REVIEW ARTICLE

Lung gene therapy—How to capture illumination from the light already present in the tunnel

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Abstract Gene therapy has been considered as the most ideal medical intervention for genetic diseases because it is intended to target the cause of diseases instead of disease symptoms. Availability of techniques for identification of genetic mutations and for in vitro manipulation of genes makes it practical and attractive. After the initial hype in 1990s and later disappointments in clinical trials for more than a decade, light has finally come into the tunnel in recent years, especially in the field of eye gene therapy where it has taken big strides. Clinical trials in gene therapy for retinal degenerative diseases such as Leber’s congenital amaurosis (LCA) and choroideremia demonstrated clear therapeutic efficacies without apparent side effects. Although these successful examples are still rare and sporadic in the field, they provide the proof of concept for harnessing the power of gene therapy to treat genetic diseases and to modernize our medication. In addition, those success stories illuminate the path for the development of gene therapy treating other genetic diseases. Because of the differences in target organs and cells, distinct barriers to gene delivery exist in gene therapy for each genetic disease. It is not feasible for authors to review the current development in the entire field. Thus, in this article, we will focus on what we can learn from the current success in gene therapy for retinal degenerative diseases to speed up the gene therapy development for lung diseases, such as cystic fibrosis.

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Introduction

In 2008, three research teams independently reported the success in clinical trials of gene therapy treating a rare form of retinal degenerative diseases called Leber’s congenital amaurosis (LCA). LCA represents a group of inherited blindness with childhood onset. The clinical success has been achieved in treating LCA2, one form of the disease, which is caused by mutations in the retinal pigment epithelium-specific 65-kDa protein gene (REP65). REP65 encodes a protein providing the isomerohydrolase activity for the retinal pigment epithelium to produce 11-cis-retinal from all-trans-retinyl esters during the visual cycle for regenerating the visual pigment after exposure to light. Without this gene function, 11-cis-retinal, the natural ligand and chromophore of the opsins of photoreceptor cells, cannot be regenerated, thus rendering the opsins incapable of capturing light or transducing it into electrical responses for initiating vision. Although this defect in light transduction has an immediate impact on visual function, retinal cell degeneration is delayed in patients, thus making target cells available for gene therapy. The three teams tested the same therapeutic approach in patients by subretinal injection of recombinant adeno-associated virus vector 2 (AAV2) expressing the RPE65 complementary DNA (cDNA). Patients with treatment showed improvements in visual function without serious adverse events. In 2012, three patients received the same treatment in their other eye and all three demonstrated improvements in visual and retinal function in their second eyes after the treatment, which was administered one-and-a-half to three-and-a-half years after their first eyes were treated. Readministration of the same gene therapy vector caused no harmful immune reactions in patients. In 2014, a gene therapy trial for another retinal degenerative disease, choroideremia, was shown to be successful. Choroideremia is an X-linked recessive disease that is caused by mutations in the CHM gene, which encodes the Rab escort protein 1 (REP1). The same gene therapy vector, AAV2, was used in this study. In addition to the eye gene therapy success, progress has been made in other fields as well. For example, as a milestone for using gene therapy as medicine, European Union approved Glybera as the first gene therapy drug for a form of lipoprotein lipase deficiency. In this case, AAV1 was used to deliver a naturally occurring functional variant of the LPL gene associated with lower rates of cardiovascular disease and increased efficiency in fat metabolism. These clinical successes provide the proof of concept that the power of gene therapy can be harnessed to benefit human beings.

However, gene therapy developments for other diseases, such as cystic fibrosis (CF) lung disease, are not as successful for eye diseases. CF is the most common monogenic fatal disorder in the Caucasian population and it is caused by recessive mutations in the gene encoding the cystic fibrosis transmembrane conductance regulator (CFTR). Although the disease affects multiple organs, including the lung, pancreas, intestine, gall bladder and reproductive organs, lung failure due to chronic infection and inflammation is currently responsible for most morbidity and mortality. Therefore, CF gene therapy studies to date have been aimed at treating the pulmonary manifestations. When the cystic fibrosis gene was identified in 1989, it appeared that this disease can be used as an ideal model for the development of gene therapy for lung diseases since airway epithelial cells where the CFTR gene is expressed are readily accessible to gene therapy vectors. Yet, all the CF clinical trials conducted so far did not show any evidence of significant therapeutic benefits brought to CF patients. Basic research in lung gene therapy developments later identified major barriers to vector delivery and sustained therapeutic gene expression. Thus, it is useful to look into what is fundamental to the successful gene therapy development for eye diseases to make lung gene therapy fruitful.

In this review article, we will first visit the early developments in CF lung therapy and look into the major challenges encountered in the lung gene therapy field. We will then review the key factors that are critical to the eye gene therapy progress to explain the possible rationale for the clinical success. We will finally discuss strategies that can be translated from the eye gene therapy field to speed up the lung gene therapy development.

Early stages of lung gene therapy developments

Because CF is a monogenic disease and the target cells in lung airway are easily accessible to gene therapy vectors, when the gene was identified, an illusion was created suggesting that lung gene therapy for CF would be available in a few years. The initial excitement inspired many scientists racing in conducting clinical trials. Both viral and non-viral gene therapy vectors were tested. One of the early clinical studies was conducted by Zabner et al in 1993 to examine the safety profile of an adenoviral (Ad) vector with nasal applications. Adenoviruses contain a linear double-stranded DNA and have been widely used as tools for gene delivery because of their ability to infect both dividing and non-dividing cells with a high efficiency, especially epithelial cells. The early generations of Ad vectors were developed by deleting the E1 region within the viral genome to prevent viral proliferation in transduced cells and/or other regions such as E3 or E4 to increase the DNA carrying capacity. There are more than 50 serotypes of adenoviruses identified so far. In this study, a serotype 2 adenoviral vector expressing the human CFTR cDNA was administered to a defined area of nasal epithelium in three patients. Although this initial study showed some functional correction in nasal epithelial cells with no vector-related adverse effects, more extensive studies later demonstrated with similar methods that there was no significant functional correction in nasal epithelia. The Ad vectors have also been tested in the lung and none of the studies demonstrated functional correction or efficacy in patients.

In addition to the early generations of Ad vectors, recombinant adeno-associated virus (AAV) vectors have also been tested in CF patients. AAV is a replication-defective parovirus that depends on a helper virus, either adenovirus or herpes virus, for its propagation during lytic infection. It has a small single-stranded DNA genome (about 4.7 kb). The advantage of AAV as a gene therapy
The results were not encouraging. Delivery methods were tested in the lung of CF patients and composed of a cationic lipid and a neutral lipid, are leoylphosphatidylethanolamine (DOPE) or cholesterol. 10

replacement therapy.

early generations of Ad vectors are not suitable for gene therapy since these vectors are considered to be safer than viral vectors. 10 Cationic liposomes, which are composed of a cationic lipid and a neutral lipid, are commonly used for gene delivery. There are many types of cationic lipids available which are often mixed with one or two commonly used neutral lipids, dioleylphosphatidylethanolamine (DOPE) or cholesterol. 10

Many clinical trials have been conducted with cationic liposomes to assess their potential for CF Therapy. The first liposome CF trial was carried out in CF patients through nasal administration and later, several other trials with various liposome formulations were conducted. 30–33 Similar delivery methods were tested in the lung of CF patients and the results were not encouraging. 34,35

Challenges that slowed the progress in the lung gene therapy

Looking back at the aforementioned clinical studies, it is now understandable; the reason for the lack of clinical progress in the lung gene therapy for CF was that major challenges to lung gene delivery were not fully appreciated at the time. These challenges include 1) lacking efficient vectors for airway gene expression, 2) host immune responses to vectors, 3) lacking efficient and safe methods for vector delivery and 4) difficulty in maintaining long-term therapeutic gene expression in airways where epithelial cells turn over. The following sections will be devoted to explain these challenges as well as progresses being made towards overcoming these challenges.

