Efficient suppression of endogenous CFTR nonsense mutations using anticodon-engineered transfer RNAs

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INTRODUCTION

Nonsense mutations or premature termination codons (PTCs) comprise ~11% of all genetic lesions, which result in over 7,000 distinct genetic diseases. Due to their outsized impact on human health, considerable effort has been made to find therapies for nonsense-associated diseases.Suppressor tRNAs have long been identified as a possible therapeutic for nonsense-associated diseases; however, their ability to inhibit nonsense-mediated mRNA decay (NMD) and support significant protein translation from endogenous transcripts has not been determined in mammalian cells. Here, we investigated the ability of anticodon edited (ACE)-tRNAs to suppress cystic fibrosis (CF) causing PTCs in the cystic fibrosis transmembrane regulator (CFTR) gene in gene-edited immortalized human bronchial epithelial (16HBEge) cells. Delivery of ACE-tRNAs to 16HBEge cells harboring three common CF mutations G542XUGA-, R1162XUGA-, and W1282XUGA-CFTR PTCs significantly inhibited NMD and rescued endogenous mRNA expression. Furthermore, delivery of our highly active leucine-encoding ACE-tRNA resulted in rescue of W1282X-CFTR channel function to levels that significantly exceed the necessary CFTR channel function for therapeutic relevance. This study establishes the ACE-tRNA approach as a potential standalone therapeutic for nonsense-associated diseases due to its ability to rescue both mRNA and full-length protein expression from PTC-containing endogenous genes.

Nonsense mutations are single-nucleotide mutations that convert a canonical amino acid codon to one of the three stop codons (UAA, UAG, and UGA). Nonsense mutations result in a premature termination codon (PTC) generally with almost complete loss of function of the affected protein. This is due to both the PTC terminating translation prematurely, resulting in a truncated protein, and nonsense-mediated mRNA decay (NMD) degrading the mRNA coding for the PTC-containing protein.1 With the advent of next-generation sequencing, more than 7,500 PTCs in nearly 1,000 different human genes have been discovered. PTCs account for close to 11% of all described mutations leading to inherited human disease.2 An abridged list of the known PTC-associated disease phenotypes includes Duchenne muscular dystrophy,3 spinal muscular atrophy,4 β-thalassemia,5 Hurler syndrome,6 Usher syndrome,7 and cystic fibrosis.8 Cystic fibrosis (CF), a common autosomal recessive genetic disease, is caused by mutations to the CF transmembrane conductance regulator (CFTR) gene.9–11 Similar to genetic diseases as a whole, PTCs account for close to 10% of mutations causing CF.12 Several therapeutic small molecules rescue clinically relevant levels of CFTR function for a growing percentage of people with CF. However, all CF therapeutic small molecules so far target full-length aberrant CFTR protein.13–15 As PTCs result in little to no full-length protein, these small molecules targeting the CFTR protein are ineffective at rescuing PTC-containing CFTR. Instead, PTC therapies must target the defective gene, post-transcriptional processing, or processes governing protein translation.

Shortly after the discovery of the CFTR gene,9–11 proof-of-concept for gene replacement was demonstrated.16,17 In the following 30 years, significant effort has been directed toward CFTR gene-replacement therapy, as this method has potential to repair the phenotypic defect for all people with CF. A number of viral and non-viral gene delivery approaches have been investigated.18 Improvements in vector design and delivery methods, a deeper understanding of lung biology and disease development, and new animal models have gotten us closer to the development of an effective mutation agnostic gene therapy for CF; however, to date, no gene-replacement therapy has made it to widespread clinical use.18 CRISPR-Cas-based technologies, including prime editing, homologous recombination following double-strand breaks, and in-frame deletions, have received considerable interest as potential nonsense mutation interventions.19–21 As with total gene replacement, the CRISPR-Cas approach requires efficient delivery, optimally to

progenitor cells, to ensure long-term repair of the nonsense mutation. Delivery of CRISPR-Cas both as an adeno-associated virus (AAV) encoding the CRISPR-Cas pair and as a CRISPR-Cas ribonucleoprotein delivered with engineered amphiphilic peptides has been demonstrated in model systems. A drawback to CRISPR-Cas approaches is that design of a new guide RNA is required for each mutation site. As with gene therapy, an efficient delivery method for clinical use has not been determined.

Small molecules, including aminoglycosides, synthetic aminoglycosides, and oxodiazoles, have been reported to act as nonsense readthrough agents. Pharmacological treatment resulting in PTC readthrough is an attractive therapeutic avenue to pursue, as this promises to allow for body-wide readthrough of PTCs. Small-molecule PTC readthrough agents generally bind to the ribosome and reduce translational fidelity. While this allows for suppression of PTCs by near-cognate tRNAs, global translational fidelity is also decreased. Of note, several of these small-molecule PTC readthrough agents have been investigated in clinical trials with generally low efficacy and strong ototoxic and nephrotoxicity. In addition, a recent study has also indicated that genome-wide natural termination codon (NTC) readthrough stimulated by the aminoglycoside G418 in vivo results in the disruption of several biological processes with detrimental cellular effects. While the promises of small-molecule nonsense readthrough agents are tantalizing, due to issues with safety, efficacy, or both, their clinical implementation has remained out of reach.

Another nonsense-suppression technology that has been explored previously is the use of nonsense-suppressor tRNAs or anti-codon-edited tRNAs (ACE-tRNAs). A recently developed library of ACE-tRNAs with the anticodons engineered via mutagenesis to suppress one of the UAA, UAG, or UGA PTC codons demonstrated efficacious in vivo PTC suppression. Previous work by ourselves and others has demonstrated the safe and efficacious ability of ACE-tRNAs to rescue PTCs in the context of cDNAs, it is still unknown whether they can take a major step forward in their therapeutic potential and rescue PTCs in a native genomic context. Heterologous overexpression of PTC-containing mRNAs as cDNAs has fundamentally limited relevance, as these constructs lack the genomic context and chromatin environment, native protein promoter, intronic sequences, and post-transcriptional processing.

To that end, we set out to demonstrate ACE-tRNA-based rescue of PTC-containing CFTR from the well-characterized, immortalized isogenic human bronchial epithelial (16HBE14o-) cell lines modified with CRISPR-Cas9 to contain the most common CF-causing PTC variants. Here, we show significant rescue of PTC-containing CFTR mRNA transcripts by real-time quantitative reverse transcription polymerase chain reaction (qRT-PCR) for three of the most common CF-causing PTCs (G542XUGA [2.5%], W1282XUGA [1.2%], and R1162XUGA [0.4%]) and significant rescue of CFTR protein function by patch-clamp electrophysiology following delivery of our best-performing ACE-tRNA. The level of ACE-tRNA-dependent rescue of functional W1282X-CFTR protein demonstrated here is well above the predicted mark for therapeutically meaningful rescue. While efficient delivery of ACE-tRNAs to affected cell types remains a hurdle to their therapeutic implementation, ACE-tRNAs display several attractive qualities as gene therapy agents, bolstering their therapeutic promise. The results presented here display an exciting step forward in an unmet medical need for patients with nonsense mutations.

