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## Therapeutic Genome Editing: Prospects and Challenges

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### Abstract

Recent advances in the development of genome editing technologies based on programmable nucleases have significantly improved our ability to make precise changes in the genomes of eukaryotic cells. Genome editing is already broadening our ability to elucidate the contribution of genetics to disease by facilitating the creation of more accurate cellular and animal models of pathological processes. A particularly tantalizing application of programmable nucleases is the potential to directly correct genetic mutations in affected tissues and cells to treat diseases that are refractory to traditional therapies. Here we discuss current progress towards developing programmable nuclease-based therapies as well as future prospects and challenges.

Of the approximately 25,000 annotated genes in the human genome, mutations in over 3,000 genes have already been linked to disease phenotypes ([www.omim.org/statistics/geneMap](http://www.omim.org/statistics/geneMap)), and more disease-relevant genetic variations are being uncovered at a staggeringly rapid pace. Now, due to sharp drops in sequencing cost, the completion of the human genome project, and the exponential growth of human genome sequencing data from diseased individuals, the role of genetics in human health has become a major focus of research, clinical medicine and the development of targeted therapeutics<sup>1</sup>. These advances in our understanding of the genetic basis of disease have improved our understanding of disease mechanisms and pointed toward potential therapeutic strategies. However, despite valid therapeutic hypotheses and strong efforts in drug development, there have only been a limited number of successes using small molecules to treat diseases with strong genetic contributions<sup>2</sup>. Emerging therapeutic strategies that are able to modify nucleic acids within disease-affected cells and tissues have potential for treatment of monogenic, highly penetrant diseases, such as severe-combined immunodeficiency (SCID), haemophilia, and certain enzyme deficiencies due to their well-defined genetics and often lack of safe, effective alternative treatments.

Two of the most powerful genetic therapeutic technologies developed thus far are gene therapy, which enables restoration of missing gene function by viral transgene expression, and RNA interference (RNAi), which mediates targeted repression of defective genes by knockdown of the target mRNA (reviewed in <sup>3,4</sup>). Gene therapy has been used to successfully treat monogenic recessive disorders affecting the hematopoietic system, such as

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SCID and Wiskott-Aldrich syndrome, by semi-randomly integrating functional genes into the genome of hematopoietic stem/progenitor cells<sup>5-7</sup>. RNAi has been used to repress the function of genes implicated in cancer, age related macular degeneration and transthyretin (TTR)-amyloidosis among others, resulting in a therapeutic effect in clinical trials ([www.clinicaltrials.gov](http://www.clinicaltrials.gov), trial numbers: NCT00689065, NCT01961921 and NCT00259753). Despite promise and recent success, gene therapy and RNAi have limitations that prevent their utility for a large number of diseases. For example, viral gene therapy may cause mutagenesis at the insertion site and result in dysregulated transgene expression<sup>6</sup>. Alternatively, RNAi use is limited to targets where gene knockdown is beneficial. Also, RNAi often cannot fully repress gene expression, and is therefore unlikely to provide a benefit for diseases where complete ablation of gene function is necessary for therapy. RNAi may also have poor specificity, posing potential safety concerns and sometimes decreasing the effectiveness of treatment<sup>8-10</sup>.

Genome editing technologies based on programmable nucleases such as meganucleases (reviewed in<sup>11</sup>), zinc finger nucleases (reviewed in<sup>12</sup>), transcription activator-like effector nucleases (reviewed in<sup>13,14</sup>), and clustered regularly interspaced short palindromic repeat (CRISPR)-associated nuclease Cas9 (reviewed in<sup>15</sup>) are opening the possibility of achieving therapeutic genome editing in diseased cells and tissues, resulting in the removal or correction of deleterious mutations or the insertion of protective mutations.

In this Review, we will describe the different nuclease-based genome editing technologies, the mechanisms by which they produce genetic changes, considerations for their uses in therapeutic settings and major challenges that will need to be addressed to realize their clinical translation. Although a large number of genome editing therapeutic efforts have focused on treatment of monogenic, highly-penetrant disorders, we will also discuss intriguing treatment strategies to apply this class of therapy to diseases such as viral infections and cancer.

## Genome Editing Technologies

Programmable nucleases enable precise genome editing by introducing DNA double strand breaks (DSBs) at specific genomic loci. DSBs subsequently recruit endogenous repair machinery for either non-homologous end-joining (NHEJ) or homology directed repair (HDR) to the DSB site to mediate genome editing.

To date, four major classes of nucleases, meganucleases and their derivatives<sup>16-19</sup>, zinc finger nucleases (ZFNs)<sup>20-28</sup>, transcription activator like effector nucleases (TALENs)<sup>29-34</sup>, and CRISPR-associated nuclease Cas9<sup>35-43</sup> have been developed to enable site-specific genome editing (Table 1). These nuclease systems can be broadly classified into two categories based on their mode of DNA recognition - ZFN, TALEN, and meganucleases achieve specific DNA binding via protein-DNA interactions, Cas9 is targeted to specific DNA sequences by a short RNA guide molecule that base-pairs directly with the target DNA, protein-DNA interactions also have a role in its targeting. Meganucleases are endonucleases with large (>14bp) recognition sites, the DNA binding domains of which are also responsible for cleavage of target sequences<sup>18</sup>.

ZFNs and TALENs are chimeric enzymes consisting of a DNA binding domain fused to the sequence agnostic nuclease domain, *FokI*<sup>20,31</sup>. Re-targeting of ZFNs and meganucleases requires protein engineering, whereas re-targeting of TALENs requires complex molecular cloning<sup>18,44,45</sup>. In contrast, the Cas9 protein is invariant and can be easily retargeted to new DNA sequences by changing a small portion of the sequence of an accompanying RNA guide that base-pairs directly with target DNA. A potential advantage of Cas9 is its ability to introduce multiple DSBs in the same cell (also referred to as multiplexing) via expression of distinct guide RNAs. All four types of nucleases have been demonstrated to achieve efficient genome editing in a wide range of model organisms and mammalian cells and efforts are now underway in both industry and academia to develop these tools as therapeutics<sup>46-49</sup>.