Efficient vectors for airway gene expression

It is apparent that all gene therapy vectors used in early clinical studies of lung gene therapy were not adequate for achieving the efficiency needed for functional correction in patients. The first generation Ad vectors used in the clinical studies could not confer sustained transgene expression in vivo. The issue of lacking sustained transgene expression from adenoviral vectors is often misunderstood because it was interpreted as the lack of vector stability in vivo. In fact, Ad vectors are stable in vivo; the reason that they cannot confer sustained transgene expression is because they elicit host immune responses (which will be addressed later), which eliminate the transduced cells in vivo. Although most Ad vectors have the E1 region deleted from their genomes to prevent viral gene expression and proliferation, leaky expression of viral genes does occur in transduced cells 42,43 thus providing antigen for the host immune system to attack the transduced cells. Therefore, early generations of Ad vectors are not suitable for gene replacement therapy.

To reduce the host immune responses, helper-dependent adenoviral (HD-Ad) vectors have been developed. 36,44 In these vectors, all the viral coding sequences have been deleted, thus eliminating the leaky viral gene expression and rendering a large DNA carrying capacity to the vectors. These vectors have also been called high capacity or gutless vectors 45 and have been shown to confer long term transgene expression when delivered to mouse livers. 46 Compared to the first generation Ad vectors, HD-Ad vectors elicited reduced levels of inflammation and conferred longer term of transgene expression when delivered to mouse lungs. 47 Our group has demonstrated that the human CFTR gene can be efficiently delivered to mouse lungs using the HD-Ad vector and that the CFTR knockout mice treated with the CFTR expressing vector are protected from acute lung infection with bacteria. 48 We have also showed that HD-Ad vectors can be used to deliver genes to the lungs of rabbits 49 and pigs. 50 We have showed that the human CFTR gene can be efficiently delivered to pig lungs. 50 However, up to date, HD-Ad vectors have not been tested in human lungs.

The reasons for the AAV vector failing to meet the efficiency required for CF lung gene therapy are different from that of Ad vectors. One of the major limitations is the small DNA carrying capacity of AAV vectors; once the CFTR cDNA is packed into the vector, there is no room for carrying DNA regulatory elements, such as cell-specific promoter or enhancer, for CFTR gene expression. 51,52 It was reported in 2008 that AAV vectors could deliver as much as 8.9 kb DNA although with a reduced efficiency, 53 but later it was found that a single AAV particle could not deliver the large reporter genes and the cell transduction in this case may be accomplished by two viral particles containing the 5’ and 3’ parts of the reporter genes. 54 In addition, the AAV2 vector is not very efficient for lung gene delivery. 55

One strategy to expand the DNA carrying capacity is to rely on trans-splicing of mRNAs in target cells. A large gene can be packaged into two AAV vectors to generate two RNA transcripts which are spliced into one functional mRNA 56,57 Since this strategy required two AAV vectors to carry each part of a gene to transduce the same cell and it is not expected that all RNA transcripts are trans-spliced, the transgene expression efficiency will likely be reduced. Although the DNA carrying capacity of AAV vectors is not easy to change, different serotype AAV vectors with better transduction efficiency for lung airway cells can be selected. For example, AAV5, AAV6, AAV9 and AAV6.2 are better vectors for lung gene delivery 58 than AAV2 that has already been used in clinical trials. In addition, vector tropism can be engineered for lung applications. 59,60

The lack of success in using nonviral vectors for clinical studies of CF lung gene therapy is understandable because, unlike the viral vectors, there is no specific mechanism for nonviral vectors to send the DNA payload into the nuclei of target cells and perpetuate the existence of the therapeutic gene in the nuclei. For gene delivery with liposomal vectors, the majority of the delivered DNA is degraded by lysosomes before entering nuclei; nuclear entry is another major barrier. 60 Although non-viral vectors might be useful for some other gene delivery applications, in our opinion, it is difficult to use them for lung gene replacement therapy due to the lack of efficiency in gene delivery. Although
Host immune responses

Host immune responses to gene therapy vectors are one of the most important challenges that were previously underestimated in early lung gene therapy development. The lung is a very sensitive immunologic organ capable of producing both strong innate and adaptive immune responses to pathogens and gene therapy vectors. Although the adaptive immune response to viral vectors was recognized early on and investigated extensively, the innate immune response was not fully appreciated. This was evident from the first and the only incidence of gene therapy related death in 1999 when a patient who received a high dose of an Ad vector via the hepatic artery succumbed to acute toxicity. Post-mortem confirmed that the patient suffered from systemic inflammation, biochemically detectable disseminated intravascular coagulation, and multiple organ failure within 98 h.66 Clearly this was the result of patient’s innate immune system reacting to the high dose of the Ad vector.

For airway gene delivery, the first innate immune response is mediated by lung macrophages that quickly engulf vector particles, reducing gene delivery to target cells. The macrophages not only take up gene therapy vectors and destroy them, but also initiate the production of proinflammatory cytokines by interacting with lung epithelial cells.67 Macrophages as well as epithelial cells express Toll-like receptors (TLRs) and RIG-I-like (retinoic-inducible gene-1-like) receptors, which recognize nucleic acid and proteins derived from viral pathogens including viral DNA, single-stranded RNA and double-stranded RNA.68 Inflammatory cytokines, such as TNF-α, IL-6, MIP-2 and MIP-1α, are dramatically induced in macrophages upon Ad vector delivery to mouse airways within 6 h.45,69 These cytokines activate airway immune cells and structural cells to produce more proinflammatory cytokines which could lead to airway damage, if the host cannot shut down the cascade. One of the important pathways involved in the induction of inflammatory cytokines is the NFκB signaling pathway.70 A variety of inflammatory cytokines, such as IL-6 and IL-8, can be induced by the activation of NFκB. In humans, a high level of IL-8, a potent neutrophil chemoattractant, can cause neutrophil infiltration that leads to tissue damage.71,72 Even by using non-viral vectors, bacterial DNA can be recognized by Toll-like receptor 9 which activates the NFκB pathway.16,45 It has been recently reported that cytosolic DNA is a danger signal that induces interferons through the production of cyclic guanosine monophosphate–adenosine monophosphate (cyclic GMP–AMP, or cGAMP).73 Other immune cells, such as neutrophils and natural killer cells, are also recruited in response to viral vector delivery, which can produce inflammatory mediators as well and cause tissue damage. The innate immune response not only leads to acute toxicity, but also enhances the adaptive immune response (will be explained later).

One major strategy to avoid the host immune responses is to avoid using unnecessary high vector dose. Since all gene therapy vectors elicit innate immune responses, the lower the dose the weaker immune responses triggered in the host. In addition, anti-inflammatory drugs may be used transiently to reduce the innate immune responses around the first few days following vector delivery. Finally, noninvasive vector delivery methods should be used to reduce host stress to dampen the host innate immune responses. These points will be further discussed later.

The adaptive immunity relies on T- and B-lymphocytes to produce cellular and humoral responses to infectious agents. In mammals, the adaptive system is developed postnataally. During the development, an extremely diverse repertoire of receptors is generated randomly and each receptor recognizing a unique antigen, is expressed on the surface of one T lymphocyte only. T cells bearing useful receptors are subsequently selected from billions of lymphocytes for clonal expansion by interacting with antigens. Professional-antigen-presenting cells, normally dendritic cells or macrophages present the antigens bound to the MHC II molecules to helper T cells to activate them. Activation of helper T cells by antigen presenting cells also requires a co-stimulatory signal, e.g., CD80 or CD86, on the surface of the antigen-presenting cell to bind to CD28 on the surface of the T cell.74 The expression of co-stimulatory molecules is regulated by innate immunity,75 and therefore, the adaptive immunity is regulated by the innate immunity. Helper T cells control other cells in the adaptive immune system, such as activation of cytotoxic T cells to destroy infected cells and B cells to produce antibodies. After elimination of pathogens, some antigen-specific clones of T and B cells remain as "memory" lymphocytes so that the adaptive immune system remembers the antigens and destroy them more quickly upon subsequent exposure.

Early work from several groups demonstrated clearly that both cellular and humoral responses are involved in adenoviral vector-mediated gene transfer in mice. Repeated delivery of viral vectors, or primary delivery to individuals with pre-existing immunity, or viral vectors expressing foreign antigens, can cause strong adaptive immune responses. Various strategies can be used to overcome the host adaptive immune responses. First of all, as described above, blocking the innate immune response can reduce the adaptive immune response. Secondly, blocking co-stimulatory pathways can be used to modulate the host adaptive immune responses. Several groups showed that an antibody against CD40 ligand or expressing CTLA4Ig, a fusion protein of cytotoxic T lymphocyte-associated protein 4 (CTLA4) and the Fc portion of immunoglobulin G (IgG), by the HD-Ad vector, improved transgene expression in rodents.