RESULTS
Development of a nonsense-suppression reporter system
While nonsense mutations cause several serious genetic diseases, here, we chose to focus on CF, using the 16HBE14o- cell line and the derivative 16HBE-gene-edited (16HBEge) cell lines with engineered R1162X-, W1282X-, or G542X-CFTR variants. The genome of each of these 16HBEge cell lines was modified via CRISPR-Cas9 to encode a UGA stop codon at the amino acid residue specified in the CFTR gene. This model retains the native CFTR transcriptional, post-transcriptional, and translational regulation. We first set out to understand which previously identified nonsense-suppressor ACE-tRNAs, whether delivered as DNA or RNA, were most effective. We also wanted to determine their persistence of activity and whether previous results were recapitulated in human epithelial cells. To do this, we wanted a flexible high-throughput method to assess nonsense suppression in different cell lines, which could be transferred to any cell line of interest for future studies. Here, we chose to use the piggyBac (PB) transposon/transposase system, which integrates the gene of interest (or cargo) specifically into genomic TTAA sites, facilitates fast selection of stable cell lines, can be used to generate both cell lines and animal models, and for which a hyperactive transposase has been generated (Figure 1A). We used a PTC-containing NanoLuc (NLuc) luciferase as a nonsense-suppression reporter due to high signal to noise and utility in high-throughput assays. We sought an assay to determine the potentially short half-lives of ACE-tRNA therapeutic agents (i.e., delivered as RNA); for this, we appended the protein destabilizing PEST domain (Figure 1B). The PEST domain has been previously shown to shorten the half-life of NLuc to ~20 min in several cell lines. Traditional NLuc assays require cell disruption, as NLuc is retained intracellularly. By appending the interleukin-6 (IL-6) secretion signal to the NLuc reporter (Figure 1B), we developed a method to assay nonsense suppression longitudinally without disrupting cells. For longitudinal nonsense-suppression studies, the cell media are assayed for PTC-containing Sec-NLuc luminescence (Figure 1B). Further, our best performing ACE-tRNA, ACE-tRNA, and ACE-tRNA library members displayed robust nonsense suppression for all three NLuc formats in 16HBE14o- cells, both as a transient co-transfection (Figures 1C–1E and S1A–S1C) and following reporter stable integration (Figures 1F and 1D). Plasmids containing four copies of the ACE-tRNA expression cassette (4×; Figure 1) were used, as previous work by ourselves and others has indicated increased nonsense-suppression efficiency with multiple nonsense-suppressor tRNAs per plasmid. These results closely track what was displayed previously...
Consistent with our previous work, the ACE-tRNA\textsubscript{Trp} possesses nearly an order of magnitude lower nonsense-suppression activity (Figures 1C–1E) than the other ACE-tRNAs shown here.\textsuperscript{30} ACE-tRNA nonsense-suppression activity relies on the action of endogenous aminoacyl-tRNA synthetases (RS) to charge the ACE-tRNA with its cognate amino acid. We have noted that, while ACE-tRNA\textsuperscript{Trp}\textsubscript{UGA} possesses poor nonsense-suppression activity, ACE-tRNA\textsuperscript{Trp}\textsubscript{UAG} possesses an order of magnitude higher suppression activity. The anticodon of tRNA\textsuperscript{Trp} has been noted to be recognized by the TrpRS.\textsuperscript{37} Likely, the altered anticodon of the ACE-tRNA\textsuperscript{Trp}\textsubscript{UGA} negatively impacts interactions between the tRNA and the TrpRS more than that of the altered anticodon of the ACE-tRNA\textsuperscript{Trp}\textsubscript{UAG}. However, we cannot rule out differences in transcription between these tRNAs or interactions with other parts of the translational apparatus causing the differences seen between the PTC suppression activity of these ACE-tRNAs. If, as we suspect, the interactions between the ACE-tRNA\textsuperscript{Trp}\textsubscript{UGA} and the TrpRS are disrupted, this would lead to a lower intracellular pool of aminoacylated ACE-tRNA\textsuperscript{Trp}\textsubscript{UGA}, which in turn is not sufficient to suppress PTCs on the ribosome to the same extent as our other ACE-tRNAs. With the poor performance of the ACE-tRNA\textsuperscript{Trp}\textsubscript{UGA} in mind, we included in our investigation the best performing ACE-tRNA\textsubscript{Leu} for rescue of W1282X, as it possesses similar activity to our other more robust ACE-tRNAs (see further discussion below).

PTC suppression by ACE-tRNA encoded in plasmid peaks 48 h following delivery to 16HBE\textsubscript{ge} cells

To assess the kinetics of nonsense-suppression rescue, we used R1162X- and W1282X-CFTR 16HBE\textsubscript{ge} cells stably integrated with PB transposon containing Sec-NLuc UGA. The cells were seeded on Transwells and transfected with a plasmid containing 4 × of our top performing ACE-tRNA\textsuperscript{Arg}, ACE-tRNA\textsuperscript{Trp}, and ACE-tRNA\textsuperscript{Leu} UGA suppressors or an empty plasmid (control). Following transfection, the Sec-NLuc luminescence was assayed daily from media collected from both the apical and basolateral sides of the Transwell. We chose to utilize cells cultured on semipermeable membranes in Transwell chambers, as Sec-NLuc can be utilized to track ACE-tRNA function longitudinally with the Transwells ultimately used in Ussing chamber experiments to measure short-circuit conductance from CFTR rescue. ACE-tRNA\textsuperscript{Arg} supported robust rescue of Sec-NLuc luminescence peaking around 2 days after transfection, in contrast to ACE-tRNA\textsuperscript{Trp}, which performed relatively poorly (Figures 2A–2D). Indeed, the result with ACE-tRNA\textsuperscript{Trp} is not altogether unexpected, as it was the poorest performing ACE-tRNA in all of the most active ACE-tRNAs identified in our screen.\textsuperscript{30} The W1282L mutant of CFTR has been previously shown to support near-wild-type (WT) channel function and displayed a similar C/B band ratio on western blots, indicating that the W1282L CFTR is trafficked and exhibits similar stability to WT CFTR.\textsuperscript{38} In the process of CFTR expression and maturation, the B band corresponds to the core-glycosylated CFTR that resides in the endoplasmic reticulum (ER), and the C band represents the mature, fully processed CFTR that has passed through the Golgi apparatus. The C/B band ratio indicates the amount of fully folded, matured, and processed CFTR protein to the amount of immature CFTR protein, with low ratios displayed for poorly folded CFTR mutants with low stability. These previous findings prompted us to test our highly performing ACE-tRNA\textsuperscript{Arg} for rescue of Sec-NLuc UGA in this assay. As with ACE-tRNA\textsuperscript{Arg}, ACE-tRNA\textsuperscript{Leu} supported robust rescue of Sec-NLuc luminescence peaking 2 days after transfection (Figures 2A–2D). Based on the results of this longitudinal study, it is likely that the decrease in luminescence rescue is based both on silencing and dilution of plasmids due to cell splitting.\textsuperscript{39,40} Importantly, the luminescence-based assay informed us on when to perform more in-depth studies on CFTR mRNA expression and channel function described in this study.

ACE-tRNAs rescue mRNA expression from endogenously encoded nonsense mutations

The CFTR transcripts in R1162X- and W1282X-CFTR 16HBE\textsubscript{ge} cells are sensitive to degradation through the NMD pathway.\textsuperscript{1,7} To that end, we determined the level of NMD inhibition elicited by transfection of ACE-tRNA plasmids. Total RNA was isolated from R1162X- and W1282X-CFTR 16HBE\textsubscript{ge} cells 2 days after transfection of either 1× or 4× ACE-tRNA\textsuperscript{Arg}, ACE-tRNA\textsuperscript{Trp}, and ACE-tRNA\textsuperscript{Leu} plasmids. We then performed real-time qRT-PCR with probes against CFTR and control TATA binding protein (TBP). ACE-tRNA\textsuperscript{Arg} and ACE-tRNA\textsuperscript{Leu}, when delivered as plasmids containing 1×, rescued significant amounts of CFTR mRNA (34.6% ± 1.0% and 38.5% ± 0.8% of WT, respectively) as compared with cells transfected with an empty plasmid (11.9% ± 0.3% and 14.8% ± 0.4% of WT, respectively; Figure 2E). Cells transfected with 4× ACE-tRNA plasmids also rescued significant amounts of CFTR mRNA (38.8% ± 1.7% and 52.5% ± 0.9% of WT, respectively), and to a greater extent than 1× ACE-tRNA plasmids for ACE-tRNA\textsuperscript{Leu}, suggesting multiple ACE-tRNAs within each delivered vector is beneficial in non-saturating conditions (Figure 2E), which is consistent with previously reported results.\textsuperscript{41} ACE-tRNA-dependent increase in CFTR mRNA expression is likely due to inhibition of NMD-dependent mRNA degradation by suppressing the PTC during pioneer rounds of mRNA translation.\textsuperscript{1} In contrast to ACE-tRNA treatment, G418 (100 μM) treatment significantly decreased WT CFTR mRNA abundance in 16HBE14o- cells (79.5% ± 1.1% of vehicle-treated 16HBE14o- cells), which is similar to a previously reported study.\textsuperscript{32} Reduction of WT CFTR mRNA expression following G418 treatment is likely due to global readthrough of NTCs and general translation miscoding,\textsuperscript{26} a feature that is not exhibited by ACE-tRNAs. Furthermore, treatment with G418 exhibited only a meager rescue of R1162X-CFTR mRNA expression (19.1% ± 0.5% of WT) compared with vehicle control (12.5% ± 0.8% of WT; Figure 2F) and did not significantly increase W1282X-CFTR mRNA expression. To our knowledge, this is the first time that delivery of nonsense-suppressor ACE-tRNAs has been directly shown to inhibit NMD and rescue mRNA from nonsense mutations in their endogenous genomic context. While we were pleased to see such robust mRNA rescue
from NMD by ACE-tRNAs, this result on the whole was not completely unexpected, as small-molecule nonsense readthrough agents have been previously shown to rescue CFTR mRNA.32