Once the DSB has been made, the lesion may be repaired by either NHEJ or HDR depending on the cell state and the presence of a repair template (**Fig. 1**). NHEJ may repair the lesion by directly rejoining the two DSB ends in a process that does not require a repair template. Although NHEJ-mediated DSB repair can be accurate, repeated repair of the same DSB by NHEJ machinery eventually results in the formation of small insertion or deletion mutations (indels) bridging the break site. Indels introduced into the coding sequence of a gene can cause frameshift mutations that lead to mRNA degradation by nonsense-mediated decay, or result in the production of nonfunctional truncated proteins<sup>50</sup>. Thus, NHEJ may be used to suppress gene function similar to RNAi, however, it may lead to permanent inactivation by introducing loss of function mutations to the gene in targeted cells.

In comparison, HDR allows researchers to use an exogenous DNA template to specify the outcome of the DSB repair<sup>22,51-56</sup>. Upon introduction of a targeted DSB, HDR machinery may use exogenously provided single or double stranded DNA templates with sequence homology to the break site to synthesize DNA that is used to repair the lesion, incorporating any changes encoded in the template DNA. For example, HDR may be used along with an appropriately designed repair template to replace a mutated gene directly, thereby restoring gene function while preserving physiological regulation of gene expression.

## Therapeutic Genome Editing Strategies

Genome editing based therapy can be achieved through a number of approaches including correction or inactivation of deleterious mutations, introduction of protective mutations, addition of therapeutic transgenes, or disruption of viral DNA.

Pathogenic mutations can be broadly classified as gain- or loss-of-function. A gain-of-function mutation, such as in the cases of the *HTT* gene for huntington disease (<http://omim.org/entry/143100>) and *FGFR3* for Achondroplasia (<http://omim.org/entry/100800>), which result in the expression of a dominant negative mutant gene, may be corrected by using NHEJ-mediated induced mutations to specifically inactivate the mutant gene while leaving the wildtype copy intact on the other allele (**Fig. 1a**). Additionally, targeting of two nucleases to make two DSBs around and insertion may be useful for treatment of nucleotide expansion disorders such as spino-cerebellar ataxia (<http://www.omim.org/entry/164400>),

huntington disease, and fredrieck ataxia (<http://omim.org/entry/229300>) (**Fig. 1B**). A combination of DSBs may also be used to edit multiple loci to achieve a therapeutic effect.

However, some gain-of-function mutations, such as the *SOD1* G93A mutation in Amyotrophic Lateral Sclerosis (ALS) (<http://omim.org/entry/147450>), are point mutations, which might not be sufficiently different from the functioning allele on the homologous chromosome to be distinguished by the current generation of programmable nucleases, potentially leading to an undesirable complete loss of function of the protein if it is corrected using NHEJ. In this case HDR could be used to change the gain-of-function allele to the wildtype sequence, recovering gene function and eliminating pathogenic activity while preserving physiological levels of gene expression (Figure 1C). Similarly, loss-of-function mutations, including those found in Tay-Sachs disease (<http://omim.org/entry/272800>) would necessitate precise sequence changes to eliminate pathogenicity, requiring HDR gene correction to revert the loss-of-function mutation to the wildtype sequence. This same logic can also be extended to mutations which protect against infectious or genetic disease which may be loss-of-function in the case of CCR5 for HIV<sup>46,57</sup> (Figure 1a) or PCSK9 for hypercholesterolemia<sup>58</sup> which require inactivation by NHEJ, or gain-of-function in the case of *APP* (p.A673T) in alzheimers disease<sup>59</sup> which would require correction by HDR.

For deleterious loss of function mutations and protective gain of function mutations, a therapeutic effect may also be achieved by introducing a copy of the wildtype gene or gain-of-function mutant respectively (Figure 1D). The therapeutic transgene may be inserted into a new locus, including identified ‘safe harbor’ loci - a region of the genome which does not lead to discernable phenotypic effects when disrupted - to recover missing gene function<sup>60-62</sup>. Gene insertion may also be used to stably confer cells novel functions that protect against disease, such as the insertion of chimeric-antigen receptors (CAR) into T-cells to target certain leukemias<sup>63</sup>. Such gene insertion strategies are similar to viral-mediated gene therapy, but with the advantage of providing better control over transgene copy number and expression levels, which may be important for gene targets whose function is sensitive to expression levels.

Programmable nucleases may also be targeted to foreign DNA, such as viral genomes, which may be integrated proviruses or maintained extrachromosomally<sup>64-68</sup>. Targeting of extrachromosomal DNA may lead to depletion of viral genomes while mutagenesis of the provirus genome at important coding sequences or regulatory regions may inactivate viral replication. Additionally, the use of multiple DSBs might be used to excise proviral genomes<sup>64</sup>. As viral sequences may bear little sequence homology to the host genome, this class of treatments may exhibit less off target effects than editing therapies targeting endogenous loci.

## Factors Influencing Therapeutic Efficacy

Genome editing has been successfully applied to a number of diseases at the preclinical level as well as in a phase I clinical trial (Table 2)<sup>46,48,49,69,70</sup>. In evaluating the feasibility of a genome editing based therapy, the therapeutic effect of the desired genetic change should first be clearly established. Subsequently, the success of a given strategy will depend

on the ease with which a therapeutic modification ‘threshold’ is achieved, a criteria that is governed by the fitness of edited cells, the DSB repair pathway utilized to edit the genome, and the efficiency of delivery of genome editing molecules to target cell types.

### Fitness of Edited Cells

If edited cells have an increased fitness relative to unedited cells, this will result in a selective advantage for edited cells, reducing the number of cells that initially needs to be edited to reverse disease symptoms (Figure 2). For example SCID-X1 is caused by mutations in the *IL2RG* gene, the function of which is required for proper development of the hematopoietic lymphocyte lineage (<http://www.omim.org/entry/300400>). Hematopoietic progenitor cells with an active *IL2RG* gene selectively expand relative to their unedited counterparts. For example in people with SCID-X1 who received viral gene therapy for SCID-X1<sup>71,72</sup>, as well in a rare affected individual who had a spontaneous correction of a SCID-X1 mutation in a T-cell progenitor<sup>73</sup>, corrected hematopoietic progenitor cells were able to overcome the hematopoietic development block and expand relative to their diseased counterparts to mediate a therapeutic effect.