In addition, "serotype switching" can be used for repeated delivery of recombinant viruses. Gene therapy is initiated with one virus serotype, then switched to a second serotype for a subsequent administration, thereby avoiding attack by neutralizing antibodies specific to the first serotype.81,82 However, the level and duration of transgene expression following serotype switching may be limited by cross-reactive cytotoxic T lymphocytes that can also target cells infected by the second serotype virus.82,83 Thus, viral vectors expressing foreign antigens, such as the first generation Ad vectors, cannot be used. Finally, since all viral
gene therapy vectors used in the future should not express any viral coding proteins, in such case transient immune modulation may be sufficient for blocking the adaptive immune response. It was shown that cyclophosphamide alone or in combination with cyclosporine A extended transgene expression mediated by the first generation Ad vector. Our group has demonstrated that transient administration of cyclophosphamide allowed readministration of HD-Ad vectors with efficient expression of transgenes.

Efficient and safe delivery to airway

How to deliver genes safely and effectively into the airway cells is an area that is not extensively studied. In addition to the immune barriers discussed above, there are physical barriers to vector delivery. At the time when the early CF gene therapy clinical studies were conducted, there was not enough information regarding physical barriers to gene delivery, such as the mucosal layer, airway innate immune cells and physical protection by airway epithelial tight junctions. The mucosal layer is the first line of airway defense and it has been shown to inhibit viral transduction. For an inflamed lung, such as in CF, excessive secretion of mucus may be even more problematic for gene delivery. In addition, as mentioned above, the airway immune cells, mainly macrophages, can take and destroy a large portion of gene therapy vectors delivered, thus preventing them by reaching the target cells. More importantly, viral vectors used in CF lung gene therapy depend on their receptors to gain entry into their target cells which are airway epithelial cells. However, cellular receptors for viral vectors, such as the coxsackie-adenoviral receptor (CAR), are located not on the lumenal side, but on the basolateral side of the airway that is not directly accessible to vectors. Since airway epithelial cells are connected by tight junction proteins, vectors delivered cannot reach the basolateral side of the airway.

There are strategies that can be considered to overcome these barriers. For reducing the inhibition from the mucus layer, mucolytic reagents, such as nacystelyn, may be used. In addition, various polycations, such as DEAE-dextran, polylysines, protamine, and branched polyethyleneimine have been shown to greatly enhance viral vector delivery to mouse lungs. To reduce the loss of gene therapy vector, lung macrophages can be depleted by gadolinium chloride or liposome/dichloromethylene-bisphosphonate. However, these approaches may be too toxic to humans even though mice tolerate them well. On the other hand, anti-inflammatory reagents such as rooperol, methyl palmitate and mangiferin, that inhibit the phagocytic activity of macrophages, may be administered by aerosolization to the airway during vector delivery. In addition, dexamethasone has been shown to reduce the phagocytic activity of pulmonary macrophages although it is not effective on peritoneal macrophages.

Although the accessibility of receptors may not be an issue with nonviral vectors, such as liposomes, it is important for gene delivery with viral vectors. Studies with animals show that reagents, such as Ca\(^{2+}\)-chelator, EGTA, can be used to break the tight-junctions transiently for enhancing viral vector delivery to the lung. It has also been shown that L-\(\alpha\)-lysophosphatidylcholine (LPC) can enhance viral vector delivery dramatically to the lung of mice and rabbits although it is not clear whether LPC breaks tight-junctions.

Since gene delivery to lung normally causes some levels of innate immune responses, it is likely that less stress put on the host with less invasive delivery method will elicit weaker innate immune reactions and produce better results of therapeutic gene expression. The delivery method itself may also affect the outcome. For example, an optimized aerosol delivery approach may give a much better vector distribution in the lung than instillation.

Sustained therapeutic gene expression in the lung airway

Sustained therapeutic gene expression is important for lung gene therapy because it is unlikely that gene therapy vectors, no matter viral or nonviral, cannot be frequently administered to the lung due to the host innate and adaptive immune responses to viral vectors and bacterial DNA in nonviral vectors. The challenge has not been paid enough attention because most of the lung gene therapy studies have been focusing on achieving efficient therapeutic gene expression. The sustained therapeutic gene is more difficult to achieve in the lung than other organs, such as the eye because the lung is an immunologically sensitive organ and the airway cells turn over. For the loss of therapeutic gene expression due to the host immune responses, strategies to overcome this problem have to be considered from all aspects of the gene therapy design, such as vector choice, vector dose and delivery methods. These issues are covered in the section of immune barriers and will not be repeated here.

Airway epithelial cells turn over naturally. In mice the average half-life of the ciliated epithelial cells is about 6 months in the trachea and 17 months in the lung. In humans, the lung epithelial cells may have similar life spans, which may be reduced in diseased conditions. Two strategies may be used to cope with the problem. The first strategy is to allow vector re-administration with a long interval, for example once a year, if the adaptive immune responses to vectors can be avoided. This strategy cannot be used if the vector expressed foreign antigens, such as the leaky expression of viral genes seen in the early generation Ad vectors. However, it is expected that all future gene therapy vectors do not express vector encoded viral genes, thus transient immune suppression may be used to control the adaptive immune problem. In fact, it has been demonstrated in mice that HD-Ad vectors can be re-administered through transient immunosuppression. The second strategy is to deliver a vector that allows a therapeutic gene to be integrated safely in the airway progenitor cells. One major concern for integration of therapeutic genes to correct genetic diseases is the risk of random insertion that could cause major side effects including cancer development. Now this problem can be solved through engineering site-specific endonucleases to select a safe site for the integration. Three types of engineered endonucleases have been studied for their potential
in genetic engineering and therapeutic development. Zinc finger nucleases (ZFNs) are the first type in this group, engineered by combining the nonspecific nuclease domain of the Fok I restriction endonuclease with a zinc finger DNA-binding domain.\textsuperscript{104,105} ZFNs have been now be used successfully for genome editing and site specific gene insertion\textsuperscript{102,106} although engineering ZFNs for a new chromosome site is still time-consuming. Recently, a new class of engineered nucleases called transcription activator-like effector nucleases (TALENs), has emerged.\textsuperscript{107–110} Like ZFNs, TALENs use the same Fok I nuclease domains for DNA cleavage. Unlike ZFNs, TALENs can be easily engineered for any new integration site. More recently, a third type of endonucleases was designed based on the CRISPR/Cas9 (CRISPR-associated) system which is involved in genome defense mechanisms in bacteria for destroying foreign DNA.\textsuperscript{111–113} This CRISPR/Cas9 system is engineered to contain a single protein, Cas9 and a small RNA.\textsuperscript{114,115} Since a relatively short target sequence (only 13 out of 20 is required) is used for determination of the cleavage site,\textsuperscript{116} its off-target effects\textsuperscript{117–119} have to be extensively characterized before it is used for gene therapy. Since HD-Ad vectors are highly efficient in gene delivery to airway cells with a large DNA carrying capacity, a single vector can carry both the engineered site-specific nuclease genes and a therapeutic gene for clinical applications.

