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Efficient ssODN-Mediated Targeting by Avoiding Cellular Inhibitory RNAs through Precomplexed CRISPR-Cas9/sgRNA Ribonucleoprotein

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SUMMARY

Combined with CRISPR-Cas9 technology and single-stranded oligodeoxynucleotides (ssODNs), specific single-nucleotide alterations can be introduced into a targeted genomic locus in induced pluripotent stem cells (iPSCs); however, ssODN knockin frequency is low compared with deletion induction. Although several Cas9 transduction methods have been reported, the biochemical behavior of CRISPR-Cas9 nuclease in mammalian cells is yet to be explored. Here, we investigated intrinsic cellular factors that affect Cas9 cleavage activity *in vitro*. We found that intracellular RNA, but not DNA or protein fractions, inhibits Cas9 from binding to single guide RNA (sgRNA) and reduces the enzymatic activity. To prevent this, precomplexing Cas9 and sgRNA before delivery into cells can lead to higher genome editing activity compared with Cas9 overexpression approaches. By optimizing electroporation parameters of precomplexed ribonucleoprotein and ssODN, we achieved efficiencies of single-nucleotide correction as high as 70% and *loxP* insertion up to 40%. Finally, we could replace the HLA-C1 allele with the C2 allele to generate histocompatibility leukocyte antigen custom-edited iPSCs.

INTRODUCTION

Human induced pluripotent stem cells (iPSCs) are important research tools for studying diseases and a promising cell source for regenerative medicine (Takahashi et al., 2007; Takahashi and Yamanaka, 2016). Recent advances in genome editing technologies enable the correction of a pathogenic mutation in patient iPSCs or the introduction of a desired genetic alteration in healthy iPSCs for studying genetic function or modulating cellular characteristics (Hockemeyer and Jaenisch, 2016; Hotta and Yamanaka, 2015). In particular, the CRISPR-associated protein (Cas9) system found in Streptococcus pyogenes has become the most widely utilized tool for genome editing (Jinek et al., 2012). Cas9 nuclease forms a complex with crRNA and tracrRNA (or single guide RNA [sgRNA]) to target a specific DNA sequence and induce a double-stranded break (DSB). The DNA is repaired by one of several DNA repair pathways, such as non-homologous end-joining (NHEJ), which introduces insertions/deletions (indels), and homologydirected repair (HDR), which requires a DNA donor template in order to make precise repairs.

Because more than half of human pathogenic mutations are single-nucleotide polymorphisms (SNPs) (Rees and Liu, 2018), the introduction of an SNP into iPSCs is of high importance. Such genetic manipulation can be utilized for correcting a pathogenic mutation or introducing a desired mutation. Among various genome editing tools, such as base editors and prime editing (Anzalone et al., 2020), the single-stranded DNA or single-stranded oligodeoxynucleotide (ssODN)-mediated HDR approach is widely used to perform precise genome editing at a desired locus. Several studies have been previously reported to achieve successful ssODN-mediated HDR in iPSCs by utilizing cell-cycle regulators (Lin et al., 2014; Yang et al., 2016), chemical inhibitors to suppress NHEJ (Ma et al., 2018; Riesenberg and Maricic, 2018; Yu et al., 2015), sib selection (Miyaoka et al., 2014, 2016), optimization of transfection conditions (Li et al., 2016; Okamoto et al., 2019), chemical modification of ssODN molecules (Carlson-Stevermer et al., 2017; Savic et al., 2018), and a stably integrated inducible SpCas9 expression vector (Bertero et al., 2016; Chen et al., 2015; Dow et al., 2015; Ishida et al., 2018), since high levels of SpCas9 expression in target cells are thought to be correlated with high levels of genome editing activity in general.

Recently, CRISPR-Cas9 ribonucleotide protein (RNP) delivery has been reported to achieve efficient indel induction and knockin in iPSCs (Dastidar et al., 2018; Kim et al., 2014; Liang et al., 2015; Martin et al., 2019; Xu et al., 2019), even though the Cas9 protein introduced into the cells is transient (typically less than 24 h). However, to the best of our knowledge, the underlying mechanisms as to why RNP transduction results in higher genome editing efficiency relative to the amount of Cas9 protein (Banan, 2020) are unknown. In addition, the ssODN-mediated knockin approach warrants further optimization and efficiency to allow more flexible and desired genome



editing outcomes, such as one-step biallelic modification and insertion of a *loxP* site, with less labor-intensive clonal isolation and genotyping processes.

In this study, we investigated intrinsic cellular factors that affect the editing efficiency of the CRISPR-Cas9/ sgRNA complex and found that cellular RNA inhibits Cas9 nuclease and sgRNA binding in vitro. Precomplexing Cas9 protein and sgRNA before delivery into cells was far more efficient at inducing indels compared with DNA plasmid-mediated delivery. Thus, we optimized an RNPbased electroporation approach in patient iPSCs with ssODNs in order to induce high levels of knockin exceeding 70%. When we compared two electroporation instruments, Lonza 4D-Nucleofector and MaxCyte, side by side, we found that the MaxCyte protocol showed a tendency toward higher knockin efficiencies under the conditions used. Furthermore, we could apply our approach to generate various iPSC lines by introducing a desired point mutation, by correcting a homozygous pathogenic mutation, or by inserting loxP sites without the need for antibiotic selection. Finally, by introducing two point mutations into the HLA-C gene, we generated a custom histocompatibility leukocyte antigen (HLA)-edited iPSC line that could evade immune surveillance from natural killer (NK) cells, suggesting that custom HLAmatching iPSCs can be generated by ssODN-mediated HDR.