In contrast, other diseases in which edited cells do not exhibit a change in fitness, the number of cells that must initially be modified to reach a therapeutic effect is higher. For example, chronic granulomatous disorder (CGD), is caused by mutations in genes encoding phagocytic oxidase proteins that are involved in the generation of reactive oxygen species by neutrophils to kill pathogens (<http://www.omim.org/entry/306400>). Dysfunction of phagocytic oxidases does not influence hematopoietic cell fitness or development thus there would probably be no preferential expansion of edited cells in this disease. Indeed, no selective advantage for gene corrected cells in CGD has been observed in gene therapy trials, leading to difficulties with long-term cell engraftment<sup>74,75</sup>.

Some diseases where edited cells do not confer a change in fitness can still have reversed diseases symptoms at low numbers of therapeutic modified cells. For example, genes that function in a non-cell-autonomous fashion may only require a small number of functioning alleles to produce enough gene product to treat disease. For instance, Haemophilia B, is caused by mutations in the gene encoding the secreted factor IX protein involved in the clotting cascade, severe disease is associated with less than 1% of normal activity, whereas restoration of at least 1% of factor IX activity prevents the most severe bleeding conditions, while higher levels of restoration will further improve other clinically relevant complications in human patients<sup>76,77</sup>. This suggests that small changes in the amount of factor IX activity through correction of alleles in even a small percentage of liver cells may be therapeutic. Indeed, a study using ZFNs to correct a mouse model of haemophilia B shortly after birth demonstrated that correction of 3-7% of mutated Factor IX alleles was sufficient to reverse disease symptoms, providing preclinical evidence for this hypothesis<sup>48</sup>.

In the case where editing imposes a fitness disadvantage, such as the correction of mutated tumor suppressor genes in cancer cells, modified cells would be outcompeted by their diseased counterparts, causing the benefit of treatment to be low. The modification threshold of this final class of diseases would be extremely high requiring many cells to be directly modified and may not be suited for genome editing therapy. Therefore given the current

state of technology genome editing therapies are most ideally suited for cases where editing confers a fitness advantage or where a small change in gene product levels can influence clinical outcomes.

### Efficiency of Genome Editing

The efficiency of NHEJ and HDR mediated DSB repair varies significantly by cell type and cell state, NHEJ is most of the time more active than HDR. This difference in activity makes treating diseases that require gene correction or insertion of a gene more challenging than those requiring gene inactivation. NHEJ is thought to be active throughout the cell cycle and has been observed in a variety of cell types, including dividing and post-mitotic cells<sup>78,79</sup>. Therefore NHEJ may be used to facilitate high levels of gene disruption in target cell populations. In contrast, HDR acts primarily during S/G2 phase, and is therefore largely restricted to cells that are actively dividing, limiting treatments that require precise genome modifications to mitotic cells<sup>80,81</sup>.

The efficiency of correction by HDR may be controlled by a number of factors. First, the nature of genome modification may influence editing rates as large HDR mediated insertions have been found to occur at a reduced rate relative to HDR mediated small deletions, insertions or substitutions<sup>61,82</sup>. Second, the exact sequence changes made through HDR may influence therapeutic efficacy, as editing events that do not destroy the nuclease recognition site may be subject to further mutagenesis by NHEJ, potentially reducing therapeutic editing rates. Third, increasing the extent of homology between the repair template and the DSB site may increase HDR, possibly by promoting the stability of D-loop intermediates formed during synthesis from a template<sup>82-84</sup>. Fourth, the topology of the HDR template may influence editing efficiency as single-stranded DNA oligonucleotides and viruses may yield higher HDR rates than double-stranded substrates<sup>85,86</sup>. Last, suppressing competing DNA repair pathways such as NHEJ has also been shown to increase HDR rates moderately<sup>87</sup> although the safety of this strategy is not known and should be carefully assessed before implementation in a therapeutic context.

In addition to these approaches, further investigations aimed at improving HDR efficiency will be necessary to address a broader range of diseases with genome editing. Furthermore, many of these approaches may be synergistic and can be implemented in combination to increase the rate HDR past the therapeutic editing threshold needed to treat many diseases. Despite the challenges associated with HDR, proof-of-concept preclinical HDR treatments have now been described for mouse models of haemophilia B and hereditary tyrosinemia<sup>48,49</sup>.

### Modes of Delivery: Ex Vivo vs. In Vivo Editing

Achieving therapeutic editing requires delivery of programmable nucleases to target cells, which can be achieved either *ex vivo*, by modifying and autologously transplanting cells, or *in vivo*, by direct application of nucleases to diseased cells in the body (Figure 3). Given that nucleases can potentially be mutagenic, the ideal delivery system would permit transient nuclease activity. Currently, nucleases can be delivered as nucleic acids encoding the desired editing system, or directly as proteins themselves. While nucleic acids and protein



are both capable of achieving transient expression in target cell types, protein is likely to provide the best control over nuclease dosage, since there is no signal amplification. Another important consideration is that DNA based nuclease expression systems pose risks of insertional mutagenesis by the vector itself. So far, a variety of delivery methods have been developed.

In *ex vivo* editing therapy the target cell population is removed from the body, modified with programmable nucleases and then transplanted back into the original host (Figure 3-top panel). This mode of therapy allows the target cell population to be manipulated with a wide range of delivery platforms such as, electroporation, cationic lipids, cell penetrating peptides, carbon nanowires and viral vectors. *Ex vivo* therapies are likely to achieve high editing rates, due to the extensive development of these delivery systems for research and gene therapy applications. Moreover, many *ex vivo* therapies afford control over the specific dosage of therapeutic molecules delivered to cells. This may be particularly important when off-target modifications are a concern, as titrating the amount of nuclease may decrease such mutations<sup>88</sup>.

However, there are two large drawbacks with *ex vivo*. First, target cells must be capable of surviving manipulation outside the body which is a challenge for many tissues because cells either fail to survive, or lose properties necessary for their function *in vivo*. Thus, *ex vivo* therapy is largely limited to tissues with adult stem cell populations amenable to culture and manipulation, such as the hematopoietic system. Second, cultured cells often engraft poorly upon re-introduction into a patient, decreasing the effectiveness of treatment. However, engraftment may be enhanced for hematopoietic cells by ablative conditioning regimens that deplete unedited cells prior to transplantation. These are clinically feasible but introduce significant risks to patients<sup>89</sup>.