**ACE-tRNA Arg delivered as RNA suppresses nonsense mutations and rescues CFTR mRNA**

We next moved to ascertain whether ACE-tRNAs delivered as RNA could also suppress nonsense mutations in 16HBE
gen cells. As with plasmid delivery, we first wanted to determine the time course of rescue in cells for ACE-tRNA as RNA delivered by nucleofection. Use of the stable PB Sec-NLuc UGA reporter 16HBE
gen cell line displayed significant rescue at day 1 but decreased sharply at day 2 (Figures S2A–S2D). As we missed the initial increase in Sec-NLuc luminescence observed previously with cDNA delivery, we likely under-sampled and missed the peak PTC suppression time point for RNA delivery. With that in mind, we shifted to the NLuc-PEST assay, as NLuc with an appended PEST sequence has a half-life of about 20 min.36 Based on results from the NLuc-PEST-UGA assay, nonsense-suppression activity of ACE-tRNA Arg delivered as RNA peaks at ~5 h and decays exponentially over the next 25 h, exhibiting a half-life of ~6.5 h (Figures 3A and S2E). Using this time course as a guide, we assayed the ability of ACE-tRNA Arg delivered as RNA to inhibit NMD at both 6 and 24 h post-nucleofection. We found R1162X-CFTR mRNA expression was significantly rescued by ACE-tRNA Arg RNA (31.9% ± 3.7% of WT) compared with mock (no tRNA) nucleofections (13.0% ± 0.6% of WT) at 6 h post-nucleofection; however, this effect was not observed at 24 h post-nucleofection (Figure 3B). ACE-tRNA delivered as RNA convincingly suppresses endogenous nonsense mutations; however, to become a viable therapy, we will likely need to extend the half-life of ACE-tRNA activity or perform frequent delivery. It also may be the case that the relatively short ACE-tRNA RNA half-life may be a product

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**Figure 1. NLuc PTC reporter platforms faithfully report PTC suppression efficiency of ACE-tRNAs in 16HBE14o- cells**

(A) Schematic illustrates the piggyBac (PB) system used in this study. Red octagons denote the presence of an in-frame PTC in the NLuc reporter. (B) Three types of NLuc PTC reporters were generated for reporting for UGA PTC suppression efficiency of ACE-tRNAs: NLuc, NLuc with protein destabilization domain (NLuc-PEST), and NLuc with secretion signal (Sec-NLuc). The wild-type NLuc PTC reporter allows for high-throughput screening of different ACE-tRNAs, the NLuc-PEST PTC reporter with short intracellular lifetime allows for the measurement of the half-life of ACE-tRNA, and the Sec-NLuc PTC reporter with secretion capacity allows for longitudinal studies. (C–F) In WT 16HBE14o- cells co-transfected with ACE-tRNAs and PTC reporters, four highest performing ACE-tRNAs suppressing UGA PTCs exhibit a similar trend of luminescence rescue for (C) NLuc, (D) NLuc-PEST, and (E) Sec-NLuc reporters. (F) The NLuc-PEST PTC reporter stably integrated in WT 16HBE14o- cells using the PB system are capable of reporting PTC suppression efficiency of transiently transfected ACE-tRNAs through luminescence. Data are presented as normalized mean luminescence ± SEM. n = 3–7, with each replicate represented as a circle. Raw luminescence of described experiments is presented in Figure S1.
Figure 2. ACE-tRNAs delivered as plasmids have transient PTC-suppression activity that significantly and robustly rescues endogenous R1162X- and W1282X-CFTR mRNA expression

(A and B) R1162X-CFTR and W1282X-CFTR cells stably expressing Sec-NLuc UGA reporter (R1162X PB Sec-NLuc UGA and W1282X PB Sec-NLuc UGA cells, respectively) were transfected with empty or 4x ACE-tRNA plasmids, and luminescence measurements were made every 24 h from media for 5 consecutive days. Transfection of (A and B) ACE-tRNAArg (solid red, n = 3), (C and D) ACE-tRNALeu (dark blue, n = 3), and (C and D) ACE-tRNATrp (light blue, n = 3) plasmids results in time-dependent PTC suppression, with the peak luminescence detected at 2 days post-transfection, in both the apical (solid lines) and basolateral (dashed lines) sides of Transwells. (A–C) Transfection of empty vectors (gray lines, n = 3) results in negligible luminescence change over the course of 5 days. Data are shown in raw luminescence. (A and C) Raw luminescence and (B and D) luminescence normalized to cells receiving empty plasmids (ACE-tRNA/Empty) display similar time course of ACE-tRNA PTC-suppression activity. (E) CFTR mRNA abundance normalized to TBP (housekeeping gene) in WT (white, n = 3), R1162X-CFTR (red, n = 6), and W1282X-CFTR (blue, n = 6) cells 48 h following (legend continued on next page)
of the target cells, where an extended ACE-tRNA RNA half-life may be observed in cells that are not actively undergoing mitosis (i.e., airway epithelial cells in vivo). Transient delivery of ACE-tRNAs as both cDNA and RNA resulted in robust NMD inhibition at early time points. This time course of PTC suppression activity is incompatible with Ussing chamber measurements of ACE-tRNA-dependent rescue of CFTR channel activity, as on average, a minimum of 4 days is needed for formation of tight junctions. We therefore pursued the possibility of generating 16HBEge cells with stably integrated ACE-tRNA expression cassettes for persistent PTC activity.

Stable genomic integration of ACE-tRNAArg rescues significant amounts of CFTR mRNA and channel function in R1162X-CFTR 16HBEge cells

Use of NLuc reporters allowed us to determine with ease a short time of action in both cDNA and RNA, likely due to rapid cell splitting of this model and dilution of ACE-tRNA plasmid. To circumvent this issue, we generated R1162X-CFTR 16HBEge cells containing stably integrated ACE-tRNAArg expression cassettes. For this purpose, we cloned 16 copies of ACE-tRNAArg along with a puromycin resistance marker into a PB transposon (Figure S3A). R1162X-CFTR 16HBEge cells stably expressing PB ACE-tRNAArg were generated in a similar manner to the stable PB NLuc nonsense reporters described in Figure 1 (see materials and methods). Of note, all studies with R1162X-CFTR 16HBEge cells stably integrated with 16 copies of ACE-tRNAArg transposon (R1162X PB 16× ACE-tRNAArg cells) were done from the total population of puromycin-resistant cells that started from >100 colonies. We chose this approach to eliminate issues with genome insertion position effects of isogenic cell lines, that started from >100 colonies. We chose this approach to eliminate issues with genome insertion position effects of isogenic cell lines, which could pose issues with deconvoluting cell-to-cell variability. Given our ability to select 16HBEge cells stably expressing ACE-tRNAArg, it assuages concerns that persistent expression of nonsense-suppressor tRNAs at therapeutic levels will result in toxicity. Indeed, we observed that cells selected with puromycin exhibited similar morphology and cell division rates over >20 passages as compared with 16HBEge and 16HBE14o- cells, demonstrating that persistent ACE-tRNA expression is not toxic. To confirm that the stably incorporated ACE-tRNAs were functional, we transfected R1162X PB 16× ACE-tRNAArg cells with an NLuc UGA reporter construct. As expected, R1162X PB 16× ACE-tRNAArg cells exhibited significant (~60-fold) NLuc PTC suppression (Figure 4A, filled red bar) as compared with stable R1162X-CFTR 16HBEge cells generated with a transposon containing puromycin resistance without ACE-tRNAs (R1162X PB Empty; Figure 4A, open red bar).

