treatment, SDS-insoluble SP-C proprotein was much more abundant in cells expressing SP-CI73T (Figure 4A) compared with cells expressing SP-CWT, SP-CL188Q (Figures 4A and 4B), or SP-CΔexon4 proprotein (Figure 4B, right). We note that the aggregates were located at the top of the gels, leaving open the possibility that not all the sample entered the gel, leading to an underestimation of the amount of SDS-insoluble SP-C. These data indicate that SP-C I73T is the most aggregate-prone of the three mutant proteins tested in this study, and that the total amount of SP-CI73T proprotein (detergent-soluble + detergent-insoluble) is greater than initially estimated (Figure 1B). Treatment with PBA promoted the accumulation of SDS-insoluble SP-C aggregates in a dose-dependent manner. This effect was most pronounced for SP-CI73T and SP-CΔexon4 after the extended exposure of Western blots (Figure 4B). At the highest dose of PBA, the amount of detergent-insoluble SP-CI73T was similar to that observed for SP-CΔexon4 proprotein (Figure 4A). Thus PBA treatment increases both soluble and insoluble forms of SP-CΔexon4.

**SP-CI73T and SP-CL188Q Traffic through Distinct Intracellular Pathways**

The detection of mature peptide in cells expressing SP-CWT or SP-CΔexon4 (Figure 1) is consistent with the trafficking of these proproteins to the endolysosomal pathway. Treatment with the lysosomotropic drugs bafilomycin A1 (Figure 5) or chloroquine (not shown) resulted in the accumulation of proprotein and inhibited the conversion to mature peptide, further supporting this conclusion. Treatment with azithromycin, a drug recently shown to inhibit lysosomal acidification (21), produced similar effects on proprotein stabilization and processing (not shown). In contrast, SP-CΔexon4 proprotein was not affected, and the mature peptide was barely detectable. This result suggests that, unlike SP-CWT and SP-CI73T, relatively little SP-CΔexon4 proprotein traffics to the lysosome. Treatment with lysosomotropic drugs also exerted no effect on SP-CΔexon4, likely because this proprotein is rapidly degraded by the proteasome and is not exported from the ER.

The differential response of SP-CΔexon4 and SP-CΔexon3 to PBA and lysosomotropic drugs likely reflects distinct intracellular trafficking patterns for the two mutant proproteins. Unlike SP-CΔexon4, SP-CΔexon3 did not accumulate in response to proteasome inhibition (not shown), and did not engage the ERAD chaperones endoplasmic reticulum–localized DnaJ homologue 4 (ERdj4) or ERdj5 (Figure E3A). Further, unlike SP-CΔexon4, the transient expression of SP-CΔexon3 did not induce ER stress (Figure E3B). These data, coupled with the finding that brefeldin A blocks the formation of band 3 in SP-CΔexon3 (Figure E1B), indicate that SP-CΔexon3 passes ER quality control, resulting in rapid export to the Golgi apparatus. Confocal microscopy with an antibody directed against the C-terminal (luminal) domain of proSP-C confirmed that SP-CΔexon3 colocalized with wheat germ agglutinin (WGA) at the surface of nonpermeabilized cells (Figure 6), as recently reported (20). Immunogold labeling further confirmed the cell-surface localization of SP-CΔexon3 and the absence of proprotein on the surface of cells expressing SP-CWT (Figure E4). Little or no colocalization of SP-C with WGA was detected in cells expressing SP-CWT, SP-CΔexon3, or SP-CΔexon4 (Figure 6). To assess the turnover of cell-surface SP-C, cells stably expressing SP-CΔexon3 were grown in the presence of doxycycline to suppress the expression of SP-C. Confocal microscopy confirmed the extended residency of SP-CΔexon3 at the cell surface...
These results suggest that an accumulation of SP-C<sup>I73T</sup> in the plasma membrane likely accounts for the unusual stability of the proprotein and the inhibition of proprotein conversion to mature peptide.

**DISCUSSION**

In this study, we used the stable expression of constructs encoding WT or mutant SP-C to model the constitutive expression of SP-C<sub>in vivo</sub>. The selection of cell lines that express similar concentrations of SP-C mRNA revealed that proprotein stability varied widely with the SFTPC mutation. Unexpectedly, both detergent-soluble and insoluble forms of SP-C<sup>I73T</sup> accumulated to a much greater extent than did the Brichos mutants, SP-C<sup>L188Q</sup> and SP-C<sup>Δexon4</sup>. SP-C<sup>I73T</sup> proprotein did not engage the ERAD chaperones previously reported to be involved in the turnover of Brichos mutants (17) or induce ER stress, as previously noted by Mulugeta and colleagues (22). These data, coupled with the results of pulse-chase experiments, suggest that SP-C<sup>I73T</sup> does not accumulate in the ER, but rather is rapidly folded and exported from the ER to the Golgi apparatus. For this reason, PBA treatment, which promotes the folding of some protein substrates in the ER, exerted no effect on SP-C<sup>I73T</sup>. Confocal microscopy detected SP-C<sup>I73T</sup> at the plasma membrane, confirming a recent report by Beers and colleagues (20). The loss of SP-C<sup>I73T</sup> from the plasma membrane occurred slowly, suggesting that cell-surface localization accounts for much of the accumulated proprotein. Consistent with this idea, relatively little proprotein was processed to mature peptide, an event that occurs in the endolysosomal pathway. Further, lysosomotropic drugs did not cause an accumulation of SP-C<sup>I73T</sup> proprotein. Collectively, these data suggest that the trafficking of SP-C<sup>I73T</sup> to the plasma membrane occurs rapidly, but that internalization and routing to the lysosome for processing/degradation constitute much slower events. Overall, the development of therapies for this “common” SFTPC mutation will require a better understanding of the molecular basis for SP-C<sup>I73T</sup> mistrafficking.