Reasons for the recent clinical success in eye gene therapy

There are several important factors that made the recent success in eye gene therapy possible. First of all, the right gene therapy vector was selected for the disease targeted. For targeting LCA2, AAV2 vector is the right choice because the therapeutic gene \textit{RPE65} is small enough for this type of vectors which are efficient in transducing retinal pigment epithelial cells.\textsuperscript{120} The cDNA of \textit{RPE65} is about 3.1 kb\textsuperscript{121} which allows AAV type vectors to have 1.6 kb room for DNA elements to control the therapeutic gene expression. For the LCA2 clinical studies, two types of DNA control elements are used. The trial study by Bainbridge et al used the human \textit{RPE65} gene promoter (1.4 kb) and the bovine growth hormone polyadenylation signal to control the \textit{RPE65} expression,\textsuperscript{2} whereas in studies by Maguire et al and Cideciyan et al, the expression of \textit{RPE65} is under the control of the chicken \(\beta\)-actin promoter together with the cytomegalovirus immediate early enhancer, the rabbit \(\beta\)-globin intro/exon junction and the SV40 polyadenylation signal. In addition to the carrying capacity good enough to meet the delivery of therapeutic gene, AAV2 is efficient in transducing retinal pigment epithelial (RPE) cells.\textsuperscript{120}

Secondly, immune barrier to AAV vector delivery is less of a problem for subretinal gene delivery because the anterior chamber of the eye is an immune-privileged site\textsuperscript{122} and AAV elicits weaker immune responses compared to other types of viral vectors.\textsuperscript{123} Immune privilege of the eye is established through five mechanisms, blood: ocular barriers, absence of lymphatic drainage pathways, soluble immunomodulatory factors in aqueous humor, immunomodulatory ligands on the surface of ocular parenchymal cells and indigenous, tolerance-promoting antigen-presenting cells.\textsuperscript{122} Since immune protection against pathogens can injure vital tissues in an innocent bystander manner, immune privilege is regarded as an evolutionary adaptation to enable local protection by immune effectors without disrupting the function of the vital tissues, such as light transduction by retinal cells.

Furthermore, RPE cells are ideal target cells for gene therapy. Essentially, all AAV vectors are efficient in transducing RPE cells although it is not totally clear whether this is due to the phagocytic activity of the RPE or the availability of the viral receptors on these cells.\textsuperscript{120} The structural position of the RPE cells allowing them to have more surface area for interacting with the vectors delivered to the subretinal space may be an advantage over other cells, such as photoreceptor cells. In addition to AAV, HD-Ad vectors can also transduce RPE in mice with a high efficiency.\textsuperscript{124} For gene therapy, it is important to have the target cells available when the vector is delivered. Although LCA2 is an early onset disease, retinal cells are well preserved in young patients,\textsuperscript{125} thus making functional rescue possible. An additional advantage for RPE cells being targets is that there is no cell turnover in the RPE layer so that the therapeutic gene expression can be sustained.

Finally, right animal models have been tested in pre-clinical studies of LCA2 gene therapy.\textsuperscript{125} For gene therapy targeting a particular disease, it is important to test the therapeutic approach in large animal models in addition to rodent models because there are major differences in organ anatomy, composition of cell types and gene function as well as tolerance to foreign substances. For example, mouse lungs do not have submucosal glands as in humans and have different epithelial compositions, such as with more Clara cells and less goblet cells in the large airways.\textsuperscript{126–130} When the mouse \textit{CFTR} gene is knocked out, there are no major lung disorders as in CF patients or other animal models for CF.\textsuperscript{131,132} Mice can tolerate endotoxin, LPS, at least 1000 time better than humans\textsuperscript{133,134} suggesting that there may be differences in tolerance to other substances such as gene therapy vectors. Before clinical tests in LCA2 patients, several research teams\textsuperscript{135–139} demonstrated proof of concept for gene replacement therapy in a canine model that is homozygous for a null mutation in \textit{RPE65} in addition to tests done in mice.\textsuperscript{140} These animal studies provided evidence for long term functional correction without apparent adverse effects to support clinical applications.

What can be learned from eye field to improve lung gene therapy

The trajectory of the eye gene therapy research sheds new light into several areas of the lung gene therapy development. First of all, we need to select the right vector for each particular lung disease. If a disease requires long term therapeutic gene expression in airway epithelial cells, such as in CF, a highly efficient vector with a large DNA carrying capacity and capability to confer long term transgene expression, should be considered to avoid frequent readministration. Frequent readministration of gene therapy vectors to lung is unlikely to be tolerated, especially for CF...
Figure 1  Schematic diagram showing viral entry and transport. This figure was reproduced from the review article by Brandenburg and Zhuang\textsuperscript{141} with permission from Nature Publishing Group. The diagram summarizes viral entry and travel in mammalian cells. Viruses attach to the plasma membrane, surf on the cell surface or along the filopodia (1–3), and bind to specific receptors before entering the cell. Viruses can directly fuse with the plasma membrane (2). They also hijack endocytic pathways, including clathrin-dependent (1), caveolin-dependent (3) or clathrin- and caveolin-independent (4) pathways for internalization. After
lungs that are under inflammatory conditions. It is likely that a viral vector, instead of nonviral vector, will have a better chance to be successful in clinical applications since current nonviral vectors cannot confer long term therapeutic gene expression. Viruses are professional gene carriers and have evolved molecular mechanisms to enter a cell and inject the DNA into cell nuclei efficiently. As shown in Fig. 1, recent studies using single-virus tracking techniques in live cells revealed many of the molecular mechanisms that different viruses use to gain entry into mammalian cells. All the advanced viral vectors do not have viral genes, but they maintain the ability to send their genomes efficiently into the nuclear compartment of target cells. In addition to the high efficiency in delivering genes to cells, the viral vector DNA is stable in transduced cells as long as host innate immune responses can be minimized. All these features of viral vectors are unmatched by nonviral vectors. Currently, the most hopeful viral vector for lung gene therapy, in our opinion, is the HD-Ad vector because its high efficiency in gene delivery has been demonstrated for transgene expression in mice, rabbits and pigs.

Second, safe and efficient delivery methods are critical for a clinical success. If the delivery method is too invasive or too stressful, it will likely enhance the host innate immune responses which will reduce the therapeutic efficacy. If the method is not efficient, the therapeutic benefits will be reduced. Over the past decade, our group has been developing safe and efficient methods for vector delivery to airways of large animals. Using the AeroProbe™ catheter and the LABneb™ control system designed by Trudell Medical International (London Ontario, Canada), we have worked out conditions for efficient gene delivery to rabbit airways. In these experiments the HD-Ad vector was formulated with 0.01% or 0.1% LPC in PBS and aerosolized to the rabbit airways through the AeroProbe™ catheter fitted into an endotracheal tube. More recently, we modified the method for efficient gene delivery to pig lungs. Since we use pigs about 30 kg in weight, we insert the AeroProbe™ catheter into a bronchoscope and aerosolize the HD-Ad vectors formulated with 0.01% LPC in PBS into pig lungs. As shown in Fig. 2, extensive human CFTR expression was observed in pig lungs one week post vector delivery. This method should be easily adapted for clinical studies since most instruments, such as the bronchoscope and ventilation machine as well as drugs for anesthesia are the same as used in humans.

Third, since lung is an immune sensitive organ, it is important to avoid the host immune responses. Even by selecting the best vector and most safe and effective delivery method, we will still face the problem of the host immune responses to viral vectors. Strategies to overcome the host immune responses are discussed early in the

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Figure 2  Expression of human CFTR protein in pig airway epithelial cells. An HD-Ad vector containing the human CFTR gene driven by the human cytokeratin 18 gene promoter was aerosolized to pig lungs. One week after delivery, lung tissues were taken and immunostaining was performed on tissue sections to visualize the human CFTR protein. Left panel shows the human CFTR protein located at the apical membrane of pig airway epithelial cells as the green immunofluorescence. The red immunofluorescence was from staining with an antibody against Zou-1 (an epithelial tight junction molecule) marking the cells as airway epithelial cells. Right panel, a section of pig lung without vector transduction was immunostained in the same way as a negative control. The blue fluorescence indicates the nuclei stain with DAPI. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
article. The most practical approaches to minimize the host immune responses are to reduce the vector dose as much as possible and to use transient immunosuppression to reduce the host immune responses.

Another critical consideration for improving lung gene therapy is to examine gene therapy vectors in appropriate preclinical animal models for the targeting disease. For gene therapy targeting CF lung disease, CF mouse models are not adequate enough for assessing the efficacy of gene therapy vectors and delivery methods, because of the major differences in lung biology between mice and humans as mentioned early. Recently, two excellent animal models (pig and ferret) for CF lung disease have been created by scientists at the University of Iowa. Unlike CF mice, knocking out the CFTR gene in pigs and ferrets results in typical CF lung symptoms, including mucus plugging and spontaneous lung infection. These animal models will be very useful for testing gene therapy vectors and delivery methods for treating CF.