*In vivo* genome editing involves direct delivery of programmable nucleases to disease affected cells in their native tissues (Figure 3 bottom panels). There are two advantages of *in vivo* editing therapy over *ex vivo* approaches. First, *in vivo* editing can be applied to diseases where the affected cell population is not amenable to *ex vivo* manipulation. Second, *in vivo* delivery has the potential to target multiple tissue types, potentially allowing for the treatment of diseases that affect multiple organ systems. These properties will probably allow *in vivo* treatment to be applied to a wider range of diseases than *ex vivo* therapies.

To date, *in vivo* editing has largely been achieved through the use of viral vectors with defined, tissue-specific tropism. Such vectors are currently limited in terms of cargo carrying capacity and tropism, restricting this mode of therapy to organ systems where transduction with clinically useful vectors is efficient, such as the liver, muscle and eye<sup>90-92</sup>. Another major potential barrier for *in vivo* delivery is the immune response that may be raised in response to the large amounts of virus necessary for treatment, this phenomenon is not unique to genome editing and is observed with other virus based gene therapies<sup>93</sup>. It is also possible that peptides from editing nucleases themselves will be presented on MHC Class I molecules to stimulate an immune response, although there is little evidence to support this happening at the preclinical level. Another major difficulty with this mode of therapy is controlling the distribution and consequently the dosage of

genome editing nucleases *in vivo*, leading to off-target mutation profiles that may be difficult to predict. To address some of the concerns associated with viral vectors, non-viral delivery systems are under active development to reduce the potential risks currently associated with the use of viral vectors and expand the range of targetable tissues (reviewed in <sup>94</sup>).

The potential clinical complications faced by therapeutic genome editing overlap significantly with those of gene therapy, which make use of similar delivery agents and result in expression of novel gene products in the host. For a more in-depth discussion of the safety concerns regarding transgene expression and viral vectors for therapy, the reader is referred to recent reviews and studies on gene therapy.<sup>95,96</sup>

## Examples of Successful Genome Editing Therapeutic Strategies

### Ex Vivo Editing Therapy

The long standing clinical expertise with the purification, culture and transplantation of hematopoietic cells has made diseases affecting the blood system such as SCID, Fanconi anemia, Wiskott-Aldrich syndrome and sickle cell anemia the focus of *ex vivo* genome editing therapy. Another reason to focus on hematopoietic cells is that, thanks to previous efforts to design gene therapy for blood disorders, delivery systems of relatively high efficiency already exist. Despite these advantages, the often low efficiency of cell engraftment upon transplantation necessitates that this mode of therapy be applied to diseases where edited cells possess a fitness advantage, so that a small number of engrafted, edited cells can expand and treat disease. One such disease is HIV, as HIV infection results in a fitness disadvantage to CD4+ T cells.

The rationale for genome editing for HIV treatment originates from the observation that individuals homozygous for loss of function mutations in CCR5, a cellular co-receptor for the virus, are highly resistant to infection and otherwise healthy, suggesting that mimicking this mutation with genome editing could be a safe and effective therapeutic strategy<sup>57</sup>. This idea was clinically validated when an HIV infected patient was given an allogeneic bone marrow transplant from a donor homozygous for a loss of function CCR5 mutation, resulting in undetectable levels of HIV and restoration of normal CD4+ T-cell counts<sup>97</sup>. Although bone marrow transplantation is not a realistic treatment strategy for most HIV patients, due to the limited numbers of CCR5-null donors and potential graft vs. host disease, HIV therapies that convert a patient's own T-cells into CCR5 null cells are.

Early studies using ZFNs and NHEJ to knockout CCR5 in humanized mouse models of HIV showed that transplantation of CCR5 edited CD4+ T cells improved viral load and CD4+ T-cell counts<sup>70</sup>. Importantly, these models also showed that HIV infection resulted in selection for cells not expressing CCR5. As a result of this promising study, genome editing therapy that knocks out CCR5 in patient T cells has now been tested in humans. In a recent phase I clinical trial, CD4+ T cells from patients with HIV were removed, edited with ZFNs designed to knockout the CCR5 gene, and autologously transplanted back into patients<sup>46</sup>. Early results from this trial suggest that genome editing through ZFNs of the CCR5 locus is



safe, although the follow up time is too short to fully understand the risks and efficacy of treatment.

Gene correction strategies have also been successfully demonstrated in a recent study in which a mutated IL2RG gene was restored in hematopoietic stem cells (HSCs) obtained from a patient suffering from SCID-X1<sup>47</sup>. First, HSCs were transduced using integration-deficient lentivirus containing an HDR template encoding a therapeutic cDNA for IL2RG. Following transduction, cells were electroporated with mRNA encoding ZFNs targeting a mutational hotspot in IL2RG to stimulate HDR based gene correction. To increase HDR rates, culture conditions were optimized with small molecules to encourage HSC division. This strategy resulted in gene corrected HSCs from the SCID-X1 patient being obtained in culture at therapeutically relevant rates. HSCs from unaffected individuals that underwent the same gene correction procedure could sustain long-term hematopoiesis in mice. HSCs are capable of giving rise to all hematopoietic cell types and can be autologously transplanted, making them an extremely valuable cell population for all hematopoietic genetic disorders<sup>98</sup>. Gene corrected HSCs could, in principle, be used to treat a wide range of genetic blood disorders making this study an exciting breakthrough for therapeutic genome editing.

### In Vivo Genome Editing Therapy

*In vivo* genome editing therapy faces similar challenges to *ex vivo* strategies and is also limited by the small number of efficient delivery systems. Inefficient modification of target loci will be compounded by any inefficiencies in delivery, making tissues lacking robust delivery platforms particularly difficult to treat with this mode of therapy. For organ systems where delivery is efficient, however, there have already been a number of exciting preclinical therapeutic successes.

The first example of successful *in vivo* editing therapy was demonstrated in a mouse model of haemophilia B<sup>48</sup>. Recovering Factor IX activity to above 1% of its levels in severely affected individuals can transform the disease into a milder form, as infusion of recombinant Factor IX into such patients prophylactically from a young age to achieve such levels largely ameliorates the most severe bleeding complications<sup>76</sup>. In addition, Factor IX is synthesized and secreted by the liver, an organ that can be transduced efficiently by viral vectors encoding editing systems.

