Barriers to inhaled gene therapy of obstructive lung diseases: A review

Namho Kim\textsuperscript{a,b,1}, Gregg A. Duncan\textsuperscript{a,c,1}, Justin Hanes\textsuperscript{a,b,c,d}, and Jung Soo Suk\textsuperscript{a,c,*}

\textsuperscript{a}The Center for Nanomedicine, Johns Hopkins University School of Medicine, Baltimore, MD 21231, USA

\textsuperscript{b}Department of Chemical and Biomolecular Engineering, Johns Hopkins University, Baltimore, MD 21218, USA

\textsuperscript{c}Department of Ophthalmology, Johns Hopkins University School of Medicine, Baltimore, MD 21231, USA

\textsuperscript{d}Departments of Biomedical Engineering, Environmental and Health Sciences, Oncology, Neurosurgery, and Pharmacology and Molecular Sciences, Johns Hopkins University, Baltimore, MD 21205

Abstract

Knowledge of genetic origins of obstructive lung diseases has made inhaled gene therapy an attractive alternative to the current standards of care that are limited to managing disease symptoms. Initial lung gene therapy clinical trials occurred in the early 1990s following the discovery of the genetic defect responsible for cystic fibrosis (CF), a monogenic disorder. However, despite over two decades of intensive effort, gene therapy has yet to help patients with CF or any other obstructive lung disease. The slow progress is due in part to poor understanding of the biological barriers to inhaled gene therapy. Encouragingly, clinical trials have shown that inhaled gene therapy with various viral vectors and non-viral gene vectors is well tolerated by patients, and continued research has provided valuable lessons and resources that may lead to future success of this therapeutic strategy. In this review, we first introduce representative obstructive lung diseases and examine limitations of currently available therapeutic options. We then review key components for successful execution of inhaled gene therapy, including gene delivery systems, primary physiological barriers and strategies to overcome them, and advances in preclinical disease models with which the most promising systems may be identified for human clinical trials.

Graphical abstract

\textsuperscript{*}Corresponding author: Jung Soo Suk, Ph.D., Center for Nanomedicine at the Wilmer Eye Institute, Johns Hopkins University School of Medicine, 400 N. Broadway, Robert H. and Clarice Smith Building, 6029, Baltimore, MD 21231, USA, Telephone: (410) 614-4526, Fax: (443) 287-7922, jsuk@jhmi.edu.

\textsuperscript{1}These authors equally contributed to this work.

Publisher’s Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.
Keywords
Respiratory gene therapy; gene delivery vector; biological barrier; preclinical model

1. Introduction
Obstructive lung diseases consist of a group of respiratory disorders characterized by airway obstruction in the lungs of affected patients. All included diseases entail severe respiratory morbidity that often results in pulmonary failure and disease-associated mortality. Current treatments generally improve patient quality of life. However, these options do not tackle the root of the disease [1], and patients are required to follow cumbersome therapeutic regimens throughout their lifetime [2]. Gene therapy has emerged as a promising alternative approach as a result of the growing number of identified genetic causes and modifiers of obstructive lung diseases. With advances in nucleic acid engineering, it is now conceivable to achieve gene transfer to specific cell types, including lung parenchymal cells [3–5], without affecting non-target cells [6]. In addition, persistent transgene expression during the life span of transfected cells can be achieved [7, 8].

Therapeutic delivery via inhalation provides direct access to the target of gene therapy for obstructive lung diseases, namely the cells lining the lung airways and airspace (i.e., alveoli), in a relatively non-invasive manner. The vast majority of preclinical and clinical studies of respiratory gene therapy have employed a pulmonary delivery strategy, including intratracheal/intranasal instillation and nebulization [9–11]. Nebulizers generate inhalable micron-sized liquid droplets that can carry hundreds of gene vectors per drop and reach virtually all areas of the lung [12]. However, despite encouraging safety and tolerability results, therapeutically effective inhaled gene therapy of obstructive lung diseases has not yet been achieved. Lack of clinical success is due to the limited ability of gene delivery vectors to overcome difficult biological barriers, which stems from a relatively poor understanding of the barriers. Improvement of this understanding may allow for more rational vector design strategies to tackle them. For example, accumulation of thickened airway mucus was recognized early on as a key pathological event of obstructive lung diseases, but overcoming it’s barrier properties has typically not been addressed in the design of inhaled therapeutic delivery systems. In addition, most clinical trials involving airway gene therapy have been pursued as a result of positive outcomes observed in animal models that lack key features of human obstructive lung diseases [13–16]. The disappointments in more than 25 prior gene therapy clinical trials for CF alone have refocused research efforts on deepening knowledge.
of key issues that have prevented success of inhaled gene therapy. This review is focused on these critical issues, including clinically-relevant gene delivery vectors, important biological barriers to successful gene transfer in the lungs, and rationally-designed approaches to overcome the barriers. Finally, we review advances in the development of improved animal models to test advanced gene transfer strategies prior to their evaluation in clinic trials.

2. Obstructive lung diseases for inhaled gene therapy

2.1. Cystic fibrosis (CF)

CF is the most common inherited genetic disorder in the US, and more than 70,000 people are affected worldwide. CF is generally caused by one of many different possible mutations to a single gene that encodes cystic fibrosis transmembrane conductance regulator (CFTR). Alteration of the CFTR ion channel protein causes abnormal ion transport between lung airway epithelial cells and the airway surface liquid (ASL), as well as similar ion transport defects in the gastrointestinal and genital tracts [17]. Pulmonary complications are the primary cause of CF-related morbidity and mortality [18]. Aberrant ion regulation in CF, including impaired chloride secretion and dysregulated sodium absorption, leads to dehydration of the ASL. The result of dehydration is a thickened mucus gel in the airways that can impair mucociliary clearance (MCC) [19]. Altered biophysical properties of airway mucus, which impair mucus detachment, may also play a role in impaired MCC in CF [20]. Reduced rates of MCC cause mucus accumulation in the airways and provide a permissive environment for chronic bacterial infection and associated inflammation, which together can cause airway obstruction, fibrosis and, eventually, pulmonary failure [19, 21]. Airway dehydration, infection and inflammation are more pronounced in females with CF, which may be due to airway-related modulatory effects of estrogen [22]. As a result, females with CF have higher mortality compared to males with CF [22].

Common inhalable treatments for CF, including recombinant human DNase (Pulmozyme®), hypertonic saline and antibiotics (TOBI®), alleviate disease symptoms, but do not address the underlying root of the problem, CFTR dysfunction. In 2012, the FDA approved the first drug that directly addresses CFTR dysfunction, Ivacaftor (VX-770; Kalydeco®) [23]. Ivacaftor demonstrated significant improvements in lung function of CF patients [24, 25]. However, only patients with a specific class of mutations respond to Ivacaftor treatment, representing only a small subset of the patient population (~5%). Another drug, lumacaftor (VX-809), showed promising in vitro correction of the most prevalent CFTR mutation, ΔF508, which represents ~70% of CF patients [26]. This finding prompted its clinical evaluation, in combination with Ivacaftor [27], and the result was published in 2015 [28]. Although the trial demonstrated only modest improvement in pulmonary function compared to Ivacaftor in G551D patients [29], the combined formulation was recently approved by the FDA under the brand name of Orkambi®.

There are more than 1900 identified CFTR mutations, many of which are not expected to be responsive to currently available CFTR drugs [27]. Inhaled CFTR gene therapy, as a means to treat the underlying cause of the disease in the lungs, could benefit CF patients regardless of their specific CFTR mutation. However, over 25 clinical trials testing viral or non-viral gene vectors have failed to show clinical benefits, largely due to inefficient gene transfer to
target cells [9, 10], including serous cells in the submucosal glands and ciliated airway epithelial cells [30]. Some viral CF gene therapy trials have been discontinued due to the generation of host immune response that renders subsequent treatments ineffective [10, 31]. It should be noted that lifetime repeated treatment is likely required for CF, as therapeutic effects will eventually fade away due to the transient nature of episomal transgene expression [32] and/or the natural lifespan of transfected cells [7]. The UK CF Gene Therapy Consortium has recently completed the only CF gene therapy clinical trial that has been active in the past decade [33]. In this study, Alton et al. demonstrated, using a non-viral gene vector, a significant, yet modest, benefit compared to placebo control. They concluded that a more potent gene delivery vector is required to make gene therapy a viable option for treating CF [33].

Clinical trials for CF gene therapy have shown evidence of CFTR transgene expression based on measurements of CFTR mRNA and changes in nasal potential difference (NPD), but no significant improvement in lung function parameters has been reported [34, 35]. This suggests that the levels of gene transfer achieved in clinical trials have been insufficient to mediate functional cure in the CF airways. Nevertheless, optimism remains as several studies have suggested that a modest level of functional CFTR protein may be sufficient to improve lung function of CF patients. An early in vitro study suggested that only ~5% of airway epithelial cells need to produce functional CFTR proteins to restore chloride ion balance in the CF lung [36]. More recently, Pickles et al. used an in vitro model of human CF ciliated airway epithelium and found that at least 25% of cells may be required to express functional CFTR proteins in order to achieve mucus transport rates comparable to those in non-CF airways [37]. Interestingly, CF patients with certain mutations, which retain ~10% of normal CFTR expression per cell, are generally not afflicted by CF lung diseases [38]. Based on these observations, modest levels of CFTR protein expression throughout the airway epithelium could normalize pulmonary function in CF lungs.

All CF gene therapy clinical trials to date have tested delivery of wild-type CFTR genes in order to provide functional proteins. However, approaches to rescue defective CFTR have been introduced in the literature, which involve miRNA [39], peptide nucleic acid [40], zinc-finger nuclease [41] and CRISPR/Cas9 [42] technologies. These studies demonstrated in vitro and/or in vivo restoration of the ΔF508 CFTR function. Recently, the CF Modifier Consortium, which combines research efforts from groups in North America and France, completed a genome-wide association study to identify genetic loci relevant to CF pathophysiology [43]. In this study, samples from 6,365 CF patients with over 8 million genetic variants were analyzed and five genetic modifier loci associated with disease severity were discovered. This finding may provide additional genetic targets and enable individualized treatment of CF.

2.2. α-1 antitrypsin deficiency

α-1 antitrypsin deficiency (AATD) is another attractive target for gene therapy since it is also a monogenic disorder. AATD is caused by mutation in the gene encoding the serine protease inhibitor (α-1 antitrypsin; AAT). In normal conditions, AAT is synthesized predominantly in the liver, secreted directly into the bloodstream, and transported to the
lungs where it protects alveolar interstitial elastin from degradation by neutrophil elastase [44]. However, reduced AAT secretion in AATD leads to protease/anti-protease imbalance and airway inflammation in the lungs. As a result, patients with AATD develop emphysema and chronic obstructive pulmonary disease (COPD; discussed in section 2.3), which is often triggered by environmental factors such as acute infection and cigarette smoking [9]. A small subset of AATD patients (<10%) develops symptomatic liver disease. It is generally accepted that mutant AAT molecules polymerize and accumulate in the endoplasmic reticulum of hepatocytes, leading to the elevation of pro-inflammatory signaling [45].

Weekly intravenous infusion of AAT protein (i.e. augmentation therapy) is the only therapeutic option for AATD lung diseases that is currently approved by the FDA [44]. This protein-based augmentation therapy is well tolerated, effective in restoring AAT serum levels to the therapeutic threshold of 11 µM, and improves lung function [44, 46]. However, the therapy is expensive and requires frequent dosing in the clinic. Thus, a gene therapy product that allows sustained production of AAT over a longer period of time may offer a significant advantage over the current therapy.

There have been four clinical trials conducted for AATD gene therapy to date [47]. The first proof-of-concept clinical trial involved intranasal administration of non-viral gene vectors carrying the AAT gene [48]. In this study, one-third of the level after AAT protein therapy was achieved, but the effect was transient. In subsequent trials, viral gene vectors were intramuscularly dosed to AATD patients, since the site or type of cells that produce AAT is irrelevant [47]. However, in a recent Phase II clinical trial, viral gene vectors administered via this route mediated AAT production at only 3–5% of the therapeutic target, necessitating an improved gene transfer strategy [49].

Directly targeting the lungs via inhalation may provide a therapeutically-beneficial level of AAT production at the site of action. Although there is no active clinical trial evaluating inhaled AATD gene therapy, inhalable AAT protein-based augmentation therapy is in progress in AATD patients with emphysema [46]. Encouragingly, an inhaled AATD gene therapy trial demonstrated superior anti-inflammatory effects in the lungs compared to intravenous protein therapy [48]. Gene therapy approaches for AATD liver diseases include the use of short hairpin RNA (shRNA) or miRNA for the knockdown of mutant AAT in hepatocytes; detailed information is available elsewhere [44, 46, 47].

