Impact of New Genome Editing Tools on iPS Cell Based Therapies

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Abstract
Embryonic stem cells (ES cells) have unique differentiation and self-renewal potential which makes them a potential candidate for cell therapy in degenerative disorders. However, the limitation of deriving ES cells from embryos and their immune-incompatibility hinder their clinical applications. Fortunately, by introduction of somatic cell reprogramming technology into induced pluripotent stem cells (iPS cells), the limitations of ES cells have been circumvented. Therefore iPS cells have been considered as a promising cell source for clinical application in regenerative medicine.

However, in many cases, disease-specific iPS cells need genome modification to replace/repair the defective gene and regain normal functions before any application. For this reason, Homologous recombination (HR) method is one of the best available options which allow proper gene editing without any undesirable side-effects associated with random integration based methods. However, the efficiency of traditional HR in mammalian cells is extremely low which makes it ineffective for gene correction in human iPS cells. Fortunately with the recent advancement of new gene editing tools which allow site specific DNA targeting, HR efficiency in iPS cells have been significantly improved. Among these new tools, ZFNs, TALENs and CRISPRs are the most promising ones. This editorial briefly reviews their advantages and shortcomings and debates the impact of these new gene editing technologies on the future of iPS based cell therapies.

Keywords
Genome editing; Human iPS cells; iPS cell therapy; Gene therapy; Induced pluripotent stem cells.

Cell Therapy and Stem Cells
Stem cells are undifferentiated cells which can self-renew and differentiate into any specialized cell types. Based on their sources, stem cells from human tissues can be classified as embryonic stem cells (ES cells) and adult stem cells. ES cells isolated from human blastocyst-stage embryos have the capacity for unlimited replication and can be differentiated into all the cell types in human body [1]. However, human adult stem cells derived from non-embryonic tissues are multipotent and can differentiate into only limited number of cell types. Therefore, transplantation of stem cells or their derivatives, known as stem cell therapy, have been considered as one of the most promising approaches to restore the damaged tissues or organs in case of many human diseases such as degenerative disorders [2-4].

However, human ES cells and adult stem cells have limitations which impede their clinical application. Among these, ethical issues and potential immunologic rejection are major obstacles which limit clinical application of human embryonic stem cells [2,4]. On the other hand, in the case of adult stem cells, only limited numbers exist in the adult tissues and they generally have poor in vitro expansion/self-renewal potentials [5]. Therefore obtaining large quantity of these cells for transplantation is a big challenge. Furthermore, they also carry immune compatibility issues in case of allogeneic transplants. For example, even in the case of hematopoietic stem cell (HSC) transplantation, the most successful example of cell therapy, over a third of patients who are eligible will not find a suitable human leukocyte antigen (HLA)-matched donor [6]. Fortunately, induced pluripotent stem cell (iPS cell) technology overcomes many of these obstacles and has opened a new era in cell therapy for many genetic and degenerative human disorders.

iPS Technology Evolution
In 2007 the groups of Yamanaka’s and Thomson’s successfully transformed human fibroblasts into human induced pluripotent stem cells (iPS cells) by over-expression of four transcription factors; Oct4, Sox2, KLF4, and c-Myc or Oct4, Sox2, Nanog, and Lin28 [7,8]. These iPS cells are similar to ES cells in most aspects, including self-renewal and differentiation potential. Therefore, Patient-derived iPS cells offer a promising solution to the major obstacles of stem cell therapy, such as the ethical concerns and immune rejection. This was a great breakthrough for science community as it can potentially provide individualized stem cell therapy for the patients [9].

However, initially both groups had applied retrovirus or lentivirus system for transgene expression, which resulted in the random integration of exogenous genes in the host genome. Even though the expressions of viral transgenes were silenced after complete reprogramming, they still had chances to be re-activated or lead to tumor formation due to random transgene integration into genome [10].

In order to solve these problems, researchers have developed several non-integrative approaches to generate safe iPS cells.
Fusaki et al. used Sendai viral vectors to generate iPS cell lines and successfully obtained iPS clones without viral RNA. However, since these viral vectors replicate constitutively, it is still difficult to remove them from host cells and isolate transgene-free clones, even at late passages [11]. Several other research groups have developed episomal vectors (such as minicircle vectors) and have applied transfection to deliver reprogramming factors into target cells [12]. Though in these cases transgene-free iPS clones could be obtained, still the efficiency is quite low.