We performed real-time qRT-PCR of total RNA isolated from R1162X PB 16× ACE-tRNAArg cells, noting a significant albeit modest ∼7% increase (16.8% ± 1.2% of WT) in CFTR mRNA as compared with mRNA from R1162X-CFTR 16HBEge cells results in significant rescue of CFTR mRNA expression

Figure 3. Delivery of ACE-tRNAArg as RNA to R1162X-CFTR 16HBEge cells results in significant rescue of CFTR mRNA expression

(A) R1162X-CFTR cells stably expressing NLuc-PEST UGA reporter (R1162X PB PEST-NLuc UGA cells) were nucleofected with and without ACE-tRNAArg RNA. Nonsense suppression mediated by ACE-tRNAArg was detected by luminescence measurements at 2, 3, 4, 6, 8, 10, 12, 18, 24, and 30 h post-nucleofection. Based on the average decay phase of luminescence by subtracting luminescence measurement with no delivered tRNA from that with ACE-tRNAArg RNA (red dotted line, y = 11,413 e−0.11x), half-life of ACE-tRNAArg is 6.5 ± 0.3 h. Raw luminescence measurements are presented in Figure S2E. (B) After 6 h (left) of delivery, ACE-tRNAArg (red filled) delivered as RNA in R1162X-CFTR cells is sufficient to significantly rescue R1162X-CFTR mRNA expression as determined by real-time qRT-PCR, from no RNA control (red open). No significant rescue of R1162X-CFTR mRNA expression was observed at 24 h (right). Data are presented as average ± SEM. All experiments contain an n = 3. Significance was determined by unpaired t test, where *p < 0.01.
Figure 4. Stable integration of ACE-tRNAArg expression cassettes in R1162X-CFTR cells significantly rescues endogenous CFTR mRNA expression and CFTR channel function

(A) Raw luminescence measurement following transfection of the NLuc UGA reporter plasmid in R1162X-CFTR cells integrated with empty transposon (R1162X PB Empty; open red bar, n = 3) and R1162X-CFTR cells stably integrated with 16x/C2 ACE-tRNAArg transposon (R1162X PB 16x ACE-tRNAArg; filled red bar, n = 3) results in significant

(legend continued on next page)
determined the extent of R1162X-CFTR functional channel rescue. To do so, we used Ussing chamber recordings of CFTR chloride short-circuit currents. Representative traces display a forskolin and 3-isobutyl-1-methylxanthine (IBMX) response over baseline, and this response is blocked by CFTRinh-172 (Figure 4C), with average forskolin/IBMX and CFTRinh-172 responses of CFTR channel function from R1162X PB 16× ACE-tRNAArg cells ranging from 5% to 7% of WT CFTR channel function from 16HBE14o- cells (Figure 4D). This level of rescued channel function exceeds that seen following treatment of 16HBE14o cells with the reference readthrough compound G418 in combination with the NMD inhibitor SMG1i (<5%) for several CFTR nonsense variants. While this indicates robust function of ACE-tRNAArg, we wanted to determine the ACE-tRNA copy number dependence of the CFTR rescue more precisely.

Using the R1162X PB 16× ACE-tRNAArg cells, we can now determine for the first time how many ACE-tRNAArg expression cassettes are needed to support significant rescue of CFTR mRNA expression and channel function. Previous studies demonstrated on average 1–3 PB transposon integrations per genome. To count the average number of ACE-tRNAArg transposon integration events per cell, we used qPCR to compare signals from the integrated transposon to standard curves generated from known quantities of target cDNA. The RPP30 gene, subunit P30 of RNaseP enzyme complex, was used as a genomic DNA (gDNA) internal control, as this is a widely accepted one-copy gene of the haploid human genome. gDNA from R1162X PB 16× ACE-tRNAArg cells was isolated, and 10 ng was used as a template for real-time qPCR. The cycle threshold (Ct) values obtained from primers and probes specific for PB puro (PB transposon puromycin resistance marker) and the RPP30 gene (RNase P) revealed that, on average, only approximately one copy of the PB transposon was integrated per genome, and therefore only 16 ACE-tRNAArg expression cassettes are integrated per genome (Figures 4F and 4G). Further experiments with 1 ng of gDNA or a probe targeted against the PB backbone using either 10 ng or 1 ng of gDNA revealed a similar average number of integration events (Figures S3B–S3D).

Indeed, we were excited to find that relatively few ACE-tRNA expression cassettes are required for significant rescue of both CFTR mRNA and channel function from endogenous CFTR genes with CF-causing PTCs, which provides further promise for use of ACE-tRNAs as therapeutics for nonsense-associated CF, as it reduces the burden of therapeutic delivery efficiency.

**ACE-tRNA expression yields significant rescue of CFTR mRNA expression in G542X-, R1162X-, and W1282X-CFTR 16HBE14o cells**

With the information that as few as 16 copies of ACE-tRNAArg support significant rescue of CFTR function, we revisited our experiments with transient transfection of plasmids encoding ACE-tRNAs, which do not recover more than 50% of the WT CFTR mRNA. Knowing that the transfection efficiency of 16HBE14o- cells is ~50%, we wanted to determine whether cDNA delivery is impacting our measurements of CFTR mRNA rescue. To account for transfection efficiency, we generated a transfection reporter plasmid containing both an EF1α-driven mNeonGreen (mNG) and four copies of either ACE-tRNAArg, ACE-tRNAArg, ACE-tRNALeu, or ACE-tRNAArg (Figure 5A). mNG-containing constructs permitted us to sort mNG-positive (Figure 5B, green curve) and negative (Figure 5B, gray curves) cells by fluorescence-activated cell sorting (FACS) 48 h after transfection and therefore determine ACE-tRNA-dependent NMD inhibition in a near-homogeneous population of cells that received ACE-tRNA-encoding plasmids. FACS analysis determined the percentage of green cells; those receiving ACE-tRNA plasmid, in fact, did not exceed 50% (Figures 5B and S4A; Table S1). ACE-tRNAArg mNG-positive G542X-, R1162X-, and W1282X-CFTR 16HBE14o cells exhibited significantly enhanced NMD inhibition over mNG-negative cells (Figure 5C). CFTR mRNA expression for ACE-tRNAArg was 41%, ACE-tRNAArg was 54%, and ACE-tRNAArg was 87% of WT CFTR. We were excited to see that ACE-tRNAArg provided strong inhibition of NMD, indicating that, when ACE-tRNAs are efficiently delivered to cells, they are exceedingly efficient at inhibiting NMD. The dependence of this result on transfection efficiency also highlights that ACE-tRNA delivery is a significant hurdle to their adoption as a therapeutic (see discussion).

**Whole-cell patch clamp reveals near-WT rescue of CFTR channel function from endogenous nonsense mutations with ACE-tRNAs**

Although we demonstrated significant and robust rescue of CFTR mRNA in G542X-, R1162X-, and W1282X-CFTR 16HBE14o cells, a rescue of 15–30% of CFTR channel function is ultimately required for a therapeutically relevant outcome. To this end, we transfected 4× ACE-tRNAArg mNG plasmids (Figure 5A) into W1282X-CFTR 16HBE14o cells and utilized the whole-cell, patch-clamp technique to directly measure CFTR channel function in mNG-positive cells 48–72 h after transfection (Figure 5D). To fully activate CFTR suppression activity of integrated ACE-tRNAArg, (B) R1162X-CFTR mRNA expression in R1162X PB 16× ACE-tRNAArg cells (filled red bar, n = 4) is significantly increased compared with R1162X PB Empty cells (open red bars, n = 4). (C) Representative short-circuit Cl− current traces recorded from WT (black trace), R1162X PB Empty (dashed red trace), and R1162X PB 16× ACE-tRNAArg (solid red trace) cells in response to forskolin (10 μM) and IBMX (100 μM), followed by CFTR inhibitor, CFTRinh-172 (20 μM). Average maximum ISC change in response to addition of (D) forskolin and IBMX and (E) CFTRinh-172 of WT 16HBE14o- (white bar, n = 21), R1162X PB Empty (open red bar, n = 14), and R1162X PB 16× ACE-tRNAArg (solid red bar, n = 9) cells is shown. (F) qPCR standard curves generated from 10-fold dilutions of PB 16× ACE-tRNAArg (red line) and RNase P (Hu_RP30) positive control; dashed blue; CDNA are shown. qPCR measurements of PB puro transgene (open red circle) and endogenous RNase P gene (open blue circle) from 10 ng input of gDNA from R1162X PB 16× ACE-tRNAArg cells are shown. Each sample was assayed in triplicate by qPCR. (G) 16× ACE-tRNAArg transgene copy number per genome in R1162X PB 16× ACE-tRNAArg cells is calculated by plotting Ct value to the standard curves. Data are presented as average ± SEM. Significance was determined by unpaired t test, where *p < 0.05, **p < 0.001, and ***p < 0.0001.