2.3. Chronic obstructive pulmonary disease

COPD is an incurable disease that is expected to be the third largest cause of death in the world by 2020 [50]. Cigarette smoking is generally accepted as the major cause of the disease, but exposure to environmental and/or work-related pollutants is also reported to be a significant factor [51, 52]. Abnormal inflammation and oxidative stress mediated by excessive inhalation of particulate matter and certain gases cause destruction of the extracellular matrix, leading to disease progression [51]. COPD is characterized by a progressive and irreversible airway limitation. This results from chronic bronchitis characterized by fibrosis, obstruction and remodeling of small airways [53]. Emphysema may also occur characterized by enlargement of airspace and destruction of lung parenchyma [53]. COPD patients with frequent exacerbations experience increased airway
inflammation, dynamic lung hyperinflation, elevated bacterial colonization in the lower airways and increased susceptibility to viral infection of the airways, rendering the lungs the major target for treatment [54].

Smoking cessation is an important part of COPD treatment, however, many COPD patients continue to suffer from the disease due to irreversible functional and anatomical alterations [55]. Current treatments for COPD reduce symptoms only, and do not arrest or reverse deterioration in lung function and architecture that accompanies moderate to severe disease. Available pharmacological treatments include bronchodilators, such as \( \beta \)-agonists and muscarinic antagonists, and inhaled corticosteroids, each of which offer short-term management of disease symptoms [56]. Other medications include mucolytic agents [57] and antibiotics [58], both recommended for patients undergoing acute exacerbations.

COPD gene therapy research has been slow due to the highly variable disease etiology and lack of good animal models, each of which limit the pace of drug development [9]. While clinically-tested AATD gene therapies may also prove useful for treatment of COPD, mutations in the AAT gene are responsible for only \(-1\text{-}3\%\) of COPD cases [51]. Several genetic loci involved in protease/anti-protease, antioxidant, or anti-inflammatory activities have been identified to exhibit polymorphisms associated with COPD [51]. However, preclinical evaluation of genetic intervention has not been extensively pursued. For example, pro-inflammatory cytokines like IL-18 and IL-1\( \beta \) have been implicated in COPD pathogenesis, but their potential role as therapeutic targets remains to be explored [59].

Recent genetic, biochemical and histological evidence suggests altered transforming growth factor beta (TGF-\( \beta \)) signaling is associated with COPD development and progression [60–62]. TGF-\( \beta \) levels are elevated in the conducting airway (i.e., bronchi and bronchioles) and airspace compartments of patients with COPD [63–65]. In addition, Podowski et al. have shown that angiotensin receptor blocker antagonizes TGF-\( \beta \) signaling and, as a result, attenuates smoke-induced lung injury and rescues lung airway and airspace architecture in a mouse model of COPD [66]. Knocking down TGF-\( \beta \) signaling in the airways and alveolar sacs via gene silencing technologies may provide similar therapeutic outcomes. Interestingly, cigarette smoking has been shown to reduce CFTR mRNA levels, CFTR protein levels and CFTR function, which contributes to mucus clearance defects in patients with COPD [67]. Thus, CFTR is another potential therapeutic target for COPD therapy.

### 2.4. Asthma

Allergic asthma is a global health problem caused by unregulated production of cytokines secreted by allergen-specific, type 2 helper T-cells (Th2 cells) [68]. Similar to COPD, both genetic predisposition and exposure to environmental irritants contribute to the development of asthma. Allergens presented to naïve T cells lead to expression of various cytokines, including IL-4, IL-5, IL-9 and IL-13, via the Th2 pathway, resulting in immunoglobulin E (IgE) production and mast cell recruitment [69]. Exposure of sensitized individuals to allergens induces release of histamine, leukotrienes and prostaglandins by mast cells in the lung airways, which promote vascular permeability, smooth muscle contraction and mucus production [69]. Subsequently, chemokines released by mast cells attract macrophages,
eosinophils, Th2 cells and basophils to the airways, triggering airway inflammation, tissue damage, and allergen hypersensitivity [69].

Anti-inflammatory medication, specifically inhaled corticosteroids, is the most frequently prescribed therapy for asthma patients [70]. β2-adrenergic receptor agonists are also frequently used to dampen the inflammatory response by relaxing the smooth muscle [71]. While effective for most of patients, corticosteroids and β2-adrenergic receptor agonists are not always effective in patients with severe disease experiencing exacerbation [72], and do not modify the course of the disease [73]. In addition, prolonged use of either medication elicits several side effects [74]. Gene therapy has emerged as a potential alternative or supplementary therapeutic approach that has been investigated in preclinical settings to date; gene silencing approaches are more common, but gene overexpression approaches are also being studied [69].

Unlike CF and AATD, but similar to COPD, there are multiple genetic loci involved in asthma development and progression. This opens many opportunities for gene therapies aimed at a host of target cells, including lung epithelial cells, smooth muscle cells, and immune and inflammatory cells [9]. Using non-viral gene delivery platforms, a broad range of targeted proteins could be simultaneously activated with a single administration, which may be an advantage compared to interventions with small molecules, such as oligonucleotide (ONT)-based knockdown approaches [75]. Repeated dosing with ONT-based therapies failed to show sufficient therapeutic benefit over time [9, 74, 75]. In contrast, plasmid DNA-based strategies could provide long-term therapeutic effects with relatively infrequent administration.

Evidence suggests that airway remodelling results in the progressive loss of lung function in asthmatics [76]. Thus, in addition to the therapies for attenuating airway inflammation, intervention of asthma-associated structural changes in the lung, including smooth muscle hypertrophy, wall thickening and collagen deposition, could reduce the rate of loss of lung function. Recently, da Silva et al. reported a proof-of-concept study where they evaluated effect of the gene encoding an active form of thymulin peptide in a mouse model of asthma [77]. Thymulin peptide has been shown to mediate anti-inflammatory and anti-fibrotic effects in several disease models by modulating T cell differentiation [78]. They found that a single intratracheal administration of a polymer-based non-viral gene vector carrying thymulin plasmid DNA effectively prevented both the inflammatory and remodelling processes in the airways, thereby providing improved airway repair and lung mechanics in a mouse model of allergic asthma. Likewise, a viral gene vector carrying antisense against a cytokine that activates the Th2 pathway, IL-4, reduced airway remodelling in a rat model despite being systemically administered [79]. Overall, multiple targets may need to be tackled to cover the broad spectrum of the diseased population, but promising preclinical studies underscore that gene therapy is an attractive strategy for patients affected by severe asthma.
3. Gene delivery platforms

3.1. Viral gene vectors

Viruses have evolved to infect and transfer genetic payloads to host cells, which makes them attractive candidates for gene therapy applications. The recent approval of adeno-associated virus (AAV) type 1 as a vector for gene therapy in patients with lipoprotein lipase deficiency in Europe (Glybera™) [80] has provided renewed optimism in virus-mediated gene therapy. Numerous clinical trials using adenovirus (AdV) or AAV as gene vectors to treat obstructive lung diseases have been completed, most of which have focused on CF due in part to its monogenic nature [81]. However, gene therapy clinical trials for CF using AdV and AAV2, dosed either intranasally or intratracheally, have failed to result in clinical benefits, and inefficient gene transfer to target cells has been cited as the primary reason [9, 10]. In addition, host immune responses to these vectors were found to limit gene expression after repeated administration of the vectors [10, 31]. However, it should be noted both AdV and AAV2 were well tolerated with limited adverse effects in these studies. Recent preclinical studies aim to more clearly define tissue tropism of the many available viral vectors and their alternative serotypes. In addition, sophisticated optimization of viral capsids and genomes (discussed in section 5.1) are further advancing the capabilities of next-generation viral gene vectors.

3.1.1. Adenovirus (AdV)—AdV was the first viral gene vector tested in inhaled gene therapy clinical trials. AdV is a non-enveloped, icosahedral capsid virion with diameter ranging from 70 – 100 nm. AdV has a genome capacity (36kb) that is much larger than typical viruses [82]. This allows full length CFTR, AATD, and other relatively large therapeutic nucleic acids to be packaged into AdV. AdV does not introduce its nucleic acid payload into the host genome, which results in transient transgene expression [83]. However, lack of DNA integration is a benefit in terms of safety since insertional mutagenesis has been observed with integrating viral vectors [84]. Importantly, the receptor that mediates AdV entry into airway epithelial cells, coxsackievirus-adenovirus receptor (CAR), resides on the basolateral side of the epithelium, which limits AdV’s potential efficacy in vivo for inhaled gene therapy [85, 86]. The innate immune responses, including generation of neutralizing antibodies against AdV, must also be addressed, as repeated administration of AdV will be required for most inhaled gene therapy applications [87, 88]. To begin to address this limitation, helper-dependent or “gutless” AdV (HD-AdV) have been produced wherein all viral DNA has been removed. Preclinical studies with HD-AdV have shown promise [89, 90]. Toeitta et al. showed that HD-AdV administration resulted in reduced inflammatory response and improved airway transduction in mice compared to AdV [91]. Importantly, Croyle et al. showed that gene transduction in mouse lungs with HD-AdV was maintained after a second administration conducted 28 days after the initial dosing [92]. Airway transduction with HD-AdV has also been demonstrated in larger animals, including the ferret [93] and pig [94], further supporting its potential for inhaled gene therapy applications.

3.1.2. Adeno-associated virus (AAV)—Recent viral gene therapy trials have shifted to the use of AAV, as it overcomes many limitations of AdV. For example, AAV provides broad
tissue tropism and more stable transgene expression with partial, site-specific integration into the host genome [10, 95, 96]. AAV is a non-enveloped, icosahedral, non-replicating capsid virus with a diameter of roughly 25 nm [97]. Infection of target cells with AAV is mediated by cell-surface associated glycans, such as sialic acids and/or heparan sulfate proteoglycans depending on the AAV serotype [98–100]. As AAV vectors have shown broad tissue tropism, debate remains over the best-suited AAV serotypes for pulmonary applications [10]. AAV2 was the first discovered serotype [95], and is the only serotype tested in clinical trials of inhaled gene therapy to date [101–103]. Although efficient gene transfer was evident in the nares of humans, lung function in CF patients was not improved [101, 102]. The disappointing outcomes with AAV2 mediated gene therapy in the lungs are likely partially due to the limited capacity of AAV2 to transduce airway epithelial cells via the apical membrane [10]. AAV2, similar to AdV, requires a receptor that is primarily expressed on the basolateral side of the airway epithelium, specifically the heparan sulfate proteoglycan receptor, in order to introduce nucleic acid payloads into airway epithelial cells [99]. A Phase II clinical trial for CF utilizing AAV2 was dropped in 2005 due to inadequate efficacy following repeated administration [103]. To this end, identification and/or engineering of AAV variants with enhanced infection capabilities via the apical membrane, lower immunogenicity, and desired tropism is likely required, especially given the therapy-inactivating immunogenicity generated by repeated administration of AAV2. In Phase I and II clinical trials for AATD, AAV1 and AAV2 dosed intramuscularly did not reach their primary therapeutic endpoints, but rather showed transient and low production of wild-type AAT [47, 49]. Inhaled administration of next generation gene vectors, packaging AAT-encoding DNA, will allow production of AAT proteins at the target of therapy, which may provide a greater therapeutic benefit to patients afflicted with AATD.

Investigations into alternative AAV serotypes, including AAV1, AAV5, and AAV6, have shown promising results, motivating their development for inhaled gene therapy. AAV5 mediated 50-fold greater gene transfer efficiency than AAV2 in air-liquid interface (ALI) culture of primary human airway epithelium (HAE) in vitro in one study [104]. The inherent tropism of AAV5 for airway epithelium results from its interaction with α2,3 N-linked or O-linked sialic acid receptors present on the apical surface of the airway epithelium [98]. Recently, a pseudotyped (hybrid) AAV gene vector, bearing AAV2 rep and AAV5 cap expression cassette (AAV2/5), achieved persistent gene transfer lasting up to 15 months in the airways and alveoli of mice following intratracheal administration (Figs. 1A, B) [105]. They also found that re-administration of AAV2/5 14 months after the initial administration did not significantly reduce the gene transfer efficacy, presumably due to the long dosing interval [105]. As a result of this dosing interval, neutralizing antibody levels were reduced by more than 50% compared to peak levels after the first administration [105]. The process and rationale of pseudotyping viral gene vectors is discussed in detail in section 5.1. Similar to AAV5, AAV1 and AAV6 require interactions with apically expressed receptors, α2,3 and/or α2,6 N-linked sialic acids [100], respectively, in order to transduce airway epithelial cells [100]. An in vitro study with ALI culture of primary HAE revealed that AAV1 exhibited orders of magnitude greater transduction efficiency than AAV2 and AAV5 when the vectors were administered to the apical side of the airway epithelium [106]. AAV1 also exhibited gene transfer efficacy following intratracheal administration in the airways of
chimpanzees that was 20-fold higher compared to that achieved with AAV5, presumably due

to a weaker T-cell response to AAV1 [107]. More recently, AAV6 was found to transduce

mouse and dog airway epithelium in vivo [108], and HAE in vitro [109], to a level

surpassing that achieved with AAV1, AAV2 and AAV5. In addition to the identification of

AAV serotypes with lung tropism, modification of AAV to overcome other key barriers and

therapy-inactivating immunogenicity is needed prior to human testing of next generation

AAV for inhaled applications.