In another attempt to improve the safety of iPS cells, Warren et al. have recently used synthetic modified mRNAs for transgene expression in somatic cells and highly efficient iPS conversion was achieved after 16-days post transfection [13]. Several other research groups have developed episomal vectors (such as minicircle vectors) and have applied transfection to deliver reprogramming factors into target cells [12]. Though in these cases transgene-free iPS clones could be obtained, still the efficiency is quite low.

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Monocytes/Macrophages with homozygous deletions of 32-bp sequences (CCR5 Δ32) in C-C chemokine receptor type 5 gene - a major co-receptor used by HIV-1 virus to infect T cells, macrophages, and other cell types- can resist HIV-1 infection [19]. Therefore, gene editing is a necessary tool for altering iPS cells to a favorable cell therapy for many diseases not limited to only genetic disorders.

Gene Editing in iPS Cells

There are two major strategies for the genome modification of patient-derived iPS cells: random integration via viral vectors and site-specific correction via homologous recombination (HR). Although gain-of-function can be achieved through correction of disease-specific iPS cells by viral vectors, genome modifications via viral vectors lead to random integration of viral genes into the host cell genome and have uncertain risks, including tumor formation [20]. This is the major obstacle for using viral vectors for gene modification in iPS cells.

Therefore, the most promising strategy is site-specific genome modification via homologous recombination (HR) mechanism. As HR has a requirement of DNA repair proteins, the homology arms in HR are defined as a homologous sequence, which can be used for HR either in vivo or in vitro. HR involves the introduction of DNA double-strand breaks (DSBs) downstream of the target sites and using homology arms with sequence identities to the genome. In the presence of a homologous sequence downstream of the DSB, the introduction of a target gene allows the repair of the DSB by HR, which generates the insertion or replacement of a specific DNA sequence [21].

Recently, a number of genome editing technologies have been developed to introduce site-specific DSB, including zinc-finger nucleases (ZFNs), transcription activator-like effector nucleases (TALENs) and the RNA-guided CRISPR-Cas nucleosome system. Here we overview each method and evaluate their applications for iPS cell gene corrections.

Zinc-Finger Nucleases (ZFNs) and iPS Cells

A ZFN is composed of three protein domains: a nuclear localization signal domain at N-terminal, a nuclease domain derived from the FokI restriction enzyme at C-terminal and a DNA-binding zinc-finger protein (ZFP) domain in between (Figure 1A). The ZFP domain contains a tetratricopeptide repeat of zinc-fingers which recognize DNA sequence in a way of one to one correspondence. The three zinc-finger repeats allow the ZFNs to recognize DNA sequence of 9-18 bp [22]. Three zinc-finger repeats can also increase binding specificity, but it will also decrease the chances to find a suitable target site. ZFN subunits with 3-6 zinc-fingers have been successfully used to recognize a DNA sequence of 9-18 bp [23].

Furthermore, dimerizations of FokI domains are required for nuclease activity of this system. When two ZFN monomers bind DNA in opposite orientations and have an appropriate spacer to permit FokI dimer formation, they form an active endonuclease complex. The one-to-one correspondence of each finger to one base pair. These repeats are highly conserved in amino acid sequence, except the amino acids at positions 12 and 13, which determine the nucleotide binding specificity of each repeat [31].

In the absence of a DNA template, two ends of DSBS are religated through the NHEJ pathway and insertion/deletion (indel) mutations are introduced at the repair sites. In the presence of a repair template, DSBS can be fixed through the HDR pathway [24]. With the combination of ZFNs and gene targeting constructs with a desired sequence flanked by two homology arms, researchers can introduce defined modifications at the interest target sites or correct the mutations in the genome. This ability makes ZFNs as a potential tool for iPS genome editing and repair.