Finally, sustained therapeutic gene expression is required for a clinical success in lung gene therapy. The most important way to maintain therapeutic gene expression is to protect the transduced cells from attack by host immune cells. This point has been discussed already. Although viral vectors are stable in lung cells, lung epithelial cell turnover does occur naturally, which will reduce the therapeutic gene expression. There are two strategies that can be considered to overcome this problem. One strategy is to re-administer viral vectors once a year. This requires transient immunosuppression to avoid host immune responses to vector readministration. In mice, this has been demonstrated. The second strategy is to integrate the therapeutic gene into progenitor cells of airway epithelium. This is a new, hot area of research and it will push gene therapy into a new stage for clinical applications.

Summary

Despite the early unsuccessful clinical trials of CF lung gene therapy, later decades of research has identified major challenges and made progress in overcoming these challenges. Analyzing the factors fundamental to the success in clinical studies of LCA2 gene therapy, we can learn a lot to improve lung gene therapy development. To address these challenges to lung gene therapy, we need to demonstrate that the host immune responses to viral vectors can be controlled effectively and sustained therapeutic gene expression can be maintained. The whole gene therapy field is still in its infancy. Even for eye gene therapy, there are still a lot of challenges, such as, efficient delivery of large genes to photoreceptor cells. With the molecular techniques under constant evolution, more and more successful gene therapy cases will emerge. It is matter of time that gene therapy will be a main approach to treating genetic diseases.

Conflict of interest

No authors have any conflict of interest to declare.

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References

1. Maguire AM, Simonelli F, Pierce EA, et al. Safety and efficacy of gene transfer for Leber’s congenital amaurosis. N Engl J Med. May 22 2008;358(21):2240–2248.
2. Bainbridge JW, Smith AJ, Barker SS, et al. Effect of gene therapy on visual function in Leber’s congenital amaurosis. N Engl J Med. May 22 2008;358(21):2231–2239.
3. Cideciyan AV, Aleman TS, Boye SL, et al. Human gene therapy for RPE65 isomerase deficiency activates the retinoid cycle of vision but with slow rod kinetics. Proc Natl Acad Sci U S A. Sep 30 2008;105(39):15112–15117.
4. Hanein S, Perrault I, Gerber S, et al. Leber congenital amaurosis: comprehensive survey of the genetic heterogeneity, refinement of the clinical definition, and genotype-phenotype correlations as a strategy for molecular diagnosis. Hum Mutat. Apr 2004;23(4):306–317.
5. Bennett J, Ashtari M, Wellman J, et al. AAV2 gene therapy readministration in three adults with congenital blindness. Sci Transl Med. Feb 8 2012;4(120):120ra115.
6. Maclaren RE, Groppe M, Barnard AR, et al. Retinal gene therapy in patients with choroideremia: initial findings from a phase 1/2 clinical trial. Lancet. Jan 15 2014;383(9923):1129–1137.
7. Bryant LM, Christopher DM, Giles AR, et al. Lessons learned from the clinical development and market authorization of Glybera. Hum Gene Ther Clin Dev. Jun 2013;24(2):55–64.
8. Yla-Herttuala S. Endgame: glybera finally recommended for approval as the first gene therapy drug in the European Union. Mol Ther. Oct 2012;20(10):1831–1832.
9. Miller N. Glybera and the future of gene therapy in the European Union. Nat Rev Drug Discov. May 2012;11(5):419.
10. Cao H, Molday RS, Hu J. Gene therapy: light is finally in the tunnel. Protein Cell. Dec 2011;2(12):973–989.
11. Griesenbach U, Alton EW. Moving forward: cystic fibrosis gene therapy. Hum Mol Genet. Oct 2013;22(11):R52–R58.
12. Prickett M, Jain M. Gene therapy in cystic fibrosis. Transl Res. Apr 2013;161(4):255–264.
13. Flotte TR, Ng P, Dylla DE, et al. Viral vector-mediated and cell-based therapies for treatment of cystic fibrosis. Mol Ther J Am Soc Gene Ther. 2007;15(2):229–241.
14. Rommens JM, Iannuzzi MC, Kerem B, et al. Identification of the cystic fibrosis gene: chromosome walking and jumping. Science. Sep 8 1989;245(4922):1059–1065.
15. Kerem B, Rommens JM, Buchanan JA, et al. Identification of the cystic fibrosis gene: genetic analysis. Science. Sep 8 1989;245(4922):1073–1080.
16. Rommens JM, Zengerling-Lentes S, Kerem B, Melmer G, Buchwald M, Tsui LC. Physical localization of two DNA markers closely linked to the cystic fibrosis locus by pulsed-field gel electrophoresis. Am J Hum Genet. Dec 1989;45(6):932–941.
17. Koehler DR, Hitt MM, Hu J. Challenges and strategies for cystic fibrosis lung gene therapy. Mol Ther J Am Soc Gene Ther. 2001;4(2):84–91.
18. Zabner J, Couture LA, Gregory RJ, Graham SM, Smith AE, Welsh MJ. Adenovirus-mediated gene transfer transiently corrects the chloride transport defect in nasal epithelia of patients with cystic fibrosis. Cell. Oct 22 1993;75(2):207–216.
19. Grubb BR, Pickles RJ, Ye H, et al. Inefficient gene transfer by adenovirus vector to cystic fibrosis airway epithelia of mice and humans. Nature. Oct 27 1994;371(6500):802–806.

20. Boucher RC, Knowles MR, Johnson LG, et al. Gene therapy for cystic fibrosis using E1-deleted adenovirus: a phase I trial in the nasal cavity. The University of North Carolina at Chapel Hill Hum Gene Ther. May 1994;5(5):615–639.

21. Knowles MR, Hohneker KW, Zhou Z, et al. A controlled study of adenoviral-vector-mediated gene transfer in the nasal epithelium of patients with cystic fibrosis. N Engl J Med. Sep 28 1995;333(13):823–831.

22. Crystal RG, McElvany NG, Rosenfeld MA, et al. Administration of an adenovirus containing the human CFTR cDNA to the respiratory tract of individuals with cystic fibrosis. Nat Genet. Sep 1994;8(1):42–51.

23. Bellon G, Michel-Calemard L, Thouvenot D, et al. Aerosol administration of a recombinant adenovirus expressing CFTR to cystic fibrosis patients: a phase I clinical trial. Hum Gene Ther. Jan 1 1997;8(1):15–25.

24. Harvey BG, Leopold PL, Hackett NR, et al. Airway epithelial CFTR mRNA expression in cystic fibrosis patients after repetitive administration of a recombinant adenovirus. J Clin Invest. Nov 1999;104(9):1245–1255.

25. Joseph PM, O’ Sullivan BP, Lapey A, et al. Aerosol and lobar administration of a recombinant adenovirus to individuals with cystic fibrosis. I. Methods, safety, and clinical implications. Hum Gene Ther. Jul 20 2001;12(11):1369–1382.

26. Perricone MA, Morris JE, Pavlica K, et al. Aerosol and lobar administration of a recombinant adenovirus to individuals with cystic fibrosis. II. Transfection efficiency in airway epithelium. Hum Gene Ther. Jul 20 2001;12(11):1383–1394.

27. Wagner JA, Messner AH, Moran ML, et al. Safety and biological efficacy of an aden-associated virus vector-cystic fibrosis transmembrane regulator (AAV-CFTR) in the cystic fibrosis maxillary sinus. Laryngoscope. Feb 1999;109(2 Pt 1):266–274.

28. Moss RB, Rodman D, Spencer LT, et al. Repeated adenovirus-associated virus serotype 2 aerosol-mediated cystic fibrosis transmembrane regulator gene transfer to the lungs of patients with cystic fibrosis: a multicenter, double-blind, placebo-controlled trial. Chest. Feb 2004;125(2):509–521.

29. Caplen NJ, Alton EW, Middleton PG, et al. Liposome-mediated CFTR gene transfer to the nasal epithelium of patients with cystic fibrosis. Nat Med. Jan 1995;1(1):39–46.

30. Gill DR, Southern KW, Mofford KA, et al. A placebo-controlled study of liposome-mediated gene transfer to the nasal epithelium of patients with cystic fibrosis. Gene Ther. Mar 1997;4(3):199–209.