A limitation of AAV is its relatively small packaging capacity for DNA payload (4.7 kb)

[97]. All CF gene therapy trials to date have used the full-length, wild-type CFTR gene to

produce functional CFTR protein. The relatively large size of CFTR cDNA (4.5 kb) has, as a

result, greatly limited the selection of important regulatory elements in plasmid design,

including promoter and enhancer components. [110]. Thus, clinical trials that have tested

AAV-mediated CFTR gene transfer have utilized a weak promoter derived from inverted

terminal repeat (ITR) [110]. ITR is a key element required for packaging therapeutic nucleic

acid payloads into the AAV capsid [111]. Several approaches have been developed seeking
to circumvent this issue. Yan et al. utilized human bocavirus virus-1 (HBoV1) capsids to

create a hybrid AAV virus with a larger packaging capacity (5.5 kb) [112]. In order to

incorporate stronger promoters into AAV, shorter versions of therapeutic genes have also

been engineered with a specific focus on CF gene therapy [113]. Specifically, truncated

CFTR genes that rescue defective CFTR, rather than synthesize the protein de novo, have

been developed [114, 115]. Cebotaru et al. demonstrated that a truncated CFTR, Δ27–264

CFTR, delivered via AAV2/5, provided Δ508 CFTR restoration in vitro in polarized HAE

with chloride currents approaching that of wild-type CFTR [115]. In an earlier report, they

also showed that expression of another truncated CFTR, Δ264 CFTR, led to an increased

level of wild-type CFTR production in the lungs of monkeys [116]. Another strategy to

overcome the packaging size limitation inherent to AAV is to split the genome between two

AAV vectors. This is achieved by either packaging overlapping genomes, which reconstruct

the full length gene through homologous recombination after viral entry, or by evenly

splitting the genome between two vector genomes that then combine after infection through

trans-splicing via heterodimerization [117]. Using the former approach, intranasal

administration of dual AAV6 vectors, carrying overlapping fragments of a reporter gene,

resulted in production of the encoded protein in the mouse lung to a level comparable to

what achieved by a single AAV6 carrying the intact gene [118]. This finding suggests that

the additional recombination step may not significantly impact the efficacy of transgene

expression.

3.1.3. Retro- and lentivirus—Retroviruses have also been investigated as viral vectors

for use in inhaled gene therapy. Unlike AdV and AAV, retroviruses are capable of fully

integrating nucleic acid payloads into the host genome via reverse transcription [119],

thereby potentially providing longer and more stable transgene expression. However,

insertion must be tightly controlled in order to avoid insertional mutagenesis that may result

from random incorporation into host chromosomal DNA [84]. In a prior clinical trial for X-

linked severe combined immunodeficiency, 2 of 4 patients who were successfully treated

with retroviral gene vectors carrying the γc gene developed treatment-related leukemia
Another limitation of retroviral vectors is their inability to infect non-dividing cells, which limits their use for inhaled gene therapy applications given the slow turnover rate of airway epithelium [7].

Other integrating viral gene vectors of interest are lentiviral gene vectors, including recombinant human (HIV) and feline (FIV) immunodeficiency viruses. In contrast to retrovirus, lentiviruses are capable of transfecting post-mitotic cells, including fully differentiated lung parenchymal cells. Lentiviral vectors have a packaging capacity large enough (8 kb) to accommodate full-length genes, such as CFTR [124]. However, lentiviruses have limited tissue tropism and, thus, capsid engineering is required to enable their use in gene therapy applications [124]. Initial preclinical studies have demonstrated efficient, persistent pseudotyped lentivirus-mediated gene expression in the lung [125–131]. For example, a recent study using pseudotyped lentivirus delivered intranasally showed gene expression in the lungs of mice that lasted up to 22 months after initial dosing without signs of toxicity or insertional mutagenesis (Figs. 1C, D) [131]. The modifications used to introduce lung tropism to lentivirus-based gene vectors are discussed in detail in section 5.1. Further investigation into lentivirus-mediated lung gene therapy is warranted given the promising results thus far. However, whether lentiviruses can efficiently overcome physiological barriers unique to inhaled gene therapy of obstructive lung diseases (discussed in section 4) remains to be investigated.

3.2. Non-viral gene vectors

The majority of inhaled gene therapy clinical trials for obstructive lung diseases to date have evaluated virus-based gene vectors. However, intrinsic limitations to their use for gene therapy over the lifetime of a patient, including therapy-inactivating immunogenicity and insufficient gene transfer in human airways to elicit clinical benefits, have spurred interests in development of synthetic systems [132, 133]. Synthetic systems, often referred to as non-viral vectors, are generally formed by multivalent electrostatic interactions between positively charged carrier materials and negatively charged nucleic acids. Unlike some viral vectors, non-viral vectors possess virtually unlimited nucleic acid packaging capacity [137], enabling the delivery of large, multiple and/or diverse nucleic acid payloads. Scale-up of non-viral vectors is relatively straight-forward compared to the complex procedures required for mass production of viral vectors [138]. Non-viral vectors can be altered to impart desired functionalities, such as the ability to penetrate through extracellular barriers [134], target specific cell types [135] and enhance intracellular delivery [136]. It is widely claimed that the primary disadvantage of non-viral vectors is that they provide lower gene transfer efficacy compared to viral vectors [139]. However, few studies have been conducted where advanced non-viral vectors have been directly compared with viruses in vivo, where physiological barriers such as the presence of thick mucus and the immune response may significantly contribute to their performance.

3.2.1. Lipid-based gene vectors—The earliest preclinical evaluation of inhaled non-viral vectors for gene therapy of the airways was conducted with lipid-based systems [140–142]. A seminal study reported by Stribling et al. reported results obtained from nebulized lipid-based gene vectors for inhaled gene therapy [142]. In their paper, aerosolization of a
system based on N-[1-(2,3-dioleoyloxy)propyl]-N,N,N-trimethylammonium chloride (DOTMA)/1,2-dioleoyl-sn-glycero-3-phosphoethanolamine (DOPE) safely mediated transgene expression in the majority of airway epithelial cells and alveolar lining cells at least for 21 days without signs of toxicity [142]. The promise provided by this and other early work [13, 143–147] enabled the prompt translation of the approach to clinical evaluation for inhaled gene therapy, with a specific focus on monogenic disorders, including CF and AATD. There have been 11 clinical trials with non-viral gene vectors for inhaled gene therapy of obstructive lung diseases to date, and all but one involved lipid-based formulations [148]. Based on promising observations in transgenic CF mouse models [13, 149], dimethylaminoethane-carbamoyl-cholesterol (DC-Chol)/DOPE was the first lipid-based system to be clinically evaluated [150]. These studies provided proof-of-concept for lipid-based inhaled gene transfer by showing partial NPD correction and a sign of CFTR transgene expression in several CF patients [11, 150]. However, similar to results with viral vectors tested in CF patients to date, the effects were modest and transient. Thus, the team tested whether repeated administration might provide more sustainable transgene expression [151]. Although clinical benefit was not achieved, the three doses of DC-Chol/DOPE carrying wild-type CFTR were well-tolerated by CF patients without any evidence of immunologic side effects [151]. In addition, gene transfer efficacy was not attenuated by repeated administration [151], unlike viral vectors [10, 31]. Other lipid-based systems, including N-[1-(2,3-dioleoyloxy)propyl]-N,N,N-trimethylammonium chloride (DOTAP) [152] and p-ethyl-dimyristoylphosphatidyl choline cholesterol (EDMPC-Chol), have been tested in humans for inhaled gene therapy of CF. Unfortunately, the results were similar at best to the earlier single dose DC-Chol/DOPE studies [11, 150]. In the first and only clinical evaluation of the inhaled gene therapy of AATD using non-viral vectors, Brigham et al. compared nasally instilled DOTMA/DOPE carrying AAT-encoding plasmid DNA to the standard-of-care for AATD, weekly intravenous injection of AAT protein [48]. The concentration of AAT in nasal lavage fluids of patients who received the inhaled AAT gene therapy was about a third of the normal value that is achieved by AAT protein therapy. However, unlike the protein therapy, inhaled gene therapy reduced the levels of IL-8, a pro-inflammatory cytokine that is elevated in AATD patients [48].

The Genzyme lipid 67 (GL67) is by far the most extensively studied lipid-based gene vector for inhaled gene therapy of obstructive lung diseases. GL67 provided 100-fold greater transgene expression compared to DC-Chol/DOPE [147], which was the first system tested clinically [150]. However, intranasal instillation of GL67, while achieving transgene expression on par with AdV, resulted in significant acute lung toxicity due to the need for high doses [147, 153]. Eastman et al. reported delivery of GL67 via aerosol significantly reduced the toxicity, but empirically determined that very high concentrations were required, which caused undesirable vector precipitation [142, 154, 155]. To address this, they modified the system with a fraction of dioleoulyphosphatidylethanolamine covalently coupled with 5 kDa polyethylene glycol (PEG) [155], yielding a formulation named GL67A. Surface stabilization with PEG was later confirmed by in vitro studies showing that exposure to physiological concentrations of CF mucus components, including albumin, mucin, and linear DNA, did not undermine the gene transfer efficacy of GL67A [156]. Clinical studies with GL67A showed signs of CFTR transgene expression, but flu-like symptoms were
observed in treated patients [11, 157]. To potentially improve the safety profile, Hyde et al. investigated the effect of unmethylated CpG on GL67A-mediated in vivo gene transfer [158]. They demonstrated that while the presence of a single CpG in plasmid DNA was sufficient to elicit a pro-inflammatory response, CpG-free plasmid DNA delivered via GL67A provided sustained transgene expression at least for 56 days without causing lung inflammation. In the meantime, GL67A was compared for in vivo gene transfer efficacy with other non-viral systems, including 25 kDa polyethylenimine (PEI)-based systems and clinically-tested PEGylated poly-L-lysine (PLL) (see section 3.2.1; [159])-based systems [14]. Among these, GL67A exhibited the highest levels of CFTR transgene expression in sheep lungs (Fig. 2A, B) [14]. However, it should be noted that the doses were not matched in this study. Recently, a clinical trial testing repeated administration of GL67A carrying CpG-free CFTR-encoding plasmid DNA was completed [160]. In this multi-dose trial, GL67A provided a significant, but moderate, improvement in the lung function of CF patients, suggesting that an improved gene delivery system is likely required to achieve therapeutically relevant outcomes [33]. Notably, monthly administration of the gene vectors to these patients was well tolerated over the course of the 12-month treatment without any detectable adverse effect [33].

3.2.1. Polymer-based gene vectors—Cationic polymers have also been explored as a means to produce non-viral gene vectors for gene therapy. Among polymer-based systems, PEI is the most extensively studied in the preclinical setting. Linear and branched PEI with molecular weights over 20 kDa have been used most frequently due to their high charge density that enables efficient complexation of nucleic acid payloads [161] and strong buffering capacity that may help the gene vectors escape acidic vesicles inside the cell [162]. Multiple groups have investigated PEI complexes with nucleic acids for inhaled gene therapy [14, 163–169]. Densmore et al. demonstrated that PEI-based gene vectors nebulized into mouse lungs produced a 10–100-fold greater pulmonary transgene expression compared to various lipid-based systems, including GL67/DOPE, DOTAP/DOPE and DC-Chol/DOPE [164]. Importantly, repeated administration at a dose interval of 56 days of 25 kDa branched PEI complexed with CpG-free plasmid DNA further boosted the transgene expression in the mouse lungs [163], suggesting that the effectiveness of repeated treatments was not limited by vector-induced immunogenicity. Nevertheless, clinical use of PEI alone as the condensing polymer in a non-viral vector system has been discouraged by some due to toxicity observed in animals caused by its high positive charge density and nonbiodegradable nature [170, 171].