Using hepatotropic adeno-associated viral (AAV) serotypes encoding ZFNs and a corrective HDR template, up to 7% gene correction of a mutated [AU: 7% of what?], humanized Factor IX gene in the murine liver was achieved<sup>48</sup>. This resulted in improvement of clot formation kinetics, a measure of the function of the clotting cascade, demonstrating for the first time that *in vivo* editing therapy is not only feasible, but also efficacious.

Building on this study, other groups have recently used *in vivo* genome editing of the liver with CRISPR-Cas9 to successfully treat a mouse model of hereditary tyrosinemia and to create mutations that provide protection against cardiovascular disease<sup>49,99</sup>. These two distinct applications demonstrate the versatility of this approach for disorders that involve hepatic dysfunction. Application of *in vivo* editing to other organ systems will be necessary

to prove that this strategy is widely applicable. Currently, efforts to optimize both viral and non-viral vectors are underway to expand the range of disorders that can be treated with this mode of therapy<sup>90,94</sup>.

## Challenges to Clinical Translation

Translating genome editing technologies to the clinic faces major challenges, primarily in terms of the safety and efficacy of these treatments. Due to the distinctly different molecular nature of these therapies compared to small molecule and biologic therapies, engineering developments in several areas will be needed to bring these tools to bear on clinical medicine.

### Increasing Efficiency of Gene Correction

Although the amount of genome modification in a target cell population required to create a therapeutic effect differs depending on the disease, the efficacy of most editing treatments will be improved with increased editing rates. As previously noted, editing rates are controlled by the activity of DSB repair pathways.

Since NHEJ-mediated DSB repair is active in most cell types and are efficient, the primary challenge to date has been to increase the efficiency of HDR. So far, applications of HDR in genome editing have been primarily limited to dividing cells, due to the selective expression of HDR machinery during cell division and its down-regulation in slowly cycling or post-mitotic cells. Cell cycle regulation has now been somewhat by-passed for slowly cycling cell types through stimulation of mitosis with pharmacologic agents *ex vivo*<sup>47</sup>. However, truly post-mitotic cells are unlikely to be amenable to such manipulation, limiting the applicability of this strategy. Nevertheless, further work that will enable precise gene correction in post-mitotic cells such as neurons are critical to developing therapeutic strategies for a large number of untreatable neurological disorders. The solution to improved HDR in neurons will likely surface as we improve our understanding of DNA damage repair mechanism in the brain, and through harnessing of heterologous mechanisms. For example, the neurotrophic Herpes Simplex Virus (HSV), which depends on single-strand annealing (SSA), a form of HDR, to replicate, supplements exogenous viral proteins to facilitate viral replication, might provide answers to achieving efficient gene correction in post-mitotic cells<sup>100</sup>.

Additional, non-HDR-based strategies may also be successful for facilitating precise gene correction in post-mitotic cells. For example, attempts have been made to completely circumvent the need for HDR through direct ligation of DNA templates containing therapeutic transgenes into targeted DSBs. Such ligation events have been successfully demonstrated using ZFN<sup>101</sup>, but the rates are low for Cas9<sup>102,103</sup>. This difference might be due to cleavage patterns between ZFN and Cas9, whereas ZFN generates a predictable 4-bp overhang and Cas9 generates a blunt cut. Future structure-guided engineering may be able to alter the cleavage pattern of Cas9 to generate a sticky ends.

## Understanding and Improving Specificity of Editing Nucleases

The specificity of genome editing tools is one of the main safety concerns for clinical application. Genetic modifications are permanent, and deleterious off-target mutations have the potential to create cells with oncogenic potential, reduced fitness, or functional impairment. Furthermore, oncogenic mutations resulting from off-target editing may lead to expansion of edited cells, thus, even low levels of off-target mutagenesis may have devastating consequences.

Two issues remain outstanding: evaluating and reducing off-target effects. A number of studies have attempted to evaluate the targeting specificity of ZFN, TALEN, and Cas9 nucleases. The limited number of studies characterizing ZFN<sup>104,105</sup> and TALEN<sup>106</sup> specificity have only highlighted the challenges of detecting ZFN and TALEN off-target activity. Of note, the two independent studies attempting to characterize the off-target profile of the same pair of CCR5-targeting ZFNs have returned distinct and non-overlapping off-target sites, which highlights the challenges associated with analysis of nuclease specificity.

Many studies have attempted to evaluate the specificity of Cas9, partly owing to the simplicity of the RNA-guided DNA targeting mechanism of Cas9, which makes it significantly easier to establish hypotheses regarding possible off-targeting mechanisms based on Watson-Crick base pairing rules. While initial bacterial<sup>39</sup>, biochemical<sup>40,41</sup>, and mammalian<sup>42</sup> experiments have suggested that the 3' 8-12bp seed region of the guide sequence can be sensitive to single base mismatches, further work have shown that this rule-of-thumb is not necessarily accurate, especially in situations where there is high concentration of Cas9 and guide RNA<sup>88,107-110</sup>. Many of these studies were carried out in cell lines and examined Cas9-mediated mutagenesis at genomic sites bearing high levels of homology to the on-target sequence and found that, unsurprisingly, subsets of highly homologous off-target sites were significantly mutated by the nuclease. However, the scope of possible off-target sites evaluated by these studies was limited to computationally predicted sites.

More recently, whole-genome sequencing of Cas9-edited cell lines revealed low incidence of off-target mutation, which suggests that Cas9-mediated genome editing may be specific<sup>111-113</sup>. Despite these studies, unbiased assessment of genome wide off-targeting using more advanced methods like direct capture of DSBs<sup>114</sup>, labeling of DSBs with oligo captures<sup>115</sup>, and techniques that can detect larger structural variations (i.e. translocations) potentially imposed by nuclease treatment<sup>116</sup> will help us further understand the true risk of mutagenesis imposed by programmable nucleases. It is worth noting that off-target effects may be cell-type specific; for example off-target effects in transformed cell lines with dysregulated DSB repair pathways may overestimate off-target effects that would be observed in healthy primary cells.