However, there are few limitations for this technology. For instance, although many of individual module fingers which target most of the possible 64 triplet-binding sites have been identified, it is still difficult to collect 64 zinc-fingers that cover all possible combinations of triplet sites. Therefore, it is still a challenge to generate ZFNs for all the target sequences. Furthermore, even if ZFNs can be successfully generated, some of ZFNs might not cleave genomic DNA efficiently [25]. Moreover, the off-target mutations introduced by ZFNs can lead to toxicity in host cells [26]. Additionally, design and construction of ZFNs is complex and time-consuming [27] as ZFNs are assembled by different zinc-fingers with similar structures. This makes it a complex and time-consuming process to generate ZFNs.

Despite these limitations, ZFNs have been used for gene correction of a number of patient-derived iPS cells. Zou et al. performed ZFNs-assisted gene correction of the β-globin mutation in the iPS cell lines derived from patients with sickle cell anemia [28]. They have successfully converted one of two mutated β-globin alleles to the wild-type. Unfortunately, after differentiation of gene-corrected heterozygous iPS cells into erythrocytes, the erythroid cells expressed the corrected allele only at the level of 25%-40% of the mutated allele. The remaining loxP sequences or a genetic mutation in HBB 3′-enhancer were speculated to be responsible for the low expression of the corrected allele [28]. ZFNs-mediated gene correction of the iPS cell lines from patients with β-thalassemia and α(1)-antitrypsin deficiency have also been reported recently [29,30]. These studies have confirmed the gene correction by DNA sequencing as well as in vitro and in vivo gain-of-function experiments. More importantly, in the case of α(1)-antitrypsin deficiency, the authors have sequenced exomes of the corrected iPS cells and its parental fibroblasts and did not detect any mutations introduced by off-target events [30]. Overall, despite of their hurdles, ZFNs are still considered as a practical tool for gene modification in iPS cells.

Transcription Activator-Like Effector Nucleases (TALENs) and iPS Cells

Like ZFNs, TALENs (Figure 1B) also contain three functional domains. But different from ZFNs, the DNA-binding domains of TALENs are derived from a class of naturally occurring DNA binding proteins, known as TALEs. TALEs consist of tandem repeats of 33-35 amino acids, which recognize DNA sequence in a way of one repeat to one base pair. These repeats are highly conserved in amino acid sequence, except the amino acids at positions 12 and 13, which are called repeat variable diresidues (RVDs). The residues of RVDs determine the nucleotide binding specificity of each repeat [31].

In the original repeat arrays, four RVDs have been identified: Asn-Asn, Asn-Ile, His-Asp and Asn-Gly, which recognize guanine, adenine, cytosine and thymine, respectively [32]. As mentioned above, the FokI domain functions as a dimer. Therefore, it requires two TALEN monomers with proper orientation and spacing to achieve DNA cleavage. The one-to-one correspondence between RVDs and base pairs make it possible to target almost any DNA sequence. Although it has been recently reported that the presence of 5-methylated cytosine at the target sites attenuate the binding affinity of TALENs, the replacement of His-Asp RVD repeat with an Asn-Gly RVD repeat can overcome this problem [33].

Consisting of only four kinds of RVD codes, TALENs can be generated within a few days which make them more attractive than ZFNs. Therefore, researches have applied it for the gene correction of disease-specific iPS cells. Ma et al. utilized TALENs to facilitate the gene correction of β-globin mutation in iPS cells from β-thalassemia patients [34]. The authors demonstrated that the gene correction did not cause off-target mutations and erythroblasts derived from corrected iPS cells expressed normal β-globin. TALENs-assisted gene correction was also successfully used to correct the NPC1 mutation in iPS cells derived from patients with Niemann-Pick type C (NPC) disease [35]. The disease-associated defects in NPC1-deficient human hepatic and neural cells were rescued in the cell types derived from corrected iPS cells. Overall, TALENs easier design compared to ZFNs makes them a favorable tool for gene correction in iPS cells.

RNA-Guided CRISPR-Cas Nucleosome System and iPS Cells

Most recently, a class of RNA-guided engineered nucleases have been developed from microbial adaptive immune systems named as clustered regularly interspaced short palindromic repeats (CRISPR)/
CRISPR-associated (Cas) systems [36,37]. These systems are used by bacteria and archaea to cleave foreign genetic elements. There are three types (I–III) of CRISPR/Cas systems [38]. Due to its simplicity, type II has been well characterized and used for genome engineering [39,40]. This system (Figure 1C) consists of three components: the nuclease Cas9, the CRISPR RNA (crRNA) array and a trans-activating crRNA (tracrRNA) which is required for the maturation of crRNA.