Interestingly, Boeckle et al. demonstrated that removal of residual free PEI polymers (polymers in a formulation that are not complexed to DNA) by size exclusion chromatography resulted in significantly reduced toxicity and increased transgene expression of gene vectors based on 22 kDa linear PEI [176]. Likewise, removing free PEI by ultracentrifugation resulted in significantly greater pulmonary transgene expression with negligible toxicity in mouse and sheep lungs following aerosolized administration of 25 kDa branched PEI-based gene vectors [169]. These findings suggest that the toxicity of PEI-based systems is largely attributed to uncomplexed polymers, and that complete removal of free PEI from formulations will be a necessary step in future clinical evaluations.
Furthermore, since low molecular weight PEI is less toxic than high molecular weight PEI, several groups have developed PEI derivatives capable of degrading into smaller subunits in physiological conditions, such as in aqueous, acidic and/or reducing environments [172–175]. Of note, albeit not for inhaled gene therapy, PEI has been tested in clinical trials for other gene therapy applications [177, 178].

Poly-L-lysine (PLL) is another well-studied cationic polymer for gene delivery applications. While PLL alone has been associated with limited transfection efficiency and cytotoxicity [160, 179], a specific PLL-PEG based gene vector, namely CK$_{30}$PEG$_{10k}$, has been found safe when administered to human nares. CK$_{30}$PEG$_{10k}$ consists of a 30-mer PLL (30 lysine residues) covalently linked to 10 kDa PEG via a cysteine residue. CK$_{30}$PEG$_{10k}$ has been shown to transfect airway epithelial cells in the lungs of mice (Figs. 2C, D) [181], perhaps owing to its ability to interact with nucleolin on the surface of airway epithelial cells [182], without eliciting significant toxicity [183]. In addition, CK$_{30}$PEG$_{10k}$ was shown to effectively complex plasmid DNA with sizes up to 20 kb [184] and to transfect post-mitotic cells [180]. The promise in the preclinical studies resulted in its evaluation in 12 CF patients who received CK$_{30}$PEG$_{10k}$ complexed with human CFTR-encoding plasmid DNA administered intranasally [159]. A majority of treated patients exhibited partial to complete NPD correction without any noted side effects [159]. The level of gene transfer to the upper airway achieved by CK$_{30}$PEG$_{10k}$ was comparable to the levels observed in trials of AAV2 at the highest titer [102, 159, 185].

4. Physiological barriers to inhaled gene therapy

4.1. Barriers in conducting airways

4.1.1. Mucus gel layer—Initial efforts to improve lung gene transfer focused primarily on efficiently overcoming cellular barriers to DNA delivery [188–190]. More recently, mucus covering the airway epithelium has been recognized as one of the greatest obstacles to overcome [191–195]. Airway mucus is primarily composed of a dense mesh of gel-forming mucin fibers, large macromolecules containing a high density of negatively charged glycans interspersed with periodic hydrophobic regions [196]. Thus, inhaled foreign materials, including gene vectors, are most often immobilized in the mucus blanket via multivalent adhesive interactions (e.g. electrostatic interactions, hydrophobic forces and hydrogen bonding) and/or steric obstruction. Gene vectors trapped in the mucus gel are cleared from the lung via mucociliary clearance (MCC) (3.6 mm/min [197]), which precludes them from efficiently reaching underlying target cells. In the lungs of people with obstructive lung diseases, including CF, COPD and asthma, mucus metaplasia and hypersecretion leads to mucus accumulation and impaired MCC, providing a permissible environment for chronic infection and inflammation [19, 21, 198–200]. Particularly in the CF airways, elevated levels of endogenous DNA and actin filaments released from necrotic neutrophils further contribute to the dense mesh structure of the airway secretions [191, 201].

The average pore size of CF mucus is 140 ± 50 nm (range: 60 – 300 nm) [202], which is markedly smaller than the average pore size of human cervicovaginal mucus secretions (340 ± 70 nm) [203]. Recently, Fahy et al. reported that elevation of oxidative stress in the lungs of CF patients increases disulfide cross-links between mucin fibers that increases mucus
elasticity [204]. The increase in mucin crosslinking density also most likely causes the mucus mesh spacing to tighten further, thereby reinforcing airway mucus as a steric barrier. It is conceivable that other obstructive lung diseases characterized by elevated oxidative stress, including COPD and asthma [205], may share this feature. The viscous drag on gene delivery vectors in the pores alone is not likely to pose a significant diffusion barrier, since the viscosity of the fluid in that fills these pores in normal airway mucus [206] or CF mucus [207] is only moderately higher than that of water. Of note, Coakley et al. have reported that estrogen reduces airway surface liquid height on the CF airway epithelium, which is restored by estrogen antagonist, tamoxifen [208]. Thus, mucus barrier properties may be more pronounced in females. Cigarette smoke coupled with progesterone exposure has also been reported to significantly elevate mucus cell metaplasia and accumulation of eosinophils in an asthma model, while progesterone alone does not [209]. MCC is likely impaired in both cases, which contributes to infection and inflammation in the airway, thereby increasing mucus barrier properties.

Gene vectors that have been used in CF clinical trials recently have been shown to be incapable of efficiently penetrating CF mucus, including AdV [210], various serotypes of AAV including AAV1, 2 and 5 [210, 211], and CK30PEG10K [212, 213]. Likewise, non-viral gene vectors based on the most widely explored cationic polymers, including PEI [214] and polyamidoamine (PAMAM) dendrimers [215], are unable to penetrate CF mucus, most likely due to the their positively charged surfaces that readily interact with negatively charged mucus constituents. It has also been shown that in vitro gene transfer mediated by lipid- and polymer-based gene vectors and AdV was significantly reduced by CF mucus, underscoring its barrier property [216, 217]. Braeckmans et al. have demonstrated that negatively charged, hydrophobic polystyrene nanoparticles strongly adhere to the CF mucus mesh network (Fig. 3A) [207]. Thus, the limited success with gene delivery systems in clinical trials to date may be at least partly attributed to their inability to overcome the mucus barrier. Given the similar pathophysiological events in the airways of patients with other obstructive lung diseases, the mucus blanket is likely to pose a similarly critical barrier to inhaled gene transfer. Indeed, diffusion of nanoparticles in COPD mucus has been shown to be hindered to a similar extent as in CF mucus [201].

Antibody trapping and/or neutralization can also contribute to inefficient gene transfer, specifically for viral gene vectors. Although antibodies diffuse in human mucus relatively unimpeded due to their small molecular size [218], the Fc region of antibodies make low affinity adhesive interactions with mucus [219]. As antibodies accumulate on the surface of pathogen, multivalent antibody interactions with the mucus mesh can trap the pathogen, thereby preventing their penetration to the underlying tissue. For example, Wang et al. recently reported that HSV serotype 1 (HSV-1) readily penetrated fresh neutralized human cervicovaginal mucus, but was trapped therein in presence of anti-HSV-1 immunoglobulin G (IgG), which protected mice against vaginal infection [220]. A significant fraction of CF patients harbor active antibodies against AAV2 (32%) and AdV (55%) [221], which may further enhance adhesive entrapment in CF mucus and/or inactivate their capacity to transduce target cells. Further, neutralizing antibodies against AAV1, 2, 5, 6, 7, and 8 are often found in the airways of healthy people and people with CF [222, 223]. Neutralizing antibodies are also produced in response to administration of viral vectors, and their levels...
are elevated upon repeat dosing [10]. Thus, it is critical to determine if the presence of antibodies affects the mucus penetration rates of AAV in these patient populations.

In addition to its role as a barrier that prevents penetration of gene vectors, mucus may impair the colloidal stability of gene vectors due to the presence of various soluble macromolecules, proteins, lipids, surfactants and ions in the airways. For example, non-viral gene vectors based on cationic polymers and lipids readily interact with negatively charged mucus constituents, resulting in large aggregates. Cationic gene vectors have been shown to aggregate in presence of albumin (the most abundant protein in airway secretions [224]), DNA or mucin [156, 225, 226]. Viruses that possess positively charged surfaces, including HSV [227] and HIV [228], may share similar fates in the mucus gel. The aggregation of gene vectors may reduce gene transfer efficacy by further hindering diffusion in mucus, especially if the aggregates become larger than mucus mesh spacings [229]. Alton et al. showed that even mucus diluted 100-fold markedly reduced gene transfer by AdV and lipid-based gene vectors [230], which is partly attributed to the altered physicochemical properties of gene vectors mediated by soluble mucus constituents [195]. Negatively charged soluble materials in mucus may also compromise non-viral gene vectors by de-stabilizing complexation of DNA with cationic carrier materials. DNA molecules released from their vector are susceptible to degradation by endogenous nucleases present in the airways [231, 232].

4.1.2. Periciliary layer (PCL)—Gene vectors trapped in the mucus gel layer of the airways are cleared by MCC or cough-driven clearance [233, 234]. In contrast, gene vectors that rapidly penetrates through the gel layer and into the periciliary layer (PCL) may be retained significantly longer in the lung, as the PCL is believed to be nearly stationary [235–237]. Button et al. found that the PCL presents a significant steric barrier to vector penetration [238]. Using ALI culture of primary human bronchial epithelial cells, they showed that fluorescent dextran probes larger than 40 nm were excluded from the PCL, while smaller probes partitioned into the PCL and penetrated further toward the epithelium as size decreased [238]. These studies demonstrated that the PCL has a fine mesh structure, as opposed to being just a watery layer as was previously suggested [239], and that nano-sized objects must pay a free energy price to penetrate the PCL [238]. Similar to the mucus gel layer, the PCL may also serve as an adhesive barrier since the meshwork is primarily composed of cell-tethered mucins [238]. Kesimer et al. showed that the PCL excluded AdV (~100 nm in diameter), but not AAV (~30 nm in diameter; serotype not specified; Fig. 3B) [240]. In CF lungs, dehydration of the PCL mediated by dysregulation of epithelial sodium channels (ENaC) on the airway epithelium can cause an osmotically-driven collapse of the PCL [238, 241]. Collapse of the PCL likely further increases the barrier function to inhaled gene vectors by making the PCL mesh tighter. Despite being relatively less explored, airway dehydration appears to be a common concern for other obstructive lung diseases, including COPD [242] and asthma [243]. Overall, gene vectors capable of penetrating the mucus gel layer but not the PCL would most likely be cleared via MCC [214] or by macrophages [244].
4.2. Barriers in airspace

4.2.1. Pulmonary surfactant—Mucus is not present in the alveolar sacs of the lungs. However, inhaled gene vectors that make it to the alveoli must retain their stability and function in presence of surfactants that are abundant in the airspace. Pulmonary surfactant is a surface-active lipoprotein complex, synthesized by type II alveolar epithelial cells, that is composed of various phospholipids, cholesterol and surfactant proteins, including SP-A, B, C and D. To date, studies exploring the effect of pulmonary surfactants on lung gene transfer have primarily involved non-viral gene vectors [170, 245–251]. Gene transfer mediated by cationic lipid-based gene vectors was significantly reduced by pulmonary surfactants [248–251]. The stability of lipid-based gene vectors in the presence of various surfactants, including Alveofact (an extract from bovine lung lavage) and Exosurf (a synthetic surfactant) and individual components of pulmonary surfactant was investigated. Negatively charged phospholipids [248–251] and/or SP-B or C [250] were found to facilitate aggregation of, or DNA release from, lipid-based gene vectors, indicative of impaired colloidal stability or disruption of DNA-lipid complexation, respectively. While to lesser extents, Alveofact has been shown to reduce gene transfer efficacy of gene vectors formulated with cationic polymers, including PEI and PAMAM dendrimer [170, 248].

Studies have also demonstrated surfactant-mediated aggregation of polymeric gene vectors [245, 246], however, DNA complexation was maintained with cationic polymer-based gene vectors, unlike lipid-based vectors, in the presence of pulmonary surfactants [248]. This carrier material-dependent effect is likely due to the difference in the nature of complexation by cationic lipids and polymers that influences vector stability in pulmonary surfactants. In addition to the disruption of complexation by negatively charged surfactants, hydrophobic lipid-surfactant interactions may render lipid-based vectors more susceptible to destabilization. Interestingly, incubation of polymeric gene vectors with pulmonary surfactants has been shown to enhance gene delivery efficiency in vitro due to improved cellular uptake and/or increased cell membrane permeability [246, 247]. Similarly, surfactants have been found to enhance AdV-mediated gene transfer to peripheral lung cells in vitro and in vivo [252, 253]; lipid recycling in pulmonary epithelial cells was suggested as a potential mechanism [253]. These findings suggest that the effect of pulmonary surfactant on respiratory gene transfer may be multimodal and may vary depending on gene vector type.