31. Porteous DJ, Dorin JR, McLachlan G, et al. Evidence for safety and efficacy of DOTAP cationic liposome mediated CFTR gene transfer to the nasal epithelium of patients with cystic fibrosis. Gene Ther. Mar 1997;4(3):210–218.

32. Zabner J, Cheng SH, Meeker D, et al. Comparison of DNA-lipid complexes and DNA alone for gene transfer to cystic fibrosis airway epithelium in vivo. J Clin Invest. Sep 15 1997;100(6):1529–1537.

33. Noone PG, Hohneker KW, Zhou Z, et al. Safety and biological efficacy of a lipid-CFTR complex for gene transfer in the nasal epithelium of adult patients with cystic fibrosis. Mol Ther. Jan 2000;1(1):105–114.

34. Alton EW, Stern M, Farley R, et al. Cationic lipid-mediated CFTR gene transfer to the lungs and nose of patients with cystic fibrosis: a double-blind placebo-controlled trial. Lancet. Mar 20 1999;353(9157):947–954.

35. Ruiz FE, Clancy JP, Perricone MA, et al. A clinical inflammatory syndrome attributable to aerosolized lipid-DNA administration in cystic fibrosis. Hum Gene Ther. May 1 2001;12(7):751–761.

36. Cao H, Koehler DR, Hu J. Adenoviral vectors for gene replacement therapy. Viral Immunol. 2004;17(3):327–333.

37. Kojaghlalian T, Flomenberg P, Horwitz MS. The impact of adenovirus infection on the immunocompromised host. Rev Med Virol. May-Jun 2003;13(3):155–171.

38. Moss RB, Milla C, Colombo J, et al. Repeated aerosolized AAV-CFTR for treatment of cystic fibrosis: a randomized placebo-controlled phase 2B trial. Hum Gene Ther. Aug 2007;18(8):726–732.

39. Wagner JA, Nepomuceno IB, Messner AH, et al. A phase II, double-blind, randomized, placebo-controlled clinical trial of IgAVCF using maxillary sinus delivery in patients with cystic fibrosis with antistomies. Hum Gene Ther. Jul 20 2002;13(11):1349–1359.

40. Berns KI, Giraud C. Biology of aden-associated virus. Curr Top Microbiol Immunol. 1996;218:1–23.

41. Gao X, Huang L. Cationic liposome-mediated gene transfer. Gene Ther. Dec 1995;2(10):710–722.

42. Shimizu K, Sakurai F, Machiti M, Katayama K, Mizuguchi H. Quantitative analysis of the leaky expression of adenovirus genes in cells transduced with a replication-incompetent adenovirus vector. Mol Pharm. Aug 1 2011;8(4):1430–1435.

43. Wivel NA, Gao P-G, Wilson JM. Adenovirus vectors. In: The Development of Human Gene Therapy. San Diego, CA: Cold Spring Harbor Laboratory Press; 1999.

44. Parks RJ. Improvements in adenoviral vector technology: overcoming barriers for gene therapy. Clin Genet. 2000;58(1):1–11.

45. Seiler MP, Cerullo V, Lee B. Immune response to helper dependent adenoviral mediated liver gene therapy: challenges and prospects. Curr Gene Ther. Oct 2007;7(5):297–305.

46. Kim IH, Jozkowicz A, Piedra PA, Oka K, Chan L. Lifetime correction of genetic deficiency in mice with a single injection of helper-dependent adenoviral vector. Proc Natl Acad Sci U S A. 2001;98(23):13282–13287.

47. Toietta G, Koehler DR, Finegold MJ, Lee B, Hu J, Beaudet AL. Reduced inflammation and improved airway expression using helper-dependent adenoviral vectors with a K18 promoter. Mol Ther J Am Soc Gene Ther. 2003;7(5 Pt 1):649–658.

48. Koehler DR, Saajan U, Chow YH, et al. Protection of Cftp knockout mice from acute lung infection by a helper-dependent adenoviral vector expressing Cftp in airway epithelia. Proc Natl Acad Sci U S A. 2003;100(26):15364–15369.

49. Koehler DR, Frndova H, Leung K, et al. Aerosol delivery of an enhanced helper-dependent adenovirus formulation to rabbit lung using an intratracheal catheter. J Gene Med. 2005;7(11):1409–1420.

50. Cao H, Machuca TN, Yeung JC, et al. Efficient gene delivery to pig airway epithelium and submucosal glands using helper-dependent adenoviral vectors. Mol Ther Nucleic Acids. 2013;2:e127.

51. Aflone SA, Conrad CK, Kearns WG, et al. In vivo model of aden-associated virus vector persistence and rescue. J Virol. May 1996;70(5):3235–3241.

52. Conrad CK, Allen SS, Aflone SA, et al. Safety of single-dose administration of an aden-associated virus (AAV)-CFTR vector in the primate lung. Gene Ther. Aug 1996;3(8):658–668.

53. Allocca M, Doria M, Petrello M, et al. Serotype-dependent packaging of large genes in aden-associated viral vectors results in effective gene delivery in mice. J Clin Invest. 2008;118(5):1955–1964.

54. Lai Y, Yue Y, Duan D. Evidence for the failure of aden-associated virus serotype 5 to package a viral genome < or = 8.2 kb. Mol Ther. Jan 2010;18(1):75–79.

55. Gruntman AM, Mueller C, Flotte TR, Gao G. Gene transfer in the lung using recombinant aden-associated virus. Curr Protoc Microbiol. Aug 2012 Chapter 14:Unit14D 12.

56. Duan D, Yue Y, Engelhardt JF. Expanding AAV packaging capacity with trans-splicing or overlapping vectors: a quantitative comparison. Mol Ther. Oct 2001;4(4):383–391.
57. Yan Z, Zhang Y, Duan D, Engelhardt JF. Trans-splicing vectors expand the utility of adeno-associated virus for gene therapy. Proc Natl Acad Sci U S A. Jun 6 2000;97(12):6716–6721.

58. White AF, Mazur M, Sorschek EJ, Zinn KR, Ponnazhagan S. Genetic modification of adeno-associated viral vector type 2 capsid enhances gene transfer efficiency in polarized human airway epithelial cells. Hum Gene Ther. Dec 2008;19(12):1407–1414.

59. Li W, Zhang L, Johnson JS, et al. Generation of novel AAV variants by directed evolution for improved CFTR delivery to human ciliated airway epithelium. Mol Ther. Dec 2009;17(12):2067–2077.

60. Pozzi D, Marchini C, Cardarelli F, et al. Mechanistic evaluation of the transfection barriers involved in lipid-mediated gene delivery: interplay between nanostructure and composition. Biochim Biophys Acta. Mar 2014;1838(3):957–967.

61. Bienenstock J. The lung as an immunologic organ. Annu Rev Med. 1984;35:49–62.

62. Dai Y, Schwarz EM, Gu D, Zhang WW, Sarvetnick N, Verma IM. Cellular and humoral immune responses to adenoviral vectors containing factor IX gene: tolerization of factor IX and vector antigens allows for long-term expression. Proc Natl Acad Sci U S A. Feb 28 1995;92(5):1401–1405.

63. Yang Y, Li Q, Ertl HC, Wilson JM. Cellular and humoral immune responses to viral antigens create barriers to lung-directed gene therapy with recombinant adenoviruses. J Virol. Apr 1995;69(4):2004–2015.

64. Kay MA, Holterman AX, Meuse L, et al. Long-term hepatic adenovirus-mediated gene expression in mice following CTLA4ig administration. Nat Genet. Oct 1995;11(2):191–197.

65. Yang Y, Nunes FA, Berenci K, Furth EE, Gonczol E, Wilson JM. Cellular immunity to viral antigens limits E1-deleted adenoviruses for gene therapy. Proc Natl Acad Sci U S A. 1994;91(10):4407–4411.

66. Raper SE, Chirmule N, Lee FS, et al. Fatal systemic inflammatory response syndrome in a orthotopic transcarbamylase deficient patient following adenoviral gene transfer. Mol Genet Metab. Sep-Oct 2003;80(1–2):148–158.

67. Lee BH, Kushwah R, Wu J, et al. Adenoviral vectors stimulate innate immune responses in macrophages through cross-talk with epithelial cells. Immunol Lett. Nov 30 2010;134(1):93–102.