In order to reduce the frequency of off-target effects, many groups are rapidly improving the targeting specificity of Cas9. For example, transformation of Cas9 into a single-strand DNA nickase that primarily generates DSBs by creating two separate single-strand breaks on opposite DNA strands via the expression of two separate guide RNAs, reduces off-target

indel formation at computationally predicted off target sites<sup>102,109</sup>. Additionally, truncation of the guide RNA as well as RNA-guided FokI nuclease based on fusion between catalytically inactive Cas9 and the FokI nuclease domain are also able to achieve improved levels of targeting specificity<sup>117-119</sup>.

Alternative genome editing strategies not involving nucleases have also been explored, and may pose a lower mutagenic risk<sup>62</sup>. AAV genomes containing transgenes flanked by homology to target loci are capable of stimulating HDR in the absence of a nuclease, albeit at lower rates<sup>48,86,120-122</sup>. Using this strategy one study targeted a factor IX cDNA to the highly expressed albumin locus, allowing for correction of the bleeding diathesis phenotype in factor IX deficient mice<sup>62</sup>. By targeting a highly expressed locus the authors could achieve 7-20% of factor IX levels, despite an HDR rate of 0.5%. Although this strategy may not be widely applicable due to the low absolute targeting rate, this and future, improved nuclease strategies should also be considered for therapeutic applications.

## Delivery

Another major challenging facing clinical translation is delivery of editing systems to target cell types. A variety of nucleic acid or protein delivery methods may be used to introduce genome editing nucleases into target cells *ex vivo* or *in vivo* (**Figure 3**). Depending on the choice of delivery method, the nucleases may either be transiently or permanently expressed in the target cell. Given that nucleases may exhibit off-target cleavage activity or trigger immune responses, the delivery system should be carefully selected.

For *ex vivo* applications, such as editing of hematopoietic stem cells, electroporation may be used to achieve transient nuclease expression through delivery of DNA-based nuclease expression vectors, mRNA, or protein. Both integration-competent and deficient lentiviral vectors have also been successfully used to drive nuclease expression. However, integrating lentiviral vectors may be less desirable as they drive constitutive expression and may result in more off-target activity. In addition, all three nuclease platform have also been demonstrated to be amenable to modifications so that proteins can be directly delivered into cells either through engineered cell-penetrating peptides or chemical conjugation<sup>106,123,124</sup>.

For *in vivo* applications, the most promising delivery systems are viral vectors, particularly adeno-associated viral (AAV) vectors, which have recently been approved for clinical use<sup>125</sup>. AAV comes in many serotypes and have been shown to have high delivery efficacy for a variety of tissue types including the eye, brain, liver, and muscle<sup>126</sup>. However, the relatively small packaging capacity of AAV vectors post some challenges for nuclease delivery. Whereas ZFNs are relatively small and a dimeric ZFN pair can be packaged into a single AAV, a dimeric TALEN pair is much larger and will likely need to be packaged into two separate AAV vectors. For Cas9, short orthologs may be packaged along with guide RNAs into a single AAV. So far, AAV-mediated nuclease expression has been demonstrated to be successful in several tissue types, including liver and brain<sup>48,127</sup>. In the case of viral mediated Cas9 delivery, which may result in constitutive expression of nuclease proteins and cause genome instability and toxicity, self-cleaving mechanisms may be used to inactivate the nuclease transgene on the delivery vector<sup>128</sup>.

Despite the potential of AAV-mediated *in vivo* nuclease expression, there are several challenges that will require further development. First, AAV-mediated nuclease expression is often constitutive and it would be more desirable to be able to shut down nuclease expression after the genome editing event has successfully occurred in the target cell. Second, patients who have already been naturally exposed to AAV will likely have developed immunity against specific serotypes. Therefore AAV may not be an appropriate delivery vehicle for these patients.

To overcome these challenges faced by viral vectors, nanoparticle- and lipid-based *in vivo* mRNA or protein delivery systems may provide an attractive alternative<sup>123,129</sup>. Delivery of nuclease mRNA via nanoparticle conjugation, or nuclease proteins will permit more precise dosage control, which has been shown to affect the level of off-target mutation rate<sup>88,124</sup>. mRNA or protein delivery will also be transient, therefore minimizing any potentially undesirable nuclease-induced toxicity. Finally, for nuclease protein delivery, especially TALENs and Cas9, which are derived from microbial origins, exposure of the microbial proteins may stimulate immune reactions. Potential strategies for circumventing immunotoxicity resulting from protein delivery may include limiting dosage and protein humanization to reduce immunogenicity<sup>130</sup>.

## Conclusion

The enormous excitement surrounding genome editing needs to be coupled with strategic planning and rigorous but enabling regulatory processes to ensure successful development of this class of potentially life-changing medicine. The technology will require a number of iterations to systematically optimize its efficacy, safety, and specificity. Despite being in its infancy, genome editing presents tantalizing opportunities for tackling a number of diseases that are beyond the reach of previous therapies. Given the accelerating pace of technological advances and broad range of basic science and clinical applications, the road ahead will undoubtedly be an exciting one.

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### Box: Using Genome Editing to Treat Non-monogenic Diseases

#### Introduction of protective mutations for complex diseases treatment

The abundance of genetic information has made it possible to identify naturally occurring mutations that confer resistance to disease phenotypes. These mutations occur in both coding and non-coding regions of the genome and have received attention as therapeutic targets for complex, non-monogenic diseases such as cardiovascular disease<sup>58,136</sup>, HIV<sup>97</sup>, Alzheimer's disease<sup>59</sup> and hemoglobinopathies<sup>137</sup>. Genome editing provides the possibility of introducing these protective mutations into patients to reverse illness.

Many known protective mutations involve loss of function alleles, which can be introduced via NHEJ-mediated gene disruption. This approach has rapidly gained traction due to the high efficiency of NHEJ in therapeutically accessible cells and treatment using this strategy is currently in clinical trials for HIV<sup>46</sup>. [AU: following paragraph removal ok to save space in the box. The HIV example is extensively discussed in the main text]

Mutations that protect against disease also lie hidden outside the coding region of the genome. GWAS and CHIP-seq were recently used together to identify non-coding mutations that may be important targets for genome editing therapy<sup>138,139</sup>. A major factor controlling sickle cell disease severity is the expression level of fetal hemoglobin (HbF), increased HbF levels decreasing disease severity. A GWAS for regions controlling HbF expression identified variation within the *BCL11A* gene, the product of which is known to negatively regulate HbF expression<sup>138,139</sup>. This variation promotes transcription factor binding within an intron that enhances *BCL11A* expression in the erythroid lineage, thereby decreasing HbF expression levels in red blood cells. TALENs were used to directly remove this intron from erythroid cells [AU:OK?] and showed that this resulted in an increase in HbF levels<sup>137</sup>. However, this study was not carried out in HSCs, the most therapeutically relevant cell population for this disease. It will be exciting to see if this approach can be extended to these cells to provide a clinical benefit for patients. Furthermore, it is worth noting that non-coding regions will likely hold therapeutically important regions: 93% of GWAS hits, disease- and trait-associated are found within noncoding regions<sup>140</sup>.