4.2.2. Alveolar macrophages—Alveolar macrophages are phagocytes residing in the airspace that play a critical role in homeostasis, host defense and tissue remodeling [254]. There are 12 –14 macrophages in each alveolus [255], which is increased in individuals who smoke regularly [256]. Alveolar macrophages engulf inhaled foreign substances directly or via an opsonin-dependent mechanism. Particles in the size range of 250 nm to 3 µm are readily phagocytosed by macrophages [257], with increasing phagocytic uptake with increase in the particle size within this range [258]. In contrast, particles smaller than 250 nm are taken up less efficiently by macrophages [257], including alveolar macrophages [259]. These findings suggest that alveolar macrophages may not pose a significant hurdle for gene vectors, since widely explored viral and non-viral gene vectors are generally smaller than 250 nm. However, aggregation of gene vectors in the presence of pulmonary surfactants is likely to impair their stability and function, which may further reduce their efficacy.

J Control Release. Author manuscript; available in PMC 2017 October 28.
surfactants may render them more susceptible to clearance by alveolar macrophages due to their increased size. For example, 20 and 110 nm silver nanoparticles delivered via aerosol to rat lungs were found as large aggregates that were engulfed by macrophages collected from bronchoalveolar lavage fluid (BALF; Fig. 3C) [260].

Other properties of gene vectors, including surface charge, composition and particle geometry, play a critical role in phagocytosis [261]. In the case of polymeric nanoparticles, studies have shown that particles with hydrophobic surfaces are more readily internalized by macrophages than those with hydrophilic surfaces, while surface charge did not appear to significantly alter the phagocytosis [261, 262]. Surfactant proteins such as SP-A and SP-D can opsonize inhaled gene vectors [263], which may facilitate opsonin-dependent phagocytosis by alveolar macrophages. Immunoglobulin (i.e. antibody) and complement are primary opsonins present in the lungs [264]. It has also been found that some viral vectors, including retrovirus and AdV, can be rapidly internalized by alveolar macrophages [265, 266]. It is possible that viral vectors are recognized by alveolar macrophages via Fc receptors on the surface of macrophages that bind to antibodies attached to the virus. Although it is a different organ, transient depletion of Kupffer cells, resident macrophages in the liver, enhanced AdV-mediated gene transfer in vivo [267]. In addition, it has been shown that complement cleavage fragment C3α and C3β opsonize non-viral gene vectors composed of cationic polymers or lipids [268], which may facilitate macrophage uptake via complement receptors.

Vector-mediated inflammation through recruitment of neutrophils and macrophages [269] may further reduce gene transfer efficiency upon repeated administration. Alveolar macrophages may pose a more challenging barrier in the lungs of patients afflicted by diseases characterized by chronic infection and inflammation, such as CF [270] and COPD [271], where alveolar macrophages are activated. It has been shown that alveolar macrophages significantly reduce retrovirus-mediated gene transfer to human bronchial epithelial cells; the inhibitory effect is elevated by lipopolysaccharide-induced macrophage activation [265].

4.3. Cellular barriers

Gene vectors that overcame the aforementioned extracellular barriers must then be taken up by target cells to introduce nucleic acid payloads to the intracellular gene expression machinery. Airway epithelial cells are a primary target for treatment of CF, and are of general importance to most obstructive lung diseases. The epithelial surface of the airways poses an additional barrier to inhaled gene therapy that is due to low efficiency of endocytosis across the apical membrane [188] and tight junctions between cells that prevent access of gene vectors to the basolateral side [272] (Fig. 3D). This is a particularly formidable challenge for AdV since its receptor, CAR, is selectively localized on the basolateral membrane of airway epithelium [86]. Preclinical studies of lentivirus-mediated gene therapy in the lungs have also shown limited transduction through the apical membrane [273].

It has been reported that the barrier property of the airway epithelium is altered by obstructive lung diseases, primarily due to structural perturbations of tight junctions [274–
In asthma, the barrier function of the airway epithelium is impaired through defective tight junction formation [274]. Likewise, cigarette smoke exposure, the major risk factor for COPD, disrupts tight junctions and increases epithelial permeability [275]. In contrast, it has been hypothesized that tight junction proliferation in CF results in increased epithelial resistance [276]. The transepithelial electrical resistance (TEER) is higher and the paracellular permeability is lower in CF airway epithelial cell cultures compared to cultures expressing wild-type CFTR [277]. However, pro-inflammatory cytokines [278] and bacterial toxins [279] reduce the permeability of tight junctions, suggesting that the barrier property of the airway epithelium may vary with disease state. Tight junction proteins in the alveolar epithelium, including claudins [280], also limit access of inhaled gene vectors to the basolateral surface. Once taken up by target cells, gene vectors must overcome several intracellular barriers, including but not limited to acidic vesicles (i.e. endosomes and lysosomes), the molecularly crowded cytoplasm, and the nuclear envelope. These barriers are shared in numerous organs and tissues in addition to the lungs and, thus, are widely reviewed elsewhere [281, 282].

5. Strategies to overcome the barriers to inhaled gene therapy

5.1. Modification of gene vectors

Tuning gene vectors to overcome one or more important physiological barriers can enhance the efficacy of pulmonary gene therapy. Inhaled gene vectors deposited on conducting airways first encounter the luminal mucus gel layer that serves as a highly adhesive and steric barrier, as described in section 4.1.1. Thus, gene vectors must be small enough to traverse through the mesh spacing of airway mucus, while possessing particle surface resistant to muco-adhesion [133]. It has been reported that nanoparticles as large as 200 nm efficiently diffuse in human airway mucus freshly collected from individuals with [211] or without [206] obstructive lung diseases, but only if particle surfaces are densely passivated with hydrophilic and neutrally charged PEG polymers. Of note, high PEG densities that yield brush conformations, as opposed to mushroom shapes, provides muco-inert particle surfaces [229, 283].

Based on these findings, polymer-based gene vectors capable of efficiently penetrating human airway mucus, namely mucus-penetrating DNA nanoparticles (DNA-MPP), have been introduced [214, 215, 284]. Suk et al. demonstrated that DNA-MPP based on PEI and PLL, unlike the otherwise identical counterparts without dense surface PEG coatings, efficiently percolated through pathological human airway mucus ex vivo [214], followed by a similar observation with DNA-MPP formulated with biodegradable poly(β-amino ester) (PBAE) (Figs. 4A, B) [284]. Likewise, the dense surface coatings with PEG corona may improve the penetration of inhaled gene vectors through another mucin-based meshwork found in PCL, if the particle diameters are small enough to fit through the PCL pores [240, 285]. The PEG surface coating may also minimize the particle aggregation in physiological environments [229], thereby providing another means of improving penetration through the steric barriers, including the mucus gel layer and PCL. Importantly, rapid ex vivo diffusion of DNA-MPP in airway mucus was translated to widespread distribution (Figs. 4C, D) and/or prolonged retention [214] in the mouse lungs, leading to approximately 25-fold
greater in vivo pulmonary transgene expression compared to leading non-viral gene vectors without dense surface PEG coatings, including PEI- and CK30PEG10k–based systems [284].

Given that several viral vectors possess muco-adhesive surfaces [210, 211], their diffusion within the airway mucus is likely enhanced by a dense surface shielding with PEG, similar to the findings with non-viral gene vectors. The enhanced lung gene transfer efficacy of AdV by surface PEG conjugation has been previously reported [286, 287]; however, the benefit of PEG in these studies was experimentally determined to be either reduced immunogenicity or resistance to the neutralization by pre-existing antibody. Thus, the effect of PEG on the ability of viral vectors to overcome key physiological barriers to inhaled gene therapy is yet to be determined. It should be noted though that interference of PEG with antibody binding will reduce Fc-mediated trapping of viral vectors in airway mucus. Other than the PEGylation approach, Schuster et al. have recently shown that AAV2 mutant possessing the capsid with reduced heparin binding exhibit significantly enhanced diffusion in human CF mucus compared to that of native AAV2 [211]. This is most likely attributed to the decrease in heparan sulfate-mediated adhesion of AAV2 to CF airway mucus rich in these proteoglycans [211].

Targeting cells of interest via specific ligands is certainly a viable approach to enhance gene transfer to the lung. Although the majority of studies involves cancer targeting due to numerous well-established pathways upregulated in cancers [288], several groups have reported feasibility of specifically targeting parenchymal cells in the lung, including airway and alveolar epithelial cells. Most of the relevant studies involve identifying targeting ligands for airway epithelial cells, reflecting the dominance of CF-related research. Following the confirmation of urokinase plasminogen activator receptor (uPAR) expression on the apical surface of differentiated HAE, Drapkin et al., coupled a 7-mer peptide derived from a respective ligand to the surface of AdV via PEG [289]. They found that the targeting AdV provide 10-fold greater gene transfer to airway epithelium in vitro compared to native and PEGylated AdV. Employing the phage display technology, Jost et al., identified a 7-mer peptide, THALWHT, that targets human epithelial cell lines [290]. Subsequently, an independent group genetically engineered AAV2 decorated with THALWHT and found that this mutant provided a significantly greater in vitro transgene expression both in undifferentiated and polarized (i.e. differentiated) HAE [291]. However, they were unable to identify any ligand harboring the sequence that targets polarized HAE and the mutant did not provide significantly increased transgene expression in vivo, implying that the sequence may not be airway-specific or may be species-dependent. Tagalakis et al. evaluated in vivo gene transfer efficacy of a lipid-based gene vector decorated with a peptide sequence previously identified to target intracellular adhesion molecule-1 (ICAM-1), a receptor for rhinovirus which causes the common cold [292]. They demonstrated that while their ICAM-1 targeting formulation provided 99% of airways with evidence of bronchial epithelial cell transfection, 73% and 38% of airways showed epithelial expression with 22 kDa PEI and GL67, respectively, following intratracheal administration [292]. Similarly, lactoferrin [293] and lactose [294] have been shown to enhance the transgene expression of PEI- and PLL-based gene vectors, respectively, in the HAE. However, both studies were conducted using immortalized cells, and thus the validity of the approach should be confirmed with ALI culture of primary HAE. There are relatively few studies describing the
targeted gene transfer to alveolar epithelial cells. In one study, β2-adrenoceptor agonist, clenbuterol (Clen), which is used as bronchodilator for COPD and asthma treatment, was incorporated onto the surface of PEI-based gene vectors [295]. The Clen-decorated system provided 14- and 3-fold higher transfection efficiency compared to non-targeted counterpart in alveolar epithelial cells in vitro and in mouse lung in vivo, respectively.

In addition to incorporating specific ligands to the surface by covalent conjugation [289] or genetic engineering [291], other approaches have been widely explored to endow viral vectors with ability to be internalized by cells of interest. Directed evolution approaches have been used to generate libraries of alternative viral capsid types and screened for tropism based on transduction efficiency in vitro or in vivo [296–298]. Rational design, specifically capsid pseudotyping, has also been used where viral capsids are replaced by those from other virus types or serotypes known to efficiently infect cells in the lung to provide or enhance lung tropism. Given the promising preclinical and clinical results, AAV has been the most extensively engineered viral vector to date using both directed evolution and rational design approaches [296–298]. The rational design approach has also been used to engineer novel AdV and lentiviral vectors for inhaled gene delivery applications [299–301].

Libraries of mutant AAV variants have been generated using a variety of methods, including error-prone polymerase chain reaction, DNA shuffling or random insertion/deletion of peptide-encoding sequences [302]. Specifically for inhaled gene therapy applications, AAV mutant vector libraries can be screened in vitro using ALI cultures of HAE and/or in vivo with suitable animal models to determine which variants successfully mediate gene transfer. After multiple rounds of selection in ALI cultures of HAE, a mutant vector with shuffled cap genes from both AAV2 and AAV5 was identified with enhanced airway tropism and mediated 100 and 10-fold greater in vitro transgene expression as compared to native AAV2 and AAV5, respectively [303]. Using a similar approach, a mutant AAV vector containing shuffled cap genes from AAV1, AAV6, and AAV9 was identified by screening in polarized HAE and showed 3-fold greater in vitro production of CFTR mRNA transcripts than native AAV6 and ~25% restoration of CFTR as compared to healthy controls [304]. By mutating single amino acids in the heparin binding domain of AAV6 capsid, Limberis et al. discovered a novel AAV6 mutant gene vector, AAV6.2, outperforming other native and mutant AAV at mediating gene transfer in mouse airways (Figs. 4E, F) and polarized HAE cultures (Figs. 4G, H) [109].