In contrast to SP-C<sup>I73T</sup>, SP-C<sup>Δexon4</sup> was the least stable of the mutant proproteins, and frequently required an overexposure of

(Figure E5). These results suggest that an accumulation of SP-C<sup>I73T</sup> in the plasma membrane likely accounts for the unusual stability of the proprotein and the inhibition of proprotein conversion to mature peptide.
immunoblots to detect its signal (Figure 1B). This result is in marked contrast to the reported tendency of SP-CΔexon4 to form aggresomes (23), a property that may well be related to the transient overexpression and saturation of the disposal pathway. The robust overexpression of SP-CΔexon4 in Type II epithelial cells of transgenic mice resulted in aggregation, cytotoxicity, and lung dysmorphogenesis (6). In the present study, the constitutive expression of SP-CΔexon4 at concentrations that did not induce ER stress resulted in the formation of very little detergent-insoluble proproteint particularly when compared with SP-CI73T. PBA treatment increased the amount of SP-CΔexon4 proprotein, but the concentration was still much lower than for SP-CWT, and was not accompanied by detectable processing to the mature peptide. Moreover, treatment with bafilomycin A1, chloroquine, or azithromycin did not cause proprotein accumulation, suggesting that little or no SP-CΔexon4 was trafficked to the endolysosomal pathway. Proteasome inhibition resulted in a robust accumulation of proprotein (7) and a stable association with ER chaperones (Figure E3A), confirming that SP-CΔexon4 is a canonical ERAD substrate. Taken together, these results suggest that this SFTPC mutation leads to severe misfolding, and that the rescue of the mutant proprotein by treatment with chemical chaperones will be very challenging.

Like SP-CΔexon4, SP-CL188Q is also an ERAD substrate (Figure E3A) (17). The constitutive expression of SP-CL188Q at concentrations that did not cause ER stress resulted in the trafficking of some proprotein to the endolysosomal pathway, as indicated by our detection of mature peptide and the accumulation of proprotein in response to bafilomycin A1, chloroquine, or azithromycin. Treatment with concentrations of PBA widely used in other in vitro studies (10, 24–26) promoted the export of SP-CL188Q from the ER, leading to increased proprotein concentrations and the enhanced conversion of proprotein to mature peptide. At a dose of 1 mM PBA, the mature peptide concentration in SP-CL188Q–expressing cells approximated that in SP-CWT cells, with a minimal aggregation of proprotein. This dose is similar to that used in two clinical trials in which the concentration of PBA in serum ranged from 0.04–2.0 mM (27, 28). Collectively, these results suggest that this SFTPC mutation

Figure 5. Trafficking of SP-C to the Golgi apparatus and endolysosomal compartments. (A) Cell lines were grown in the absence of doxycycline and treated with bafilomycin A1 for 24 hours. Cell lysates (30 μg) were analyzed by SDS-PAGE/Western blotting. Blots were sequentially probed for proSP-C (top), mature SP-C peptide (middle), and actin (bottom). (B) Quantitation of the effects of bafilomycin A1 on proSP-C (n = 3) and mature peptide concentrations (n = 2).

Figure 6. SP-CI73T is localized to the cell surface. Cells were grown on coverslips in the absence of doxycycline, fixed in situ with 4% paraformaldehyde, and processed for confocal microscopy without permeabilization. Cells were incubated with antibody directed against the C-terminus of proSP-C (top row) or wheat germ agglutinin (WGA) to label the cell surface (middle row). No signal was detected when cells were incubated with antibody directed against the N-terminus of proSP-C, confirming that the Brichos domain is extracellular (not shown).
is amenable to rescue by PBA, and may offer a viable treatment option for patients.

The results of the present study provide support for the screening of other Brichos domain mutants to determine which mutants may be amenable to PBA therapy. However, this study also underscores the need for further investigation of the long-term effects of PBA treatment. PBA caused an increase in SDS-insoluble proprotein in cells expressing SP-C(L188Q) Mutant proSP-C, including SP-C(L188Q), can form amyloid-like fibrils (15) related to the loss of chaperone activity of the Brichos domain (14, 29) and subsequent transition of the α-helical transmembrane domain to a β-sheet structure (30, 31). Artificial β-sheet proteins that form amyloid-like fibrils have been shown to sequester proteins required for critical cellular functions (32). In the present study, the accumulation of SDS-insoluble proprotein was not associated with cytotoxicity (data not shown). However, the effect of extended PBA treatment on SP-C(L188Q) aggregation/cytotoxicity remains to be evaluated. Overall, analyses of PBA treatment in mice expressing SP-C(L188Q) (currently in development) will provide important insight into the risks and benefits of this potential therapy.

In conclusion, PBA treatment substantially restored intracellular trafficking and processing of the SP-C(L188Q) proprotein to its mature peptide. Whether this correction will be sufficient to prevent or slow pathogenesis remains to be established, as do any potential adverse effects of prolonged PBA treatment. Given the propensity of all SP-C proproteins, including SP-C(W1, to form detergent-insoluble aggregates, the use of drugs promoting the simultaneous proprotein stabilization and inhibition of processing, such as chloroquine and azithromycin (33, 34), should be carefully reevaluated.

Author disclosures are available with the text of this article at www.atsjournals.org.

References