#### Programmable Nucleases as Antivirals

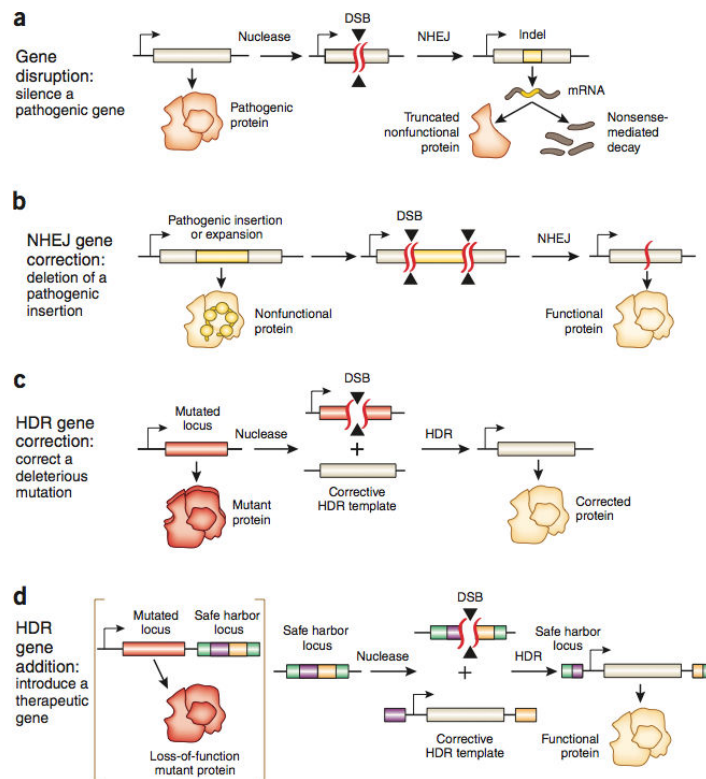
In addition, programmable nucleases may be developed as anti-viral strategies.

In principle, nucleases may be used to target viral sequences for cleavage and subsequent destruction. Additionally, NHEJ based mutagenesis of elements critical for viral fitness could render latent viruses incapable of propagating infection. Alternatively, multiplexed nucleases like Cas9 could be used to excise provirus from the genomes of infected cells, leading to their degradation by cellular nucleases.

Efforts to develop genome editing nuclease for antiviral therapy have largely focused on HIV, where large reservoirs of latent provirus can persist in the presence of anti-retroviral therapies and serve to re-activate infection once treatment is stopped. The long-terminal repeats (LTRs) of HIV drive viral gene expression and are critical for viral fitness. By

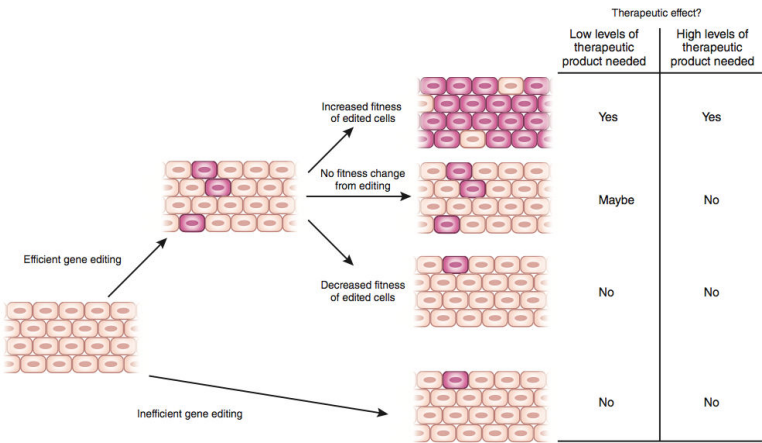


targeting Cas9 to cleave LTR sequences, a study recently demonstrated the possibility of mutating the proviral LTR and significantly reduces expression of HIV genes in T cells <sup>64</sup>. Although an exciting discovery, there are several additional challenges to translate this *in vitro* result to the clinic. Likely the biggest challenge will be delivering nucleases to all HIV carrying cells in an infected individual to eliminate all of the latent provirus. Currently, there are no therapeutic platforms capable of delivering genome editing nucleases to the majority of T cells. Similar strategies have been seen with HPV <sup>141</sup>, and HBV <sup>65,66</sup> but most infectious diseases face the same problem as HIV: extremely efficient delivery of genome editing tools is likely required to achieve complete removal of viral infection.



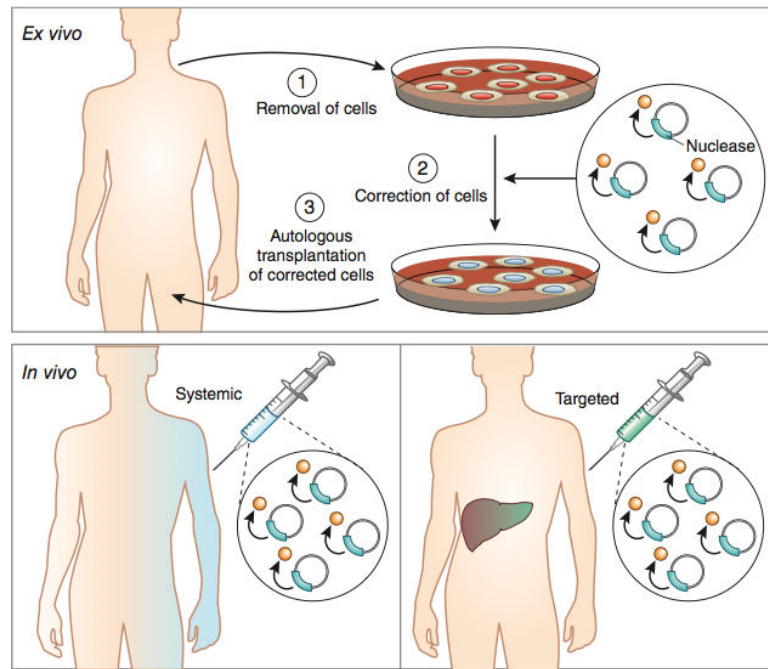
**Figure 1. Types of Therapeutic Genome Modifications**