Using the pseudotyping approach, hybrid viral vectors have been engineered that contain envelope or capsid proteins from other viruses with alternative tissue tropisms to change gene transfer properties [296–298, 301]. AdV and AAV hybrid vectors have been generated with capsid proteins from alternative viral serotypes that use receptors expressed on the apical surface of airway epithelium for entry. For example, a pseudotyped AdV vector was engineered to incorporate AdV35 fibers into the AdV5 capsid, redirecting tissue tropism towards airway epithelium with viral entry mediated by CD46 receptors expressed on the apical membrane [305]. A pseudotyped AAV vector containing an AAV2 genome packaged in AAV5 capsid proteins (i.e. AAV2/5) allowed for targeting of apically-expressed sialic acid receptors [116]. Using aerosolized AAV2/5, Fischer et al. demonstrated efficient gene
transfer to the airways of monkeys to a level 20-fold greater than what achieved in historical AAV2 studies [116].

Incorporating components from the capsid of other respiratory viruses, such as bocavirus [112], baculovirus [9] and Sendai virus (SeV) [9, 130, 306], is also a common strategy to introduce lung tropism. In addition to improving the packaging capacity, AAV2 pseudotyped with human bocavirus-1 (HBoV1) capsid demonstrated 5.6- and 70-fold greater efficiency at transducing polarized HAE culture in vitro compared to native AAV1 and AAV2, respectively [112]. They were able to transduce via the apical surface of primary HAE isolated from ΔF508/ΔF508 homozygous CF patient, resulting in ~30% restoration of CFTR-mediated chloride currents [112]. Initial inhaled gene transfer studies with lentivirus focused on vectors pseudotyped with vesicular stomatitis virus G glycoprotein (VSV-G) [125–129]. However, this approach was generally developed to broaden the tropism of lentivirus rather than to specifically design a system for inhaled gene therapy applications [124]. It was later found for VSV-G-pseudotyped lentivirus that pre-treatment with compounds disrupting tight junction is required for gene transfer to the airway epithelium, suggesting that the transduction was achieved through the basolateral surface [125, 129]. To bypass the need for adjuvant treatments, hybrid lentiviral vectors were engineered by incorporating envelope proteins from SeV or baculovirus that have demonstrated tropism for the apical surface of airway epithelium [9]. Alton et al. were able to apically transduce in vitro in polarized primary CF HAE cultures and in vivo in CF mouse nasal epithelium using SeV-pseudotyped SIV [130]. The transgene expression in mouse nasal epithelium lasted up to 1 year after a single administration with no sign of immune response (Figs. 1C, D). This approach has significantly advanced the use of lentiviral vectors for inhaled gene therapy applications.

5.2. Modulation of biological barriers

Adjuvant agents that reduce airway and/or cellular barrier properties provide a relatively simple means to enhance the efficacy of inhaled gene therapy, potentially without the need of modifying gene vectors. However, it is crucial to ensure that any adjuvant approaches do not cause significant toxicity or disrupt normal lung function. This is of even greater concern for patients afflicted with obstructive lung diseases having impaired lung function. To our knowledge, specific strategies to reduce the barrier properties of the airspace are yet to be introduced, and as described in section 5.1, gene vector modification serves as the primary method to address this issue.

The most widely explored approach in this category is the use of mucolytic agents that degrade primary macromolecular components of airway mucus. Two primary compounds of selection may be recombinant human DNase (rhDNase, dornase alfa, Pulmozyme®) and N-acetyl cysteine (NAC, Mucomyst®), which are currently or previously utilized in the clinic to help CF patients clearing accumulated mucus in their airways. NAC’s mode of action is to cleave disulfide intermolecular crosslinks between mucin fibers, which can significantly reduce the viscoelasticity of airway mucus [307]. As previously discussed, airway secretions in patients with obstructive diseases can also carry high levels of DNA, further enhancing its
barrier properties. Enzymatic degradation of DNA by rhDNase reduces entanglements within the mucus gel to further decrease the viscoelasticity [307, 308].

In the context of inhaled gene therapy, NAC and rhDNase can enlarge the mesh spacings of airway mucus, reducing the physical obstruction of inhaled gene vectors. For example, NAC was shown to increase the average pore size of CF mucus from 145 ± 50 nm to 230 ± 50 nm (Figs. 5A, B) [309]. It has been reported that pre-treatment with NAC leads to more rapid diffusion of leading gene vectors, including AAV1 [211] and CK30PEG10k –based system [213], through CF mucus. Accordingly, NAC significantly improved CK30PEG10k –mediated gene transfer in the lungs of a lipopolysaccharide-induced mouse model of mucus hypersecretion [213]. Likewise, a NAC derivative was shown to enhance gene transfer efficacy of AdV [310] and non-viral gene vectors [216], including EDMPC-Chol and PEI-based systems, in the mouse lung in vivo and in an ex vivo sheep trachea model, respectively. Suk et al. previously demonstrated that diffusion of nanoparticles as large as 200 nm in CF mucus was significantly improved by NAC, but the effect was greater when nanoparticles possessed muco-inert surface coatings [309]. This finding suggests that simultaneously modulating gene vectors and the mucus barrier may synergistically improve mucus penetration and thus the efficacy of inhaled gene therapy. In addition to clinically used agents, Yuan et al. showed methyl 6-thio-6-deoxy-α-D-galactopyranoside reduced levels of reactive oxygen species in CF mucus and provided a greater reduction of disulfide crosslinks in CF mucus compared to NAC [204]. A novel alginate oligosaccharide compound, currently in a Phase Iib clinical trial for CF, was also shown to widen the pores within CF mucus by disrupting mucin-DNA interactions [311]. While safety must be confirmed, these newly developed mucolytic agents may also be useful as adjuvants for improving the penetration of gene vectors through the airway mucus barrier.

Osmotic agents are another type of agent that may effectively reduce the barrier property of the airway mucus blanket. In particular, hypertonic saline is a clinically used osmotic agent to rehydrate the airways of CF patients and improve MCC [312]. Although it is yet to be tested, hypertonic saline may be used as an adjuvant to inhaled gene therapy, as it can potentially reduce the barrier property of airway mucus by diluting this gel layer. Improved MCC is likely achieved by the restoration of the collapsed PCL in the CF airways (Figs. 5C, D) [238], and thus inhaled hypertonic saline treatment may also render the PCL more permeable to inhaled gene vectors. Graeber et al. demonstrated that 3% and 7% hypertonic saline administered via aerosol greatly reduced mucus accumulation in the airways of a ENaC-overexpressing transgenic mouse model of obstructive lung diseases (to be discussed in section 6.1) [313], presumably by increasing airway hydration and improving MCC. Mannitol (Bronchitol™) is another hypertonic osmotic agent that hydrates airways [314] and thus may also be useful as an adjuvant to inhaled gene therapy. The kinetics of hydration by inhaled hypertonic agents must be carefully evaluated in relevant preclinical or clinical settings in order to determine an adequate time interval between the pre-treatment and gene vector administration. Administration of gene vectors in hypotonic solution as a vehicle may also improve gene vector penetration through the mucus barrier via convective flow generated by the osmotic gradient established between the airway lumen and epithelial cells. Ensign et al. demonstrated that by using hypotonic solution as delivery vehicle, muco-inert (i.e. densely PEGylated) nanoparticles were able to rapidly penetrate the luminal mucus.
layer and reached the immediate surfaces of vaginal [315] and colorectal [316] epithelium in vivo. However, the effect and safety of this approach is yet to be established for delivery of inhaled therapeutics, including gene vectors.

Use of hypotonic vehicle solution has shown potential of reducing the cellular barrier to inhaled gene therapy as well. Huang et al. demonstrated that hypotonic shock enhanced uptake of plasmid DNA by nasal epithelium in vivo via the regulatory volume decrease (RVD) mechanism [317]. Specifically, they showed that sodium- and sucrose-based hypotonic vehicles provided enhanced transgene expression in the mouse nasal epithelium compared to isotonic vehicles with the greater effect observed with lower osmolality. Similarly, other studies have shown that fluorescently labeled compounds administered to airway epithelium in hypotonic vehicles were efficiently internalized by cells, whereas the same molecules in isotonic vehicle were not [318–320]. In addition to the observations in conducting airways, Sawa et al. showed an enhanced transgene expression in the lung airspace of rats when plasmid DNA was intratracheally instilled in hypotonic, rather than isotonic, vehicle solution [321]. RVD occurs after cell swelling driven by hypotonic shock [322], stimulating the fusion of numerous intracellular vesicles with the plasma membrane to prevent cell lysis [323, 324]. The response involves release of intracellular ions and subsequent water loss by osmosis [325], leading to the internalization of excess apical and basolateral membrane to reform the lost intracellular vesicles. During this endocytic process, particulates, including inhaled gene vectors, in the vicinity of the apical membrane can be taken up by epithelial cells. The RVD effect has been shown to last up to 30 minutes [317, 319]. Potential safety concern resulting from hypotonic shock would need to be addressed before clinical implementation of this approach.

As discussed earlier, several viral vectors possess natural tropism towards the basolateral surface of airway epithelium [85]. Thus, transient disruption of epithelial tight junctions may enhance transgene expression mediated by these vectors. Further, both viral and non-viral gene vectors may benefit from this approach due to the low rates of endocytosis across the apical membrane [188]. It has been shown that pre-treatment with fatty acid surfactants, including polidocanol (PDOC), sodium caprate (C10) and lysophosphatidylcholine (LPC), can increase paracellular permeability by transiently opening the epithelial tight junctions (Figs. 5E, F). Transduction by AdV was greatly enhanced in vivo in mouse nasal and airway epithelium pre-treated with PDOC [326] and C10 [327]. Likewise, pre-treatment with LPC has been shown to enhance VSV-G-pseudotyped lentivirus transduction in vitro in polarized HAE culture [273] and in vivo in marmoset [125] and ferret [129] airways. Calcium-chelating agents, such as ethylene glycol-bis(beta-aminoethyl ether)-N,N,N',N'-tetraacetic acid (EGTA), can also be used as adjuvants to reduce the intracellular concentration of calcium ions, thereby disrupting calcium-dependent formation of tight junction protein complexes [328]. Pre-treatment with EGTA has been shown to enhance efficacy of AdV and retrovirus gene transfer in vitro in polarized HAE cultures and in vivo in rabbit tracheal epithelium [329]. However, there have been concerns over the safety of this approach and potential adverse effects. To investigate this, Johnson et al. evaluated the safety of EGTA, C10 and sodium laurate (C12), in vitro in primary HAE cultures and in vivo in the lungs of mice where they found evidence of causing a minimal to mild inflammatory response, based on histopathological analysis [330]. However, this study revealed that EGTA increased cell
counts and the levels of pro-inflammatory cytokines in BALF, while C10 and C12 altered airway responsiveness after methacholine stimulation in vivo in the lungs of mice [330]. LPC has been used most extensively as an adjuvant to VSV-G-pseudotyped lentivirus-mediated inhaled gene therapy to enhance gene transfer with minimal adverse effects reported [125, 129].

6. Preclinical models

As discussed in previous sections, we have now learned through past successes and failures of preclinical and clinical studies the potential hurdles towards achieving therapeutically relevant inhaled gene therapy of obstructive lung diseases. Relevant preclinical models are needed in order to reliably evaluate newly developed strategies to better predict their performances in clinical trials. First, those models should replicate the physiological barriers found in the lungs of respective diseases to confirm whether the approaches of interest provide the benefits as designed. In addition, general pathology of respective human diseases should be reproduced in the animal models for assessing the pharmacodynamics (i.e. therapeutic gene transfer efficacy). In this section, we discuss the most advanced animal and tissue culture models available for evaluating gene delivery strategies.

6.1 Cystic Fibrosis

Development of genetic animal models has greatly enhanced our understanding of CF lung pathophysiology. The CFTR-null (i.e. knockout) mouse was the first type of CF animal model established that results in a complete loss of functional CFTR [331–333]. This was followed by development of transgenic mouse models that produce low levels of CFTR [334, 335] and specific-mutant CFTR including but not limited to ΔF508 [336], G551D [337] and G480C [338]. However, spontaneous generation of CF lung phenotype with mucus accumulation as well as chronic infection and inflammation is rare in knockout and transgenic mouse models with only one reported case [339]. This may in part be due to the lack of submucosal glands in the lower airways beyond the trachea of mouse lungs [340, 341]. Of note, submucosal glands are the primary source of the airway mucus secretion in the human lungs [200] and plays critical roles in innate immunity by secreting antimicrobials [342]. A recent study also found an alternative chloride channel is present in the mouse lung that can compensate for the lack of CFTR activity [93].