Figure 5. Expression of ACE-tRNAs results in significant and robust increase in CFTR mRNA and channel function

(A) Schematic of ACE-tRNA mNG plasmid. (B) Forty-eight hours following transfection of WT, G542X-, R1162X-, and W1282X-CFTR cells with plasmids encoding ACE-tRNAs and mNeonGreen (mNG), expression of fluorescent protein was determined by FACS to sort ACE-tRNA-transfected cells (mNG positive) from non-transfected cells. (legend continued on next page)
channels, we pre-incubated cells in forskolin (20μM) and IBMX (100μM) for 5 min before starting the recording. We propose that measuring CFTR function under these conditions, where likely saturating amounts of ACE-tRNA expression cassettes are delivered, provides a theoretical upper limit to ACE-tRNA-dependent PTC suppression efficiency of endogenous CFTR. Here, ACE-tRNA<sup>Gly</sup> and ACE-tRNA<sup>Arg</sup> mNG-positive R1162X- (Figure 5D, red trace; 19.9 ± 6.8 pA/pF at +80 mV; n = 6) and W1282X-CFTR (Figure 5D, blue trace; 26.7 ± 4.3 pA/pF at +80 mV; n = 11) 16HBE<sub>14o</sub> cells exhibited CFTR<sub>ina</sub>-172-sensitive chloride current magnitudes that were 75% and 91% of those measured in WT 16HBE14o- cells (Figure 5D, black trace; 29.3 ± 6.2 pA/pF at +80 mV; n = 7), respectively. As expected, R1162X- (Figure 5D, dashed red trace; n = 8) and W1282X-CFTR (Figure 5D, dashed blue trace; n = 5) 16HBE<sub>14o</sub> cells receiving empty mNG plasmids exhibited no significant CFTR function. The average CFTR<sub>ina</sub>-172-sensitive chloride current magnitudes are also shown in Figure 5E with individual experiments plotted to not obfuscate the considerable cell-to-cell variability of CFTR channel expression in all conditions, including WT 16HBE<sub>14o</sub> cells. These results are consistent with Ussing chamber measurements of R1162X PB 16 × ACE-tRNA<sup>Gly</sup> cells (Figure 4B), where CFTR channel function appears to track CFTR mRNA expression.

Western blot analysis indicates expression of full-length and fully glycosylated CFTR from endogenous nonsense mutations with ACE-tRNAs

To provide further evidence of successful suppression of endogenous CFTR PTCs and expression of full-length CFTR protein following transfection of ACE-tRNAs, we performed western blot analysis of R1162X- and W1282X-CFTR 16HBE<sub>14o</sub> cells transfected with 4× ACE-tRNA<sup>Gly</sup> mNG and 4× ACE-tRNA<sup>Arg</sup> mNG plasmids, respectively. Western blot analysis indicates robust readthrough of CFTR PTCs occurs only in the presence of ACE-tRNAs with rescue of full-length, successfully trafficked, CFTR, as evidenced by the presence of the fully glycosylated band C form and the core glycosylated band B CFTR (Figure 5E). Glycosylated CFTR has been shown to bind strongly to the lectin wheat germ agglutinin (WGA). To that end, we subjected solubilized membrane samples from R1162X- and W1282X-CFTR 16HBE<sub>14o</sub> cells transfected with 4× ACE-tRNA<sup>Gly</sup> mNG and 4× ACE-tRNA<sup>Arg</sup> mNG plasmids to WGA precipitation. Subsequent western blot analysis of proteins eluted from WGA agarose again indicates robust expression of full-length, PTC-containing CFTR only following transfection of ACE-tRNAs (Figure 5E). Quantification of normalized C band intensity for ACE-tRNA<sup>Gly</sup>,W1282X (45% ± 9% of WT; n = 3) and ACE-tRNA<sup>Arg</sup>,R1162X (29% ± 2% of WT; n = 3) revealed robust rescue of full-length trafficked CFTR as compared with the untransfected controls W1282X- (2% ± 1% of WT; n = 3) and R1162X-CFTR (6% ± 2% of WT; n = 3). When considering that the cells collected for western blot analysis are unbiased for transfection efficiency (mNG-positive and negative cells were not sorted) and given a transfection efficiency of ~40% (Figure 5B), these results are in close agreement with the amount of CFTR rescue demonstrated by whole-cell, patch-clamp results.

DISCUSSION

Several studies have used nonsense-suppressor tRNAs to address PTCs in certain disease-related genes, including β-globin in <i>Xenopus laevis</i> oocytes, xeroderma pigmentosum group A in human cells, chloramphenicol acetyltransferase in transgenic mice, CDH1 in human cell lines, and CFTR in human cells and <i>X. laevis</i> oocytes.30,31 We previously demonstrated using mass spectrometry of the full-length protein product of PTC readthrough that ACE-tRNAs in their current form insert the correct amino acid, providing seamless rescue of the PTC.30 While these studies indicated that ACE-tRNAs are competent to promote suppression of PTCs in mRNA produced from cDNA, they do not fully recapitulate the hurdles seen in the rescue of genuine PTCs in their genomic context. NMD is a eukaryotic translation-dependent mRNA quality control pathway that selectively degrades mRNA transcripts containing PTCs. Here, we demonstrate CFTR mRNA rescue from 16HBE<sub>14o</sub> cells containing PTCs in the endogenous genomic context. Electrophysiological...
assays, including Ussing chamber short-circuit current recording and patch clamp, are considered the “gold standard” for evaluating CFTR functional rescue in drug development.32 Using these assays, we demonstrate protein rescue of PTC-containing CFTR, providing proof of principle that ACE-tRNAs when delivered as cDNA or RNA can rescue native PTCs. Based on our work here, we note that protein rescue generally tracks with mRNA rescue from NMD. Here, we have demonstrated a rescue of 75%–90% of WT CFTR channel function at two PTC positions when selecting for cells with efficient delivery of ACE-tRNAs. Further, the western blot results demonstrate significant ACE-tRNA-dependent rescue of CFTR protein, to levels consistent with patch-clamp results. Assuming an efficient delivery modality is forthcoming, this indicates the upper limit to the therapeutic window for ACE-tRNAs is well above what is necessary for a therapeutically relevant treatment of nonsense-associated CF. While our results are promising thus far, a larger sampling of CF-causing PTCs need to be examined to determine whether all PTC-ACE-tRNA pairings will provide therapeutically meaningful rescue of CFTR channel function.

While this study represents PTC rescue in one of the most faithful PTC models used to date when assaying for function of ACE-tRNAs, a fundamental limitation is that this is an immortalized cell line and does not fully recapitulate the delivery issues or potential safety issues realized in vivo. A notable concern about the safety of ACE-tRNA is that nonsense-suppressor tRNAs may cause read-through of NTCs to toxic effect. In our previous work, we determined via ribosome profiling that readthrough of NTCs is minimal in comparison to PTCs.30 Further, in the context of genetic code expansion, stable expression and viral delivery of nonsense-suppressor tRNAs have been shown to be tolerated in mice.47,48 Our ability to generate cell lines stably expressing 16 copies of ACE-tRNAs with similar growth rates and morphology to WT parents cells lends further credibility to tolerably low levels of toxicity due to NTC readthrough. It should also be noted that ACE-tRNAs inherently possess a lower level of global NTC readthrough. Unlike the majority of small-molecule PTC readthrough agents developed to date, ACE-tRNAs only suppress one of the three stop codons. A further advantage of ACE-tRNAs over small-molecule PTC readthrough agents highlighted here is that ACE-tRNAs provide a la carte rescue; that is, the amino acid encoded at the PTC can be specified (i.e., Leu at W1282X). While results shown by ourselves and others indicate the safety of ACE-tRNAs, the DNA-delivery methods used here require the use of a transfection marker to control for efficient DNA delivery and cannot be directly translated to the clinic. This highlights a remaining roadblock of delivery to the clinical implementation of ACE-tRNAs; however, it should be noted that tRNAs as gene therapy cargo possess a number of attractive qualities.31 We have shown here that ACE-tRNAs can be delivered as RNA or DNA, for which there are many therapeutically relevant delivery avenues to pursue.

As demonstrated here, ACE-tRNAs delivered as RNA are competent to rescue the expression of endogenous PTC-containing R1162X-CFTR mRNA. Given that CFTR protein rescue generally tracks with mRNA rescue, it follows that ACE-tRNA delivered as RNA will rescue protein from genes containing endogenous PTCs. However, due to technical limitations of the model cell line used here, we have not demonstrated CFTR protein rescue directly using ACE-tRNA as RNA. While ACE-tRNA delivered as RNA can go through multiple rounds of aminoacylation and translation, it is not a renewable PTC-suppression resource like ACE-tRNA delivered as DNA. Therefore, it is likely that repeated delivery of ACE-tRNA would be needed to sustain a therapeutic level of target protein expression and that modifications to the RNA may be necessary to extend the half-life of the ACE-tRNA therapeutic agent. However, because ACE-tRNAs are only ~76 nt in length, many of the existing mRNA and short RNA delivery technologies can likely be adapted for delivery of ACE-tRNAs. These delivery methods include but are not limited to cationic or ionizable lipids and lipid-like materials and polymers and nanoparticles.49 In contrast to ACE-tRNA delivered as RNA, delivery as DNA likely will result in augmented persistence of activity.