68. Sinn PL, Burnight ER, McCray Jr PB. Progress and prospects: prospects of repeated pulmonary administration of viral vectors. Gene Ther. Sep 2009;16(9):1059–1065.

69. Zsengeller Z, Otake K, Hossain SA, Berclaz PY, Trapnell BC. Internalization of adenovirus by alveolar macrophages initiates early proinflammatory signaling during acute respiratory tract infection. J Virol. Oct 2000;74(20):9635–9667.

70. Baldwin Jr AS. The NF-kappa B and I kappa B proteins: new discoveries and insights. Annu Rev Immunol. 1996;14:649–683.

71. Cao HB, Wang A, Martin B, et al. Down-regulation of IL-8 expression in human airway epithelial cells through helper-dependent adenoviral-mediated RNA interference. Cell Res. 2005;15(2):111–119.

72. Matsushima K, Morishita K, Yoshimura T, et al. Molecular cloning of a human monocyte-derived neutrophil chemotactic factor (MDNCF) and the induction of MDNCF mRNA by interleukin 1 and tumor necrosis factor. J Exp Med. Jun 1 1988;167(6):1883–1893.

73. Sun L, Wu J, Du F, Chen X, Chen ZJ. Cyclic GMP-AMP synthase is a cytosolic DNA sensor that activates the type I interferon pathway. Science. Feb 15 2013;339(6121):786–791.

74. Mays LF, Wilson JM. The complex and evolving story of T cell activation to AAV vector-encoded transgene products. Mol Ther. Jan 2011;19(1):16–27.

75. Fearon DT, Locksley RM. The instructive role of innate immunity in the acquired immune response. Science. Apr 5 1996;272(5258):50–53.

76. Scaria A, St George JA, Gregory RJ, et al. Antibody to CD40 ligand inhibits both humoral and cellular immune responses to adenoviral vectors and facilitates repeated administration to mouse airway. Gene Ther. Jun 1997;4(6):611–617.

77. Wilson CB, Embree LJ, Schwalter D, et al. Transient inhibition of CD28 and CD40 ligand interactions prolongs adenovirus-mediated transgene expression in the lung and facilitates expression after secondary vector administration. J Virol. Sep 1998;72(9):7542–7550.

78. Liang Z, Feingold E, Kochanek S, Clemens PR. Systemic delivery of a high-capacity adenovector expressing mouse CTLA4ig improves skeletal muscle gene therapy. Mol Ther. Sep 2002;6(3):369–376.

79. Yamashita K, Masunaga T, Yanagida N, et al. Long-term acceptance of rat cardiac allografts on the basis of adenovirus mediated CD40ig plus CTLA4ig gene therapies. Transplantation. Oct 15 2003;76(7):1089–1096.

80. Mastrangeli A, Harvey BG, Yao J, et al. “Sero-switch” adenovirus-mediated in vivo gene transfer: circumvention of anti-adenovirus humoral immune defenses against repeat adenovirus vector administration by changing the adenovirus serotype. Hum Gene Ther. Jan 1996;7(1):79–87.

81. Sumner-Jones SG, Gill DR, Hyde SC. Lack of repeat transduction by recombinant adeno-associated virus type 5/5 vectors in the mouse airway. J Virol. Nov 2007;81(22):12360–12367.

82. Mack CA, Song WR, Carpenter H, et al. Circumvention of anti-adenovirus neutralizing immunity by administration of an adenoviral vector of an alternate serotype. Hum Gene Ther. Jan 1 1997;8(1):99–109.

83. Kaplan JM, Pennington SE, St George JA, et al. Potentiation of gene transfer to the mouse lung by complexes of adenovirus vector and polycations improves therapeutic potential. Hum Gene Ther. Jul 1 1998;9(10):1469–1479.

84. Cao H, Yang T, Li XF, et al. Readministration of helper-dependent adenoviral vectors to mouse airway mediated via transient immunosuppression. Gene Ther. Feb 2011;18(2):173–181.

85. Pickles RJ, Fahrner JA, Petrella JM, Boucher RC, Bergelson J. Retargeting the coxsackievirus and adenovirus receptor to the apical surface of polarized epithelial cells reveals the glycolaxyl as a barrier to adenovirus-mediated gene transfer. J Virol. Jul 2000;74(13):6050–6057.

86. Arcasoy SM, Latoche J, Gondor M, et al. MUC1 and other sialoglycoconjugates inhibit adenovirus-mediated gene transfer to epithelial cells. Am J Respir Cell Mol Biol. Oct 1997;17(4):422–435.

87. Bals R, Xiao W, Sang N, Weiner DJ, Meegalla RL, Wilson JM. Transduction of well-differentiated airway epithelium by recombinant adeno-associated virus is limited by vector entry. J Virol. Jul 1999;73(7):6085–6088.

88. Worgall S, Leopold PL, Wolff G, Ferris B, Van Roijen N, Crystal RG. Role of alveolar macrophages in rapid elimination of immune response to inhaled antigen by alveolar macrophages. Am J Respir Cell Mol Biol. Oct 1997;17(4):422–435.

89. Singh B, de la Concha-Bermejillo A. Gadolinium chloride and the degree of ovine lentivirus-induced lymphoid interstitial pneumonia. Int J Exp Pathol. Jun 1998;79(3):151–162.

90. Thepen T, McMenamin C, Oliver J, Kraal G, Holt PG. Regulation of immune response to inhaled antigen by alveolar...
macrophages: differential effects of in vivo alveolar macrophage elimination on the induction of tolerance vs. immunity. Eur J Immunol. Nov 1991;21(11):2845–2850.

92. Guzdak A, Turyna B, Allison AC, Sladek K, Koj A, Roopor, an inhibitor of cytokine synthesis, decreases the respiratory burst in human and rat leukocytes and macrophages. Mediators Inflamm. 1997;6(1):53–57.

93. Cal P, Kaphalia BS, Ansari GA. Methyl palmitate: inhibitor of phagocytosis in primary rat Kuffer cells. Toxicology. Jun 1 2005;201(2–3):197–204.

94. Garcia D, Delgado R, Ubeira FM, Leiro J. Modulation of a macrophage function by the Mangifera indica L. extracts Vimang and mangiferin. Int Immunopharmacol. May 2002;2(6):797–806.

95. Nakamura Y, Murai T, Ogawa Y. Effect of in vitro and in vivo administration of dexamethasone on rat macrophage functions: comparison between alveolar and peritoneal macrophages. Eur Respir J. Feb 1996;9(2):301–306.

96. Chu Q, St George JA, Lukason M, Cheng SH, Scheule RK, Nakamura Y, Murai T, Ogawa Y. Effect of in vitro and in vivo administration of dexamethasone on rat macrophage functions: comparison between alveolar and peritoneal macrophages. Eur Respir J. Feb 1996;9(2):301–306.

97. Limberis M, Anson DS, Fuller M, Parsons DW. Recovery of airway cystic fibrosis transmembrane conductance regulator gene function in mice with cystic fibrosis after single-dose lentivirus-mediated gene transfer. Hum Gene Ther. Nov 1 2002;13(16):1961–1970.

98. Rawlin LS, Hogan BL. Ciliated epithelial cell lifespan in the mouse trachea and lung. Am J Physiol Lung Cell Mol Physiol. Jul 2008;295(1):L231–L234.

99. Hacein-Bey-Abina S, Von Kalle C, Schmidt M, et al. LMO2-associated clonal T cell proliferation in two patients after gene therapy for SCID-X1. Science. Oct 17 2003;302(5644):415–419.

100. Hacein-Bey-Abina S, Hauer J, Lim A, et al. Efficacy of gene therapy for X-linked severe combined immunodeficiency. N Engl J Med. Jul 22 2010;363(4):355–364.

101. Xu L, Zhao P, Mariano A, Han R. Targeted myostatin gene replacement in eukaryotes. Curr Opin Genet Dev. Feb 1 2012;22(1):306–310.

102. Xu L, Zhao P, Mariano A, Han R. Targeted myostatin gene replacement in eukaryotes. Curr Opin Genet Dev. Feb 1 2012;22(1):306–310.