The crRNA array is a long array of direct DNA repeats separated by short variable sequence spacers. These spacers encode guide RNAs, which direct Cas9 to the target DNA through Watson-Crick base-pairing interaction. As the crRNA array can have more than one guide RNAs, targeting multiple sites at once can also be achieved. crRNA and tracrRNA form an active nuclease which this complex recognizes the protospacer-adjacent motif (PAM) site and initiates the crRNA-DNA interaction. In other words, the Cas9 system targets a 20-bp DNA sequence followed by a PAM site at its 3' end. The most popular Cas9, Streptococcus pyogenes cas9 (SpCas9) binds to the 5'-NGG-3' PAM sequence, and 5'-NAG-3' with a lower frequency, while the Cas9 derived from other species recognize different PAM sequences [41].

Cas9 contains two nuclease domains: RuvC and HNH, each of which cuts a different DNA strand and generate double strand breaks (DSBs) together. In order to improve the on-target specificity of Cas9 nuclease, two different kinds of strategies have been developed. In first strategy, a Cas9 nickase that cuts a single strand of DNA was generated by inactivating RuvC nuclease domain via point mutation (D10A) [42,43]. Directing by a pair of guide RNAs, which their target sites have proper space and orientation, the Cas9 nickase can generate a DSB efficiently (Figure 1D). As this strategy a) doubles the length of target sequence and b) the DNA single-strand breaks created by the nickases at off-target sites can be repaired via the high-fidelity base excision repair (BER) pathway, the nicking strategy improves target specificity by up to 1500 time relative to the wild-type Cas9 [42]. It has also been shown that gRNAs with 2-3 bp shorter sequence could further reduce off-target effects induced by Cas9 nickases, which might be a result of more sensitivity to mismatches [44].

In the second strategy, a catalytically inactive Cas9 was also generated and fused with a FokI nuclease domain (Figure 1E). This FokI-dCas9 nuclease was guided by a single gRNA. As The FokI domains must dimerize to acquire nuclease activity, the cleavage of FokI-dCas9 nuclease was guided by a single gRNA. As The FokI generated and fused with a FokI nuclease domain (Figure 1E). This FokI-dCas9 nuclease was guided by a single gRNA. Although in this case the frequency of indel mutations induced by a pair of FokI-dCas9 molecules to the target locus. Although in recent studies, only few mutagenesis at off-target site have been reported but most of these studies only examined off-target mutations at few predicted sites. Therefore, chances of mutagenesis at many other off-target sites still remain unknown. The extent of off-target side effects should be fully evaluated before the clinical application of genome editing technology. Meanwhile, further improvements in binding and cleavage specificity of the genome editing tools will minimize the off-target events and enhance their potential therapeutic application. These make the future of iPSC cell therapy very exciting and challenging road map to complete.

Concluding Remarks

So far, ZFNs, TALENs and CRISPR-Cas nucleases have been successfully used for genome modification of a number of disease-specific iPSCs, indicating that they are powerful tools for site-specific gene targeting and iPSC cell therapy will benefit from them in the future. Among these, CRISPR-Cas nucleases have shown more promises for future genome editing in human iPSCs cells due to their easy and fast design as well as their relative precision- especially in case of Cas9-nickase version- and efficacy for gene editing.

However, the biggest concern about all above mentioned tools is their safety, as all of these genome editing tools generate DSBs at the target sites as well as off-target sites, which will introduce indel mutations. Although in recent studies, only few mutagenesis at off-target site have been reported but most of these studies only examined off-target mutations at few predicted sites. Therefore, chances of mutagenesis at many other off-target sites still remain unknown. The extent of off-target side effects should be fully evaluated before the clinical application of genome editing technology. Meanwhile, further improvements in binding and cleavage specificity of the genome editing tools will minimize the off-target events and enhance their potential therapeutic application. These make the future of iPSC cell therapy very exciting and challenging road map to complete.

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