When ACE-tRNAs are efficiently delivered as DNA, we demonstrate here rescue of both PTC-containing CFTR mRNA and near 100% rescue of CFTR protein, far exceeding the therapeutic threshold for functional CFTR rescue of 15%–30%.34 Due to the small size of ACE-tRNA expression cassette (~250 nt), there are a number of avenues available to deliver ACE-tRNA DNA expression cassettes. The small size of the ACE-tRNA expression cassette allows for optimization of cassette copy number with a minimal vector size, which in turn has been shown to increase intracellular mobility, nuclear entry, and gene therapy agent expression.30–32 Naked DNA delivery is an attractive gene therapy approach as it has inherently low immunogenicity, can be designed to remain episomal or integrate, and can be paired with several technologies, including nanoparticle formulations, liposomes, pressure injection, and electroporation.33–38 When plasmid DNA is made resistant to silencing, several studies have shown extended expression (months) in murine lung following plasmid DNA delivery by polyethyleneimine/DNA complexes,39 cationic liposome/DNA complexes,57 and DNA electroporation.58

The small expression cassette size of ACE-tRNAs also makes them amenable to delivery by a variety of viral vectors. Viral delivery is an attractive gene therapy approach, as viral cargo delivery is usually much more efficient than non-viral delivery and has the advantage of cell-type-specific transduction. There are a number of candidate viral vectors, some of which include adenovirus, AAV, and lentivirus.39 Recently, the US Food and Drug Administration (FDA) approved the use of the first AAV therapeutic for total gene-replacement therapy of RPE65 (retinal pigment epithelial-specific 65 kDa protein) for treatment of the inherited retinal disorder Leber congenital amaurosis type-2 (Luxturna). This demonstrates the possibility of a therapeutic route to viral delivery of ACE-tRNAs. Of note, ACE-tRNAs are particularly amenable to AAVs, as their small size allows for multiple expression cassettes despite the relatively small cargo capacity of AAVs (~4.7 kb). Indeed, for many diseases associated with nonsense mutations, AAV total gene replacement is not viable due to the large
size of the gene in question. In contrast, ACE-tRNA-mediated PTC suppression is independent of target gene size, meaning ACE-tRNAs delivered as an AAV can rescue gene targets of any size. As we demonstrated here that integrating up to 16 copies of ACE-tRNA<sup>AV</sup> was well tolerated in 16HBE<sub>140</sub> cells, the use of integrating lentiviral vectors for one-time delivery of ACE-tRNAs is another attractive approach.

In summary, nonsense mutations cause ~11% of all serious genetic diseases, including CF, for which there are currently few therapeutic options, even given their unitary mechanism. Our results suggest that nonsense suppressor ACE-tRNAs are capable of rescuing both PTC-containing mRNA from NMD and functional protein expression. Coupled with an efficient RNA or DNA delivery technology, ACE-tRNAs may provide relief for a variety of genetic diseases in the future.

**MATERIALS AND METHODS**

**Materials for molecular biology**

Unless otherwise specified, all molecular biology reagents were obtained from New England Biolabs. For all PCR reactions, Q5 High-Fidelity 2X Master Mix (NEB M0492L) was used, Gibson assembly was performed using NEBuilder HiFi DNA Assembly Master Mix (NEB E2621L), Golden Gate reactions used T4 DNA ligase (NEB M0202L) and restriction enzymes from NEB, gel purifications were performed using the Monarch DNA Gel Extraction Kit (NEB T1020L), and minipreps were performed using the Monarch Plasmid Miniprep Kit (NEB T1010L). NEB 5-alpha competent E. coli (NEB C2987H) were used for all bacterial transformations unless otherwise noted. All oligonucleotides were synthesized by Integrated DNA Technologies. All cDNA preparations were performed using the NucleoBond Xtra Midi EF kit (Macherey Nagel Ref. 740420.50).

**DNA constructs and plasmid generation**

The PB transposon plasmid with PB inverted terminal repeats (ITRs) flanking an ampicillin resistance marker, pBR322 origin; short ubiquitin C promoter (Ubc)-puromycin N-acetyltransferase-SV40 pA; and EcoRV, PmlI, PmeI, and SmaI multiple cloning site in pUC57 was synthesized (Genscript). The hyperPB transposase sequence<sup>AV</sup> was synthesized as gBlocks (IDT, Coralville, IA, USA) and Gibson Assembled into pcDNA3.1(+). For further specifics of plasmid cloning methods, see supplemental materials and methods.

**tRNA RNA synthesis**

ACE-tRNA<sup>AV</sup> used as RNA was synthesized by IDT. The sequence synthesized was ArgTGAct9tRNA6/no intron (Arg-TCT → TGA-3:1). 5′-GGGCUUGUGUGGCGCAUUGAUGCGCC UUGGACU/C/AAUUGGUAAGGUUGGGUGCUAGUCCAC CAGAGUCC-3′.

The ACE-tRNA was not pre-treated or pre-folded and was suspended in diethyl pyrocarbonate (DEPC)-treated ddH<sub>2</sub>O at 1 mg/mL.

**Cell culture**

Parental 16HBE140- human bronchial epithelial cells, which were generated by Dieter Gruenert (University of California, San Francisco),<sup>AV</sup> and 16HBE<sub>140</sub> cells with engineered CFTR variants, CFF-16HBE<sub>140</sub> CFTR G542X, CFF-16HBE<sub>140</sub> CFTR R1162X, and CFF-16HBE<sub>140</sub> CFTR W1282X cells, were obtained from the Cystic Fibrosis Foundation Therapeutics Lab (Lexington, MA, USA).<sup>AV</sup> HBE cells were cultured in minimum essential medium (MEM) (Gibco; no. 11095-080) supplemented with 10% fetal bovine serum (Gibco; no. 26140-079) and 1% penicillin/streptomycin/glutamine (Gibco; no. 10378-016) in a 37°C/5% CO<sub>2</sub> humidified incubator. Plates were coated with coating solution (LHC-8 basal medium [Gibco; no. 12677-027], 1.34 μL/mL BSA 7.5% [Gibco; no. 15260-037], 10 μL/mL bovine collagen solution, type 1 [Advanced BioMatrix; no. 5005–100ML] and 10 μL/mL fibronectin from human plasma (Thermo Fisher Scientific; no. 33016-015]) for 2 to 3 h at 37°C/5%/ CO<sub>2</sub> and stored at 4°C after thorough removal of coating solution.

**Transfection and nucleofection of HBE cells**

HBE cells were transfected with Lipofectamine LTX and Plus reagent (Invitrogen by Thermo Fisher Scientific; no. 15338-100) according to manufacturer’s guidelines with slight modifications. Briefly, on the day of transfection, 500,000 cells per well were plated in a 6-well plate and incubated in a cell-culture incubator for 20–30 min. 2.5 μg of DNA and 2.5 μL of PLUS reagent were diluted in 125 μL of Opti-MEM (Gibco; no. 31985-034), and 5 μL of Lipofectamine LTX reagent was diluted in Opti-MEM. Diluted CDNA and Lipofectamine LTX reagent were combined, mixed thoroughly, and incubated for 5 min at room temperature. The transfection mixture was then added dropwise to pre-plated cells. The media containing transfection mix was replaced with normal cell culture media 16–24 h post-transfection. HBE cells were nucleofected with the Amaxa 4D-Nucleofector system (Lonza, Basel, Switzerland) using the Amaxa SG cell line 4D-Nucleofector X Kit S (Lonza; no. V4XC-3032). Each well of a black 96-well plate was pre-warmed in a 37°C/5% CO<sub>2</sub> incubator until an aliquot of cell culture media was pre-warmed in a 37°C/5%/ CO<sub>2</sub> water bath. For each reaction, 200,000 cells were centrifuged at 90 × g for 10 min, resuspended in 20 μL of supplemented SG 4D-Nucleofector X solution, and mixed with 1 μg of RNA (maximum 10% of final sample volume). A total volume of 20 μL was transferred to a well of Nucleocuvette Strip. The nucleofection process was started by placing the strip into the 4D-Nucleofector X Unit and running the CM-137 Nucleofector program from the 4D-Nucleofector Core Unit. Cells were incubated at room temperature for 10 min and resuspended with 80 μL of pre-warmed cell culture media. Fifteen microliters of nucleofected cells were plated in pre-incubated cell-culture plate and maintained in a humidified 37°C/5% CO<sub>2</sub> incubator until analysis.