103. Joung JK, Sander JD. TALENs: a widely applicable technology for targeted genome editing. Nat Rev Mol Cell Biol. Jul 2011;12(7):435–445.

104. Urnov FD, Rebar EJ, Holmes MC, Zhang HS, Gregory PD. Nucleic Acids Res. Oct 11 2013;41(20):8204–8212.

105. Guzdek A, Turyna B, Allison AC, Sladek K, Koj A, Roopor, an inhibitor of cytokine synthesis, decreases the respiratory burst in human and rat leukocytes and macrophages. Mediators Inflamm. 1997;6(1):53–57.

106. Cal P, Kaphalia BS, Ansari GA. Methyl palmitate: inhibitor of phagocytosis in primary rat Kuffer cells. Toxicology. Jun 1 2005;201(2–3):197–204.

107. Garcia D, Delgado R, Ubeira FM, Leiro J. Modulation of a macrophage function by the Mangifera indica L. extracts Vimang and mangiferin. Int Immunopharmacol. May 2002;2(6):797–806.

108. Chu Q, St George JA, Lukason M, Cheng SH, Scheule RK, Nakamura Y, Murai T, Ogawa Y. Effect of in vitro and in vivo administration of dexamethasone on rat macrophage functions: comparison between alveolar and peritoneal macrophages. Eur Respir J. Feb 1996;9(2):301–306.

109. Hockemeyer D, Wang H, Kiani S, et al. Genetic engineering of human pluripotent cells using TALE nucleases. Nat Biotechnol. Aug 2011;29(8):731–734.

110. Cermak T, Doyle EL, Christian M, et al. Efficient design and assembly of custom TALEN and other TAL effector-based constructs for DNA targeting. Nucleic Acids Res. Jul 2011;39(12):e82.

111. Bhaya D, Davison M, Barrangou R. CRISPR-Cas systems in bacteria and archaea: versatile small RNAs for adaptive defense and regulation. Annu Rev Genet. 2011;45:273–297.

112. Terns MP, Terns RM. CRISPR-based adaptive immune systems. Curr Opin Microbiol. Jun 2011;14(3):321–327.

113. Wiedenheft B, Sternberg SH, Doudna JA. RNA-guided genetic silencing systems in bacteria and archaea. Nature. Feb 16 2012;482(7385):331–336.

114. Jinek M, Chylinski K, Fonfara I, Hauer M, Doudna JA, Charpentier E. A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity. Science. Aug 17 2012;337(6096):816–821.

115. Deltcheva E, Chylinski K, Sharma CM, et al. CRISPR RNA maturation by trans-encoded small RNA and host factor Rnase III. Nature. Mar 31 2011;471(7340):602–607.

116. Jinek M, East A, Cheng A, Lin S, Ma E, Doudna J. RNA-programmed genome editing in human cells. Elife. 2013;2:e00471.

117. Yang H, Wang H, Shivalila CS, Cheng AW, Shi L, Jaenisch R. One-step generation of mice carrying reporter and conditional alleles by CRISPR/Cas-mediated genome engineering. Cell. Aug 27 2013;154(4):1370–1379.

118. Yoon RS, Tu M, Chatterjee S, et al. CRISPR RNA targeting beta-globin and CCR5 genes have substantial off-target activity. Nucleic Acids Res. Aug 11 2013;41(20):9584–9592.

119. Hsu PD, Scott DA, Weinstein JA, et al. DNA targeting specificity of RNA-guided Cas9 nucleases. Nat Biotechnol. Jul 21 2013;31(9):827–832.

120. Vandenbergh LE, Auricchio A. Novel adeno-associated viral vectors for retinal gene therapy. Gene Ther. Feb 2012;19(2):162–168.

121. Hamel CP, Tsilou E, Pfeffer BA, Hooks JJ, Detrick B, Redmond TM. Molecular cloning and expression of RPE65, a novel retinal pigment epithelium-specific microsomal protein that is post-transcriptionally regulated in vitro. J Biol Chem. Jul 25 1993;268(21):15751–15757.

122. Streilein JW. Ocular immune privilege: the eye takes a dim but practical view of immunity and inflammation. J Leukoc Biol. Aug 2003;74(2):179–185.

123. Petsis-Silva H, Linden R. Advances in gene therapy technologies to treat retinitis pigmentosa. Clin Ophthalmol. 2014;8:127–136.

124. Wu L, Lam S, Cao H, et al. Subretinal gene delivery using helper-dependent adenoviral vectors. Cell Biol. 2011;11(1):15.

125. Buch PK, Bainbridge JW, Ali RR. AAV-mediated gene therapy for retinal disorders: from mouse to man. Gene Ther. Jun 2008;15(11):849–857.

126. Wang X, Zhang Y, Ambersen A, Engelhardt JF. New models of the tracheal airway define the glandular contribution to airway surface fluid and electrolyte composition. Am J Respir Cell Mol Biol. Feb 2001;24(2):195–202.

127. Pack RJ, Al-Ugaily LH, Morris G, Widdicombe JG. The distribution and structure of cells in the tracheal epithelium of the mouse. Cell Mol Biol. Aug 1 1983;128(2 Pt 2):S41–S52.

128. Moller GG, Mariassy AT, Lollini LO. Structure as revealed by single-pair of TALE nucleases. Mol Ther Nucleic Acids. 2013;2:e112.

129. Jinek M, East A, Cheng A, Lin S, Ma E, Doudna J. RNA-programmed genome editing in human cells. Elife. 2013;2:e00471.

130. Yang H, Wang H, Shivalila CS, Cheng AW, Shi L, Jaenisch R. One-step generation of mice carrying reporter and conditional alleles by CRISPR/Cas-mediated genome engineering. Cell. Aug 27 2013;154(4):1370–1379.

131. Sun X, Yan Z, Yi Y, et al. Adeno-associated virus-targeted disruption of the CFTR gene in cloned ferrets. J Clin Invest. Apr 2008;118(4):1578–1583.
132. Rogers CS, Stoltz DA, Meyerholz DK, et al. Disruption of the CFTR gene produces a model of cystic fibrosis in newborn pigs. *Science*. Sep 26 2008;321(5897):1837–1841.

133. Sauter C, Wolfensberger C. Interferon in human serum after injection of endotoxin. *Lancet*. Oct 18 1980;2(8199):852–853.

134. Taveira da Silva AM, Kaulbach HC, Chuidian FS, Lambert DR, Suffredini AF, Danner RL. Brief report: shock and multiple-organ dysfunction after self-administration of *Salmonella* endotoxin. *N Engl J Med*. May 20 1993;328(20):1457–1460.

135. Acland GM, Aguirre GD, Bennett J, et al. Long-term restoration of rod and cone vision by single dose rAAV-mediated gene transfer to the retina in a canine model of childhood blindness. *Mol Ther*. Dec 2005;12(6):1072–1082.

136. Acland GM, Aguirre GD, Ray J, et al. Gene therapy restores vision in a canine model of childhood blindness. *Nat Genet*. May 2001;28(1):92–95.

137. Jacobson SG, Acland GM, Aguirre GD, et al. Safety of recombinant adeno-associated virus type 2-RPE65 vector delivered by ocular subretinal injection. *Mol Ther*. Jun 2006;13(6):1074–1084.

138. Narfstrom K, Katz ML, Bragadottir R, et al. Functional and structural recovery of the retina after gene therapy in the RPE65 null mutation dog. *Invest Ophthalmol Vis Sci*. Apr 2003;44(4):1663–1672.

139. Narfstrom K, Katz ML, Ford M, Redmond TM, Rakoczy E, Bragadottir R. In vivo gene therapy in young and adult RPE65−/− dogs produces long-term visual improvement. *J Hered*. Jan-Feb 2003;94(1):31–37.

140. Dejneka NS, Surace EM, Aleman TS, et al. In utero gene therapy rescues vision in a murine model of congenital blindness. *Mol Ther*. Feb 2004;9(2):182–188.

141. Brandenburg B, Zhuang X. Virus trafficking — learning from single-virus tracking. *Nat Rev Microbiol*. Mar 2007;5(3):197–208.

142. Pelkmans L, Puntener D, Helenius A. Local actin polymerization and dynamin recruitment in SV40-induced internalization of caveolae. *Science*. Apr 19 2002;296(5567):535–539.