The specific type of genome editing therapy depends on the nature of the mutation causing disease. **a**, In gene disruption, the pathogenic function of a protein is silenced by targeting the locus with NHEJ. Formation of indels on the gene of interest often result in frameshift mutations that create premature stop codons and a non-functional protein product, or non-sense mediated decay of transcripts, suppressing gene function. **c**, HDR gene correction can be used to correct a deleterious mutation. A DSB is induced near the mutation site in the presence of an exogenously provided, corrective HDR template. HDR repair of the break site with the exogenous template corrects the mutation, restoring gene function. **d**, An alternative to gene correction is gene addition. This introduces a therapeutic transgene into either the native or a non-native locus in the genome. A DSB is induced at the desired locus and an HDR template containing homology to the break site, a promoter, a transgene and a polyadenylation (polyA) sequence is introduced to the nucleus. HDR repair recovers gene function in the target locus albeit without true physiological control over gene expression. **b**, In NHEJ gene correction two DSBs targeted to both sides of a pathogenic expansion or insertion may be resolved by NHEJ, causing a deletion of the intervening sequences to mediate therapy. This form of treatment would require multiplexed targeting of disease causing mutations.



**Figure 2. Factors Influencing Therapeutic Efficacy**

For a genome editing therapy to be efficacious, enough cells carrying the desired genome modification must exist in a tissue to reverse disease. If editing is efficient, treatment will create a population of cells carrying the desired genomic modification (depicted in pink). Depending on whether the editing event creates a fitness change in target cells, edited cells will proportionally increase, or decrease relative to unedited cells (depicted in brown) over time in tissues. Proportionally high levels of cells carrying therapeutic genome modifications in a disease-affected tissue are likely to result in a therapeutic effect. However, if low levels of a secreted gene product are needed to reverse disease, then successfully editing a small number of cells may be therapeutically efficacious.



**Figure 3. *Ex Vivo* vs. *In Vivo* Editing Therapy**

In *ex vivo* editing therapy cells are removed from a patient, edited and then re-engrafted (top panel). For this mode of therapy to be successful, target cells must be capable of survival outside the body and homing back to target tissues post-transplantation. *In vivo* therapy involves genome editing of cells *in situ* (bottom panels). For *in vivo* systemic therapy, delivery agents that are relatively agnostic to cell identity or state would be used to effect editing in a wide range of tissue types. For example systemic delivery of AAV serotype 8 vectors has been used in preclinical models to target liver tissue with high efficiency. Alternatively, *in vivo* therapy, may also be achieved through local injection of viral vectors to the affected tissue, such as the eye, brain, or muscle.

**Table 1**

Comparison of Different Programmable Nuclease Platforms.

	<b>Zinc Finger Nuclease</b>	<b>TALEN</b>	<b>Cas9</b>	<b>Meganuclease</b>
Recognition site	Typically 9 to 18 bp per ZFN monomer, 18 to 36 bp per ZFN pair	Typically 14 to 20 bp per TALEN monomer, 28 to 40bp per TALEN pair	22bp (20bp guide sequence + 2bp PAM sequence for <i>S. pyogenes</i> Cas9); up to 44 bp for double nicking	Between 14 and 40 bp
Specificity	Small number of positional mismatches tolerated	Small number of positional mismatches tolerated	Positional and multiple consecutive mismatches tolerated	Small number of positional mismatches tolerated
Targeting constraints	Difficult to target non-G-rich sequences	5' targeted base must be a T for each TALEN monomer	Targeted sequence must precede a PAM [AU: please define]	Targeting novel sequences often results in low efficiency
Ease of engineering	Difficult, may require substantial protein engineering	Moderate, requires complex molecular cloning methods	Easily re-targeted using standard cloning procedures and oligo synthesis	Difficult, may require substantial protein engineering
Immunogenicity	Likely low, as ZFs are based on human protein scaffold. FokI is derived from bacteria and may be immunogenic	Unknown, protein derived from <i>Xanthomonas sp.</i>	Unknown, protein derived from various bacterial species	Unknown, meganucleases may be derived from many organisms including eukaryotes
Ease of ex vivo delivery	Relatively easy through methods such as electroporation and viral transduction	Relatively easy through methods such as electroporation and viral transduction	Relatively easy through methods such as electroporation and viral transduction	Relatively easy through methods such as electroporation and viral transduction
Ease of in vivo delivery	Relatively easy due to small size of ZFN expression cassettes, allows use in a variety of viral vectors	Difficult due to the large size of each TALEN and repetitive nature of DNA encoding TALENs, leading to unwanted recombination events when packaged into lentiviral vectors	Moderate: The commonly used Cas9 from <i>S. pyogenes</i> is large and may impose packaging problems for viral vectors such as AAV, but smaller orthologs exist.	Relatively easy due to small size of meganucleases, allows use in a variety of viral vectors.
Ease of multiplexing	Low	Low	High	Low

**Table 2**

Examples of applications of genome editing to therapeutic models.

Disease Type	Nuclease Platform Employed	Therapeutic Strategy	References
Hemophilia B	ZFN	HDR-mediated insertion of correct gene sequence	48
HIV	ZFN and CRISPR	NHEJ-mediated inactivation of CCR5	46,69,70,131
DMD	CRISPR and TALEN	NHEJ-mediated removal of stop codon, and HDR-mediated gene correction	132,133
HBV	TALEN and CRISPR	NHEJ-mediated depletion of viral DNA	65,66
SCID	ZFN	HDR-mediated insertion of correct gene sequence	47
Cataract	CRISPR	HDR-mediated correction of mutation in mouse zygote	134
Cystic fibrosis	CRISPR	HDR-mediated correction of CFTR in intestinal stem cell organoid	135
Hereditary tyrosinemia	CRISPR	HDR-mediated correction of mutation in liver	49