It has been found that defective CFTR leads to hyperabsorption of sodium through dysregulated ENaC channels present on the apical surface of CF airway epithelium [343–345], and has been implicated as an initiating pathological event in the CF lung. Based on these findings, Livraghi-Butrico et al. have shown that transgenic mice overexpressing β-subunit of ENaC spontaneously established CF-like lung disease, which resembles the pathological features of human CF lungs, including airway inflammation and dehydration as well as mucus obstruction [346–348]. Chronic lung infection, a hallmark of CF, has been established in normal mice and transgenic mouse model with specific CFTR mutation, including R117H, S489X, Y122X and ΔF508, using Pseudomonas aeruginosa-laden agarose beads to prolong the bacterial residence time within the lung [349, 350]. Interestingly, inflammatory response to P. aeruginosa infection did not appear to depend on mutation type.
of the CF transgenic mouse models included in the study [349]. Mice intranasally challenged with endotoxin derived from *P. aeruginosa* have been shown to establish lung inflammation, mucus cell metaplasia and mucus hypersecretion [213], providing another relevant mouse model that mimics common features of the CF lung.

Larger genetic animal models have been developed using pig [351, 352], ferret [353–355] and rat [356] as background species, offering physiological features better reflecting the lungs of human CF patients [357–359]. It has been shown that pigs with mutated CFTR and/or CFTR knockout develop lung defects resembling those found in human CF lungs, including inflammation, impaired bacterial clearance, airway remodeling, and mucus hypersecretion (Figs. 6A, B) [351, 352]. A CFTR-null ferret model also shares features similar to humans with CF, showing altered airway chloride transport, mucus hypersecretion and propensity for lung infection [353–355]. These larger animal models are ideal for testing inhaled gene therapy, as they better recapitulate the physiological barriers and pathological events established in the human CF lungs. However, larger animal models are relatively costly, and accompany greater ethical challenges, and thus may be best suited for the later stages of preclinical development. A recent development of CFTR-knockout rats provides a model with more extensive development of submucosal glands in the trachea which may lend itself towards development of CF-like lung phenotype [356], thereby potentially serving as an alternative to aforementioned models.

### 6.2 α-1 antitrypsin deficiency

The most widely employed animal models for AATD to date were established in the 1980’s [360–366]. These models are transgenic mice constitutively expressing a mutated version of an essential anti-protease, human AAT (hAAT), with a glutamate-to-lysine mutation at position 342 of the hAAT [367]. The mouse models harboring mutant hAAT exhibit low levels of hAAT in serum, caused by accumulation of the mutant protein in the endoplasmic reticulum of hepatocytes, similar to what is found in AATD patients [363]. However, none of these studies clearly demonstrated the development of emphysema, a key pathological characteristic in the lungs of human AATD patients. Thus, an advanced preclinical model that more closely mimics AATD lung phenotype is needed to reliably evaluate therapeutic outcomes of inhaled gene therapy. For example, emphysema can be established in mouse and rat lungs by disrupting the protease/anti-protease balance via an instillation of different types of proteases into the lungs, including, but not limited to, papain, human neutrophil elastase, porcine pancreatic elastase and galactosamine [368–371]. It should also be noted that imbalance in protease activity is a common characteristic of a broader disease, COPD [372]. However, the lack of significant inflammation and airway changes [373] limit their use for studies that require general pathology of COPD, including chronic bronchitis. Other clinically relevant COPD animal models are discussed in section 6.3.

### 6.3 Chronic Obstructive Pulmonary Disease

There are a number of mouse-based COPD models, including cigarette-smoke (CS), protease-induced models, and genetic models. The most widely utilized models are generated by exposing rodents including mice and guinea pigs to CS [372, 374]. CS is exposed daily either only to the nose or whole body [375] for an extended period of time,
usually weeks to months, to establish COPD-like pathology, including elevated oxidative stress [376], chronic inflammation [377], small airway remodeling and/or emphysema (i.e. enlargement of the airspace compartment; Figs. 6C, D) [378]. The nose-only administration allows for more direct control of amount of CS exposure compared to whole body exposure, but mandates prolonged restraint of the animals, which may cause stress. Using either administration modality, the effects of CS exposure can be closely monitored based on levels of CS exposure markers such as carbon monoxide-hemoglobin complexes (i.e. carboxyhemoglobin) and nicotine metabolites (e.g. cotinine) present in serum [375].

Using nose-only CS exposure, Beckett et al. recently demonstrated that an 8-week exposure of CS to mice twice per day and 5 times a week resulted in COPD-like pathology, including airway inflammation, emphysema and impaired lung function (determined by forced oscillation and forced maneuver techniques), which was not resolved during the following 4 weeks of CS cessation [379]. The lungs of this model also became susceptible to acute bacterial infection, as evidenced by decreased clearance of *Streptococcus pneumoniae* and influenza virus by 2-fold in comparison with wild-type mice [379], presumably replicating pulmonary exacerbation often observed with COPD patients [380]. However, mucus accumulation and airway obstruction are not established in this model, despite the presence of goblet cell metaplasia and airway remodeling [379]. As a result, this model may be limited in evaluating therapeutic intervention of the chronic bronchitis phenotype.

CS models have been also generated using other larger species such as guinea pigs [381, 382]. Guinea pig-based CS models have been favored by several groups due to its unique advantages, including relatively close resemblance to human lung physiology and anatomy [383–385] as well as lung alveolarization at birth which may be beneficial for pre- and neonatal studies [386]. In addition, hallmark features of COPD including inflammation [387, 388], goblet cell metaplasia [389, 390], small airway remodeling [391, 392], airway obstruction [393] and emphysema [394], have been successfully demonstrated in guinea pigs. COPD has been extensively studied over the years with the use of various CS-induced animal models reported by numerous groups, but development of a standard CS exposure protocol seems necessary as it is currently difficult to directly compare studies by different groups.

The COPD animal models based on genetic variations include natural mutant, transgenic and knockout models. Some of these models may offer the advantage of consistency, however they do not always reflect the COPD pathogenesis specific to the lungs and may adversely affect other organs [373]. Interestingly, some naturally occurring mutant strains of C57BL/6 mice, such as tight skin and pallid mouse models, are known to have abnormally large airspace, lower serum level of AAT and impaired alveolar septa [369, 395]. Tight skin mouse model is characterized by a mutation in the fibrillin-1 gene, which affects formation of elastic fibers, leading to abnormal airspace development and progressive alveolar enlargement with age [396, 397]. Pallid mouse model is known to have lower level of AAT in serum and gradually develops mild emphysema late in life [369]. The pallid mouse model may also be utilized for AATD studies [398]. Transgenic and knockout models may be more useful for studying a particular pathway of the COPD pathogenesis, as exact roles of specific genes can be elucidated in these models.
6.4 Asthma

The ovalbumin (OVA)-challenged mouse model is by far the most widely utilized preclinical model for allergic asthma, exhibiting the characteristic Th2-type immune response [399]. The models are generally established by intraperitoneal sensitization with OVA, followed by repeated intratracheal OVA challenges, but exact dosing schedule varies among individual studies [400]. The model reliably recapitulates asthma-like pathophysiology, including recruitment of eosinophils, elevated levels of OVA-specific IgE and Th2 cytokines, mucus accumulation by mucous and goblet cell hyperplasia, airway hyperresponsiveness and airway remodeling (Fig. 6E) [401, 402]. Although the OVA mouse model is the most widely used, there are questions regarding its clinical relevance as OVA is not a naturally occurring allergen. Hence, it may not properly mimic how asthmatic patients become sensitized to allergens [403]. Numerous groups using this model have reported that chronic asthmatic symptoms do not exacerbate over time; the symptoms are actually ameliorated by natural development of tolerance and/or termination of OVA challenge [404, 405]. More recently, the development of tolerance to repeated challenges has been addressed by optimizing amount and frequency of OVA administration [77, 400]. Although not as frequently used, rats and guinea pigs are also utilized as background species of OVA models. Rat models share similar immunological cascade as the mouse model when sensitized with OVA and may perhaps be used as an alternative [406]. Guinea pigs, while closely resembling the human lung anatomy and physiology, have been reported that IgG is the primary antibody produced post-sensitization and the baseline level of eosinophils is high, which may somewhat limit their use [406, 407].

Mouse models have been also developed using naturally occurring allergens such as house dust mite (HDM) [408]. Many groups favor HDM models over OVA models as it yields pathophysiology similar to that exhibited by the more widely-studied OVA-sensitized model, while also having greater environmental/clinical relevance [409]. For example, Johnson et al. administered HDM intranasally for 5 days per week for up to 7 weeks and observed eosinophilic inflammation, elevated levels of Th2 cytokines, as well as airway hyperresponsiveness and remodeling [401, 409]. Additionally, in contrast to OVA-challenged animal models, re-exposure of HDM to mice resulted in sustained inflammatory response, including elevated eosinophil and Th2 effector cell counts, suggesting that tolerance to HDM did not develop [409]. However, after cessation of HDM exposure, airway inflammatory response fully recovered, while alterations in airway remodeling and hyperreactivity persisted [409]. HDM consists of a mixture of allergen subtypes including Der p I and Der p II and other biomolecules such as endotoxins, proteases, proteins and peptides, making it difficult to identify the component(s) responsible for specific allergic responses [401, 410, 411]. Moreover, many commercially available HDM extracts are different in composition and proteolytic activities [412], and thus direct comparison of various HDM studies can be difficult.

Use of transgenic mice overexpressing GATA3 transcription factor may be considered. GATA3 is known to play significant roles in Th2 cell differentiation and activation by controlling Th2-driven cytokine production [413]. Several groups demonstrated that mice with GATA3 overexpression, in comparison to its normal counterparts, showed upregulation
of Th2 cytokines, airway remodeling, airway smooth muscle hyperplasia, eosinophilic inflammation and subepithelial fibrosis, upon sensitization and challenge with allergens [414–417]. Overexpression of GATA3 alone induces production of Th2 cytokines and leads to lung inflammation and mucus hypersecretion, but not to the extent exhibited after allergen challenge [413]. Sensitization of GATA3-overexpressing transgenic mice with even relatively low amount of OVA or HDM cause further increases in airway inflammation, while no significant inflammation was observed in wild-type C57BL/6 mice [413]. In another study, Ano et al. sensitized and subsequently challenged GATA3-overexpressing transgenic mice with OVA and were able to observe enhanced airway inflammation and goblet cell hyperplasia [414]. A Phase IIa trial of an inhaled DNAzyme targeting GATA3 mRNA have been recently initiated for Th2-driven asthma [418], underscoring the relevance of these transgenic mouse models.

Larger animal models based on non-human primates, ponies, and Basenji greyhounds have been developed since they develop natural allergies as well as persistent respiratory allergic responses; however, they have not been widely explored primarily due to cost issues [419].

6.5 Human Tissue Culture Models

Therapeutic efficacy and safety of novel delivery strategies must be ultimately demonstrated in preclinical animal models prior to their clinical evaluation. However, recent studies have shown species-specific differences in tropism of AAV by comparing transduction in vitro in ALI cultures of human lung epithelium to those of mouse [420], monkey [359], ferret [359] and pig [421] lung epithelium. These findings underscore the importance of confirming the efficacy in relevant human cell-based models as compared to preclinical models. Studies in human tissue culture models are also beneficial for high-throughput screening of gene vector candidates and further optimization of vector design. Human lung epithelial cells grown in ALI cultures have the ability to differentiate into polarized epithelial layers, developing key physiological barriers including tight junctions between neighboring cells and mucus secreted from the apical surface within ~4 weeks [422, 423]. Ussing chamber experiments can be conducted to confirm the presence of intact tight junctions determined by the measured TEER [422, 423] and also to assess CFTR correction via quantifying the chloride conductance [26, 115, 277].

ALI cultures have been generated from lung epithelial-derived cell lines, including Calu-3, CFBE, NuFi-1, and CuFi-1 (for complete list of available cell lines, see [424]). However, concerns arise around the use of ALI cultures established with a single cell type, as there are several different cells present in the lung (i.e. ciliated epithelial cells, goblet cells, etc.) [424–426]. Mucus production and PCL development may also vary greatly depending on the cell lines used [424–426]. Thus, the properties of relevant delivery barriers encountered by gene vectors may not fully recapitulate their characteristics in the human lung. Primary ALI cultures are an attractive alternative as they are established with airway cells directly collected from human lung explants and thus retain many features of the physiological human lung epithelium [238, 241, 427–430]. Another benefit is that epithelial cells can be harvested directly from specific patients of interest for screening gene delivery strategies, potentially enabling personalized medicine. Shortcomings of primary ALI cultures include
limited availability of fresh explants, short cell lifespan, and variability among patient donors. It should be also noted that submucosal glands, an important target tissue in obstructive lung diseases, are not present in ALI cultures regardless of the source of constituting cells.