**Stable cell line generation**

To generate stable HBE cell lines, the PB transposon vector expressing gene of interest (NLuc PTC reporter or ACE-tRNA) was co-transfected...
with the PB transposase vector. For each well of a 12-well plate, 250,000 cells were seeded on the day of transfection and transfected with 0.5 μg of PB transposon vector and 0.5 μg of empty vector as a negative control or 0.5 μg of PB transposon vector and 0.5 μg of PB transposase vector using Lipofectamine LTX and Plus reagent as described above. Stable cells were selected with 0.5 μg/mL of puromycin (InvivoGen; no. ant-pr-1) 1 to 2 days post-transfection until no viable cells were detected in the negative-control condition.

**FACS**

For FACS experiment, cells were washed with Dulbecco’s phosphate-buffered saline (DPBS) (Gibco; no. 14190-144) twice, dissociated with TrypLE Express (Gibco; no. 12604-013), pelleted at 1,000 rpm (~500 × g) for 5 min, resuspended with DPBS with calcium and magnesium (Gibco; no. 14040-133), and strained with 40 μm Cell Strainer. Cells were then sorted using the BD FACSaria II cell sorter (BD Biosciences), and data were analyzed by using FCS Express 7 flow cytometry software (De Novo Software, Pasadena, CA, USA).

In order to sort ACE-tRNA-expressing cells from ACE-tRNA-not-expressing cells, HBE cells were transfected with the plasmid-containing ACE-tRNA and mNG fluorescent protein in one plasmid. Approximately 36–48 h after transfection, cells were prepared as described above for FACS experiment. Empty-vector-expressing cells were used to determine the gate for mNG-negative and positive cells. Each population of mNG-negative and positive cells was then sorted into a 15-mL conical tube using the BD FACSaria II cell sorter. Total RNA was isolated from sorted cells and used for reverse transcriptase and quantitative PCR (RT-qPCR) as described below.

**Luciferase reporter assay**

PTC readthrough by ACE-tRNA was assessed by measuring NLuc luciferase activity using the Nano-Glo Luciferase Assay System (Promega; no. N1130) on a Synergy2 multi-mode microplate reader (BioTek Instruments, Winooski, VT, USA). Using Lipofectamine LT and Plus reagent as described above, HBE cells were co-transfected with NLuc, NLuc-PEST, or Sec-NLuc PTC reporters and ACE-tRNA, or HBE cells stably expressing PTC reporters were transfected with ACE-tRNA. In each well of a black 96-well plate, 15 μL of DPBS was added for cells expressing NLuc and NLuc-PEST PTC reporters, whereas 15 μL of media from transfected cells were transferred for cells expressing Sec-NLuc PTC reporter. Then, 15 μL of reconstituted Nano-Glo Luciferase Assay Reagent (1:50 of Nano-Glo Luciferase Assay Substrate to Nano-Glo Luciferase Assay Buffer) was added to each well for luminescence recording using a Synergy2 multi-mode microplate reader. Each condition was assayed in triplicate, and cells expressing empty vector instead of ACE-tRNA were included as a negative control. Luciferase activity was displayed as raw NLuc luminescence or normalized NLuc luminescence to background luminescence.

**qRT-PCR**

Total RNA was isolated from cells with the Monarch Total RNA Miniprep Kit (NEB; no. T2010S) according to manufacturer’s recommen-


dations. RNA quantity and quality were measured with a NanoDrop OneC Spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA). Subsequently, one-step RT-qPCR was performed on the QuantStudio 3 Real-Time PCR System (Applied Biosystems, Waltham, MA, USA) using the Luna Universal One-Step RT-qPCR Kit (NEB; no. E3005L or no. E3006E) according to manufacturer’s instructions and then analyzed with QuantStudio Design & Analysis Software v.1.5.1 (Applied Biosystems). Briefly, the reaction mixture contained 10 μL 2× Luna Universal Probe One-Step Reaction Mix, 1 μL 20× Luna WarmStart RT Enzyme Mix, 1 μL 20× CFTR-specific primers and probe, 1 μL TBP-specific primers and probe, 100–500 ng of template RNA, and nuclease-free water in a final volume of 20 μL. The amplification conditions were as follows: one cycle of 10 min at 55°C and one cycle of 3 min at 95°C followed by 40 cycles of 15 s at 95°C and 30 s at 60°C. Fluorescence was detected at the end of the 60°C extension step. Each sample was quantified in triplicate, and a no-template control (NTC) reaction, with just nuclease-free water, was included as a negative control. The fold difference in CFTR gene expression normalized to TBP was calculated using the comparative Ct method, 2−ΔΔCt. The following target-specific primers and probe used in RT-qPCR experiments were purchased from IDT: CFTR probe, CFTR primer 1, and CFTR primer 2 and endogenous control TBP-specific TBP probe, TBP primer 1, and TBP primer 2 (sequences in Table S2). The final concentration of primers was 500 nM each and probe was 250 nM.

**Ussing chamber studies of HBE cells**

HBE cells were seeded onto 6.5 mm Transwell Permeable Support with 0.4 μm Polycarbonate Membrane Inserts (Corning; no. 3413) pre-coated with collagen from rat tail (Sigma-Aldrich; no. C7661-25MG) 5 to 6 days before assay and maintained in normal cell culture media at 37°C and 5% CO2. Cells were grown until the formation of electrically tight epithelial monolayer was confirmed by measuring transepithelial resistance with EVOM2 Epithelial Voltohmmeter (World Precision Instruments, Sarasota, FL, USA). Short-circuit current (Isc) was measured under voltage-clamp condition to determine CFTR channel function. Cell monolayers were mounted in Ussing chambers (Physiologic Instruments, San Diego, CA, USA) filled with Ringer’s solution (in mM: 135 NaCl, 5 HEPES, 0.6 KH2PO4, 2.4 K2HPO4, 1.2 MgCl2, 1.2 CaCl2, and 5 D-glucose adjusted to pH 7.4) at 37°C and gassed with compressed air. After baseline record-
ings, cells were treated from the apical side with 100 μM amiloride to block epithelial sodium channels and then with 100 μM 4,4’-disothiocyanostilbene-2,2’-disulfonic acid (DIDS) to block Ca2+-activated chloride channels. Then, the solution in the apical side was replaced with a low Cl− solution (in mM: 135 Na-glucconate, 5 HEPES, 0.6 KH2PO4, 2.4 K2HPO4, 1.2 MgCl2, 1.2 CaCl2, and 5 D-glucose adjusted to pH 7.4). CFTR-mediated Cl− current was measured by adding CFTR activators, 10 μM forskolin and 100 μM 3-isobutyl-1-methyuxanthine (IBMX), and CFTR inhibitor, 20 μM CFTR inhibitor-172 (CFTRinh-172), to the apical side. Data were acquired with Acqure & Analyze 2.3 software (Physiologic Instruments). The maximum change in Isc in response to additions of CFTR activators...
and CFTR inhibitor were quantified as measures of functional CFTR Cl\textsuperscript− transport activity.

**Macroscopic recordings of CFTR currents**

Cells were transfected using Lipofectamine LTX Reagent with PLUS Reagent (Invitrogen) with a modified reverse transfection as described above. W1282X-CFTR 16HBE14o- cells were transfected with either a plasmid containing four copies of ACE-tRNA\textsuperscript{A\textsuperscript{rg}} with mNG or empty plasmid encoding only mNG. Results were compared with WT 16HBE14o- cells transfected with empty mNG plasmid. After transfection, cells were lifted using Versene (Gibco; no. 15040-066) and plated on 35-mm dishes coated with human fibronectin (as previously described) at low density, and cells in isolation were patched 2 to 3 days post-transfection. Cells were continuously perfused with external solution (in mM: 145 TEA-Cl, 10 CaCl\textsubscript{2}, and 10 HEPES adjusted to pH 7.4 with TEA-OH) supplemented with 10 \mu M forskolin (Sigma) and 100 \mu M IBMX (Sigma) for maximal activation of CFTRs; dishes of cells were incubated in this solution for at least 5 min prior to recording. Green cells were patched using thick-walled borosilicate glass pipettes (Sutter, Novato, CA, USA) pulled to approximately 7 MΩm using a vertical puller (PC-100, Narishige, Amityville, NY, USA). Pipettes were filled with internal solution (in mM: 30 CsCl, 110 Cs-aspartate, 2.5 CaCl\textsubscript{2}, 2.5 MgCl\textsubscript{2}, 10 HEPES, 5 CsEGTA, and 3 mM MgATP adjusted to pH 7.4 with CsOH). Internal and external solutions were formulated to eliminate non-chloride contaminating currents (i.e., TEA and cesium to block potassium currents and no sodium present to eliminate sodium currents) and feature a reversal potential for chloride of approximately −36 mV. Currents were amplified using an Axon 200B amplifier (Axon Instruments, Hamden, CT, USA), sampled at 10 kHz and filtered at 2 kHz using a Digidata 1440A, and recorded using pClamp 10 (Molecular Devices, San Jose, CA, USA). Whole-cell capacitance and series resistance were compensated (approximately 70%), and a linear ramp voltage protocol from −80 mV to +80 mV (200 ms) was then applied from the holding potential (−40 mV) to collect current traces at 0.5 Hz for 60 sweeps. Per fused solutions were then rapidly exchanged using a fast-step perfusion barrel (VC-8, Warner Instruments) to an external solution containing 20 \mu M CFTR\textsubscript{inh}-2 to block CFTR currents, and the ramp voltage protocol was continued at 0.5 Hz for 180 more sweeps. Recordings were analyzed using Clampfit 10, whereby peak outward current near +80 mV was divided by cell capacitance, as determined by the membrane test function in pClamp, to yield current densities (pA/pF). Peak outward current density at +80 mV was then plotted versus time using Prism 9 (GraphPad) for comparison of treatment groups and controls. Only experiments where a gigaohm seal was achieved before break-in and had a measured reversal potential that fell within 7 mV of −36 mV were used in analysis.