Recent developments in nasal brushing-derived primary lung epithelial ALI cultures could provide a more readily accessible tissue source [431]. However, there are concerns over its use as a surrogate tissue source as studies have shown marked differences in genomic profiles of primary human nasal and bronchial epithelial cells [432, 433]. Primary human organoids differentiated from patient-derived stem cells are also an attractive in vitro model for high-throughput screening [434]. Specifically for CF, the organoid’s degree of swelling in response to the treatment with a well-documented CFTR activator, forskolin, was found to correlate with CFTR function, providing a means to assess CFTR activity and/or correction [435]. The organoid systems remain viable after long periods of storage in liquid nitrogen and could be passaged up to 40 times, beneficial for long-term studies. More recently, this approach was expanded to human lung organoid systems consisting of proximal lung epithelium surrounded by mesenchymal smooth muscle tissue and have been shown to generate airway-like features with both club and ciliated cell development [436]. However, the barrier properties of organoid systems have not been thoroughly characterized and may differ from those observed in the human lung.

7. Conclusion

Over two decades of preclinical and clinical evaluations of inhaled gene therapy have provided valuable lessons building towards the ultimate goal of developing curative treatments for patients with obstructive lung diseases. We have now established better understanding of disease pathology, genetic targets and physiological barriers. The accumulated knowledge has driven the development of advanced gene delivery systems, nucleic acid engineering tools and human disease-like preclinical models by scientists, engineers and clinicians in various settings. A recently completed clinical trial of inhaled CF gene therapy has left some questions to be addressed, but has certainly rejuvenated the field with more clinical evaluations anticipated in the near future. Through this review, we hope to provide a comprehensive overview of the knowledge gained and encourage collaborative efforts for realizing therapeutically effective gene therapy of obstructive lung diseases. We began our review by overviewing the types of obstructive lung diseases and potential genetic targets for respective diseases. We then highlighted gene delivery systems, primarily those tested in clinical trials of inhaled gene therapy, followed by challenging physiological barriers that have hampered translation into the clinic. We next introduced strategies to overcome those hurdles and preclinical disease models resembling the pathological lung environments of diseased human patients. Future preclinical studies of the next-generation of gene vectors in animal models that present all these critical features will help better predict therapeutic outcomes and thus facilitate clinical development of inhaled gene therapy.
Acknowledgments

The National Institutes of Health (R01HL127413) and Cystic Fibrosis Foundation provided funding for this work. The content is solely the responsibility of the authors and does not necessarily represent the official view of the NIH. We thank Julie Shade for contributing to the artwork used in the graphical abstract.

References


J Control Release. Author manuscript; available in PMC 2017 October 28.


133. Suk JS. Could recent advances in DNA-loaded nanoparticles lead to effective inhaled gene therapies? Nanomedicine (Lond). 2016


J Control Release. Author manuscript; available in PMC 2017 October 28.


investigation; a journal of technical methods and pathology. 2007; 87:893–902. [PubMed: 17592477]


435. Dekkers JF, Wiegnerink CL, de Jonge HR, Bronsveld J, Janssens HM, de Winter-de Groot KM, Brandsma AM, de Jong NWM, Bijvelds MJC, Scholte BJ, Nieuwenhuis EES, van den Brink S,

Figure 1. Viral vectors for inhaled gene therapy

(A, B) A recombinant AAV2/5 (AAV2 rep gene, AAV5 cap gene) delivered by intratracheal instillation demonstrates long-lasting gene expression (up to 15 months) in both conducting airways and alveoli of mice. (A) Immunohistological staining of β-galactosidase (β-gal) expression in alveoli and conducting airways 1 month post-administration. (B) Bioluminescence imaging of firefly luciferase expression in the lung and nose of mice at 1, 3, 6, 12, and 15 months post-administration. Reprinted from [105] with permission of Mary Ann Liebert, Inc.

(C, D) A simian immuno-deficiency virus pseudotyped with the respiratory pathogen Sendai virus (F/HN-SIV) demonstrates sustained transgene expression in the nose and lungs of mice after intranasal administration, lasting 22 months post-administration. (C) Fluorescent microscopy images of GFP expression mediated by F/HN-SIV in mouse lungs. (D) Bioluminescence imaging of firefly luciferase expression in the mouse lungs and noses at 2 and 22 months post-administration. Reprinted from [131] with permission of the American Thoracic Society. Copyright © 2016 American Thoracic Society.

J Control Release. Author manuscript; available in PMC 2017 October 28.
Figure 2. Non-viral gene vectors for inhaled gene therapy

(A, B) Localization of gene expression in sheep lungs treated with GL67A carrying human CFTR (hCFTR)-expressing plasmid DNA. (A) Dual labeling of two hCFTR epitopes (G449, Texas red; MATG1061, FITC) of lung sections. Arrows indicate epithelial cells that are positive with both antibodies. (B) Dual labeling of cytokeratin and G449 (anti-cytokeratin antibody, red; G449, FITC). Reprinted from [14] with permission of Macmillan Publishers Ltd. (C, D) Localization of gene expression in the (C) medium (20× magnification) and (D) small airways (40× magnification) in the mouse lungs. Lungs harvested from animals that received 100 µg of β-gal expressing plasmid DNA compacted with CK30 PEG10k were fixed, sectioned, and immunohistochemically stained for the bacterial β-galactosidase protein 2 days after intratracheal administration. Reprinted from [181] with permission of Macmillan Publishers Ltd.
Figure 3. Primary physiological barriers to inhaled gene therapy

(A) Mucus: A confocal image showing 89-nm polystyrene nanoparticles (PS NP) trapped via adhesive interactions within CF mucus. Reprinted from [207] with permission from Elsevier.

(B) The periciliary layer (PCL): Adenovirus (AdV, blue arrows) is excluded from the PCL while adeno-associated virus (AAV, red arrows) penetrates into PCL and reaches underlying epithelium. Reprinted from [240] by permission from Macmillan Publishers Ltd.

(C) Alveolar macrophages: an SEM image showing aerosolized 110 nm silver nanoparticles (Ag NP) accumulating in macrophages collected from bronchoalveolar lavage fluid (BALF) after being administered to rats. Reprinted from [260] by permission from Oxford University Press.

(D) Epithelial cell tight junctions: A confocal image (top: xy view, bottom: xz view) showing AdV restricted to the apical side of human airway epithelium due to the
presence of tight junctions which prevents the access to receptors required for cell entry.
American Thoracic Society.
Figure 4. Modification of gene vectors to overcome physiological barriers
Representative trajectories of (A) conventional DNA nanoparticles (DNA-CP) and (B) mucus-penetrating DNA nanoparticles (DNA-MPP) based on biodegradable PBAE polymers in freshly expectorated CF mucus. Representative images of gene vector distribution in large airways following intratracheal administration of (C) DNA-CP and (D) DNA-MPP. Reprinted from [284] with permission from PNAS. Copyright © 2016 National Academy of Sciences, USA. (E, F) In vivo and (G, H) in vitro transduction of AAV6 and a mutant variants, AAV6.2, engineered with a single amino acid substitution in the heparin
binding domain. Comparison of intratracheally administered (E) AAV6 and (F) AAV6.2 showing a stronger β-gal transgene expression in the mouse lung with AAV6.2. Comparison of (G) AAV6- and (H) AAV6.2–mediated GFP expression in ALI cultures of primary HAE. AAV6.2 treated cultures show stronger GFP expression as well as transduction of both ciliated and nonciliated cells (H, inset). Reprinted from [109] by permission from Macmillan Publishers Ltd.
Figure 5. Modulating physiological barriers to inhaled gene therapy

(A, B) N-acetyl cysteine (NAC) treatment increases mucus mesh spacing, thereby facilitating gene vector penetration through airway mucus. Reprinted from [309] with permission from Nanomedicine as agreed by Future Medicine Ltd.

(C, D) Reducing the osmotic pressure (OP) of mucus that can be achieved inhaled hypertonic saline rehydrates and restores the collapsed PCL. Reprinted from [238] with permission from AAAS.

(E, F) Sodium caprate (C10) disrupts tight junctions in epithelial layer allowing gene vectors to access the basolateral compartment where specific receptors required for viral gene
Figure 6. Animal models of obstructive lung diseases
Airway obstruction develops in CF pig due to highly viscoelastic mucus (A; white arrow) in the airway where mucus cytology further revealed presence of neutrophils, macrophages, and bacteria (B). Reproduced from [351] with permission from AAAS. Airspace in (C) healthy guinea pigs and (D) guinea pigs exposed to cigarette smoke (CS) for 6 months that show airway enlargement characteristic of emphysema in COPD. Reproduced from [374] with permission from APS. (E) Mucous cell hyperplasia, subepithelial fibrosis, and smooth
muscle hypertrophy in ovalbumin (OVA)-induced allergic asthma mouse model. Reproduced from [77] with permission from Elsevier.
Table 1

<table>
<thead>
<tr>
<th>Gene vector type</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Clinical Development Stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenovirus (AdV)</td>
<td>Large nucleic acid packaging capacity (36 kb) compared to AAV [81]</td>
<td>Transient transgene expression [82]</td>
<td>Clinical trials for CF [9]</td>
</tr>
<tr>
<td></td>
<td>Non-integrating; no concerns over insertional mutagenesis [82]</td>
<td>Requires basolaterally-expressed receptors for cell entry [84,85]</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Immune response limits efficacy upon re-administration [9]</td>
<td></td>
</tr>
<tr>
<td>Adeno-associated virus (AAV)</td>
<td>Non-pathogenic [10]</td>
<td>Limited nucleic acid packaging capacity (4.7kb) [96]</td>
<td>Clinical trials for CF [100–102] and AATD [46,48]</td>
</tr>
<tr>
<td></td>
<td>Multiple serotypes (e.g. AAV1, 5 and 6) able to enter airway epithelial cells via apical cell surface receptors [94]</td>
<td>Immune response limits efficacy upon re-administration [10]</td>
<td></td>
</tr>
<tr>
<td></td>
<td>More stable gene expression compared to AdV [94,95]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lentivirus</td>
<td>Long-term, stable gene expression [118]; lower dosing frequencies are achievable [130]</td>
<td>Safety concerns over insertional mutagenesis [119]</td>
<td>Pre-clinical [124–130]</td>
</tr>
<tr>
<td></td>
<td>Capable of transfecting post-mitotic cells [121,122], unlike retrovirus [120]</td>
<td>Requires vector engineering for lung applications [123]</td>
<td></td>
</tr>
<tr>
<td>Lipid- and Polymer-based</td>
<td>Facile chemical modification [137]</td>
<td>Cytotoxicity [153, 173]</td>
<td>(Lipid) Clinical trials for CF and AATD [148]</td>
</tr>
<tr>
<td></td>
<td>Minimal constraint in packaging capacity [151, 184]</td>
<td>No intrinsic tropism [139]</td>
<td>(Polymer) Clinical trials for CF [35]</td>
</tr>
<tr>
<td></td>
<td>Large-scale production [186]</td>
<td>Occasional stability issue [137]</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Variety of FDA-approved drug delivery formulations [187]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Controlled release is achievable [139]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Table 2
Physiological barriers to inhaled gene therapy of obstructive lung diseases and strategies to overcome them

<table>
<thead>
<tr>
<th>Regions of Lung</th>
<th>Primary Barriers</th>
<th>Gene Vector Modification Strategies</th>
<th>Barrier Modulation Strategies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducting Airway</td>
<td>Adhesive and/or steric trapping of inhaled gene vectors by mucus [202] and PCL [238]</td>
<td>Engineer gene vectors with surface coatings to reduce adhesive interactions with mucus (e.g., PEG [214, 215, 284], mutant AAV2 [211]) Sizes below characteristic mucus (≤50 nm) [202] and PCL (≤80 nm) [238] mesh pore size</td>
<td>Mucus-altering agents (e.g., rhDNase [307, 308], NAC [309]) to increase mucus mesh pore size</td>
</tr>
<tr>
<td>Airspace</td>
<td>De-stabilization of inhaled gene vectors by pulmonary surfactants [248–251] Uptake of inhaled gene vectors by alveolar macrophages [257–267]</td>
<td>Engineer gene vectors with surface coatings (e.g., PEG [214, 215, 284]) to enhance particle stability and/or to reduce macrophage uptake</td>
<td>None reported</td>
</tr>
<tr>
<td>Cellular</td>
<td>Low endocytic rate on apical surface [188] Restricted access to basolateral surface due to the presence of tight junctions [86, 273, 280]</td>
<td>Introduce targeting ligands to enhance apical transfection [288–295] Engineer viral gene vectors to endow apical epithelial tropism (e.g., pseudotyping [296–298, 301], directed evolution [302])</td>
<td>Tight junction disrupting agents (e.g., C10 [327, 330], C12 [330], EGTA [328, 330], LPC [125, 129, 273], PDOC [326]) to provide an access to the basolateral surface Hypotonic vehicles to enhance uptake by the RVD mechanism [317–324]</td>
</tr>
</tbody>
</table>