**Quantification of stably integrated copy number using quantitative real-time PCR**

Genomic ACE-tRNA\textsuperscript{A\textsuperscript{rg}} integration of the cell population was quantified by performing quantitative real-time PCR. Genomic DNA (gDNA) was extracted using the Monarch Genomic DNA Purification Kit (NEB; no. T3010S) according to manufacturer’s instructions. Genomic integrants of the PB transposon expressing 16 copies of ACE-tRNA\textsuperscript{A\textsuperscript{rg}} cassette (PB 16× ACE-tRNA\textsuperscript{A\textsuperscript{rg}}) were detected by targeting either the puromycin cassette (puro) or PB backbone from the PB transposon vector. Puro-specific puro probe, puro primer 1, and puro primer 2 and PB-backbone-specific PB backbone probe, PB backbone primer 1, and PB backbone primer 2 were purchased from IDT. The gDNA content was normalized based on the amplification of the Ribonuclease P (RNase P) gene using RNase P probe purchased from Applied Biosystems and RNase P primer 1 and RNase P primer 2 purchased from Invitrogen. The final concentration of primers was 500 nM each and probe was 250 nM. Real-time PCR was performed on the QuantStudio 3 Real-Time PCR System using the PrimeTime Gene Expression Master Mix (IDT; no. 1055772) according to manufacturer’s guidelines and then analyzed with QuantStudio Design & Analysis Software v.1.5.1. Briefly, the reaction mixture contained 10 \mu L 2× PrimeTime Gene Expression Master Mix, 1 \mu L 20× puro- or PB backbone-specific primers and probe (target gene), 1 \mu L RNase P-specific primers and probe (reference gene), DNA template, and nuclease-free water in a final volume of 20 \mu L. PCR cycling conditions were one cycle of 3 min at 95°C followed by 40 cycles of 15 s denaturation at 95°C and 1 min annealing and extension at 60°C. A standard curve was constructed by serial dilutions of a PB 16× ACE-tRNA\textsuperscript{A\textsuperscript{rg}} plasmid with puro- or PB backbone-specific primers and probe and Hs_RPP30 Positive Control (IDT; no. 10006622) with RNase-P-specific primers and probe. The number of ACE-tRNA\textsuperscript{A\textsuperscript{rg}} copies in 10 ng or 1 ng DNA from cells stably integrated with ACE-tRNA\textsuperscript{A\textsuperscript{rg}} was calculated by plotting cycle threshold (C\textsubscript{T}) to the linear equation from the standard curve. All qPCR reactions were run in triplicate. An NTC reaction with just nuclease-free water was included as a negative control, and 10 ng or 1 ng DNA from cells not integrated with ACE-tRNA\textsuperscript{A\textsuperscript{rg}} was assayed to confirm specificity of puro- and PB-backbone-specific primers and probe.

**Western blot analysis**

16HBE14o- cells were grown and transfected as described above. Following 36 h, the cells were scraped in ice-cold PBS supplemented with protease inhibitor cocktail (EDTA-Free; MedChemExpress, Monmouth Junction, NJ) and pelleted at 7,000 × g for 8 min, the supernatant aspirated, and pellets stored at −80°C. The cell pellet was vigorously dounced in membrane separation buffer (100 mM sucrose, 150 mM NaCl, 1 mM dithiothreitol [DTT], 1 mM EDTA, protease inhibitor cocktail, and 50 mM Tris-HCl [pH 7.4]) and centrifuged at 100,000 × g to separate total membranes from the soluble cytosolic proteins. Pellets were solubilized in buffer (1% Triton X-100, 250 mM NaCl, protease inhibitor cocktail, and 50 mM Tris-HCl [pH 7.4]) and rocked for 1 h at 4°C, and insoluble cell debris was pelleted at 14,000 rpm. Protein concentration in clarified cell lysate was determined by BCA assay (Pierce BCA Protein Assay Kit, Thermo Scientific, Waltham, MA) using a Synergy H4 Hybrid Microplate Reader (BioTek). For wheat germ agglutinin precipitation, 120 \mu g of Triton X-100 solubilized membrane protein was added to 25 \mu L of WGA agarose resin (Vector Laboratories,
Burlingame, CA) in 1 mL of WGA wash buffer (10 mM HEPES [pH 7.5], 150 mM NaCl, and protease inhibitor cocktail) and incubated rocking overnight at 4°C. The resin was washed 3× in 1 mL ice-cold WGA wash buffer, and the WGA-bound protein was eluted in 50 μL of 5× Laemml sample buffer (LSB) (15% SDS, 575 mM sucrose, 325 mM Tris-HCl [pH 6.5], 5% β-mercaptoethanol, and 0.002% bromophenol blue) with a 30-min incubation at 37°C. Equal volume of LSB eluate from the WGA resin was used for western blot analysis. Equal total protein amounts were loaded on a 3% stacking gel and 10% resolving gradient SDS-PAGE with 4% stacking gel in the presence of 0.45% β-mercaptoethanol, separated at 120 V, and transferred to 0.45 μm Immobilon-FL PVDF (Merck Millipore, Tullagreen, Carrigtwohill, Co. Cork, Ireland). PVDF was immunoblotted using anti-CFTR antibody (1:1,000, University of North Carolina CFTR Antibody, 1:2,000, sc-48345, Santa Cruz Biotechnology, Dallas, TX) in Intercept Protein-Free Blocking Buffer (LI-COR Biosciences, Lincoln, NE) and imaged on an LI-COR Odyssey Imaging System. Quantiﬁcation of bands was conducted using ImageJ (v.1.52q). CFTR C band intensity was normalized to Na+/K+-ATPase actin band intensity for each blot. The normalized CFTR C band intensity for each of W1282X- and R1162X-CFTR was then compared with the normalized WT CFTR C band control on the blot to give the percent rescue as compared with WT.

Statistical analysis

Data are presented as mean ± standard error of the mean (SEM). GraphPad Prism 7 software (GraphPad, San Diego, CA, USA) was used for all statistical analysis and plotting except for Figure 5D, which was plotted using GraphPad 9. Comparisons between two groups were performed using an unpaired Student’s t test. Multiple comparisons were analyzed using a one-way analysis of variance (ANOVA) followed by Tukey’s multiple comparisons test. p < 0.05 was considered statistically significant.

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.omtn.2022.04.033.

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AUTHOR CONTRIBUTIONS

J.J.P., W.K., and J.D.L. designed the study, J.J.P., W.K., M.T.S., K.M.E., and J.D.L. performed experiments. J.J.P., W.K., M.T.S., and J.D.L. analyzed the data and constructed the ﬁgures. J.J.P., W.K., and J.D.L. wrote the manuscript. All authors read and revised the manuscript.

DECLARATION OF INTERESTS

J.D.L. is a co-inventor of the technology presented in this study and receives royalty payments related to the licensing of the technology from the University of Iowa. PCT/US2018/059065, ﬁled November 2, 2018; METHODS OF RESCUEING STOP CODONS VIA GENETIC REASSIGNMENT WITH ACE-rRNA; Inventors - University of Iowa – Inventors J.D.L. and Christopher A. Ahern pertains to the tRNA sequences used in this study.

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