RESULTS

Intracellular Cas9 Protein Levels Do Not Correlate with Genome Editing Outcomes

We first evaluated genomic DNA cleavage activity in HEK293T cells by comparing the transfection of various forms of Cas9 and sgRNA: lipofection of plasmid DNA vector, lipofection of Cas9 protein and in vitro transcribed (IVT) sgRNA, and stable expression of Cas9 and sgRNA from doxycycline (Dox)- and dexamethasone (Dex)-inducible CRONUS piggyBac vector (Ishida et al., 2018). Interestingly, we found that, although RNP transfection resulted in 13-fold lower levels of Cas9 protein compared with plasmid DNA (Figure 1A), the cleavage activity of RNP was 1.5-fold higher than that of plasmid DNA transfection by T7E1 analysis (Figure 1B). Furthermore, when RNP lipofection and the CRONUS piggyBac vector-mediated Cas9-inducible expression were compared, cleavage activity was comparable, even though the amount of Cas9 protein in cells was 20-fold more abundant using the CRONUS system. However, the high cleavage activity was compromised when recombinant Cas9 and IVT sgRNA were transfected separately ("RNP separate" in Figure 1B). From these observations, we hypothesized that Cas9 might be inactivated or inhibited by an intracellular molecule in mammalian cells.

Endogenous RNA Potentially Interferes with sgRNA Binding to Cas9

To search for cellular components that may interfere with Cas9 cleavage activity in cells, we performed an in vitro cleavage assay using a DNA fragment as a substrate for Cas9/gRNA cleavage. We initially hypothesized that an excess amount of genomic DNA might sequester Cas9 protein, but the addition of genomic DNA prior to sgRNA showed almost no inhibition (Figure S1). We also tested cellular protein fractions from cell lysates and observed no significant inhibition. Interestingly, when total cellular RNA from human cells was preincubated with Cas9 protein before the addition of IVT sgRNA (DMD1), DNA cleavage activity of Cas9 was inhibited in a dose-dependent manner, whereas no inhibition was observed when sgRNA was added prior to cellular RNA (Figure 1C). This suggested that total cellular RNA may have an inhibitory effect by preventing sgRNA from complexing with Cas9 protein. Preincubation with non-targeting sgRNA as an antagonist also strongly reduced the subsequent cleavage activity (Figure S1B), suggesting sgRNA switching is rare once sgRNA binds to Cas9 protein.

A time-course experiment revealed that Cas9 cleavage inhibition from total cellular RNA occurred immediately and was sustained. In contrast, genomic DNA exerted no apparent inhibitory effect, similar to the no-inhibitor negative control (Figure S1C).

Inhibition of Cas9 by Cellular RNA Is Reversible by RNase Treatment

To determine whether the inhibition effect by cellular RNA is reversible, we incubated Cas9 protein with inhibitory RNAs (cellular RNAs or antagonist sgRNA) and then performed RNase treatment to see if removal of the inhibitory RNA component from Cas9 would allow sgRNA to complex again and DNA cleavage to occur. After the addition of RNase inhibitor to protect sgRNA from RNase, we added sgRNA and proceeded with the *in vitro* cleavage assay (Figure 1D). We found that RNase treatment managed to reverse the inhibitory effect of cellular RNA as well as that of antagonist sgRNA. This confirms that oligomeric RNA, not monomeric nucleotides, is a key molecule of the inhibitory effect of Cas9. Interestingly, when we examined the various sizes of IVT RNA on Cas9 activity inhibition, larger RNA fragments showed similar inhibition activity compared with bulk cellular RNA (Figure S1D), suggesting that the inhibition effect is sequence independent, but size dependent.

We also performed an RNA pull-down assay to quantify the amount of sgRNA bound to Cas9 in the presence or absence of inhibitory cellular RNA by qRT-PCR. The results showed that the presence of inhibitory cellular RNA significantly (p = 0.004 by Welch's t test) reduced





Figure 1. Expressed Cas9 Protein in Cells Can Be Inhibited by Cellular RNA

(A) Intracellular Cas9 protein amount in HEK293T cells was measured by western blot after various transfection methods: transient transfection of plasmid DNA vector by Lipofectamine 2000 (Plasmid DNA, 48 h post-transfection), lipofection of Cas9 protein using Lipofectamine CRISPRMAX (4 h post-transfection) either premixed with sgRNA (RNP pre-mix) or transfected separately (RNP separate), or stably integrated CRONUS piggyBac vector with Dox/ Dex induction (CRONUS[+], 48 h post-induction) or residual expression (CRONUS [-]). Cas9 band intensities are represented as means \pm SD (n = 3 experiments). Oneway ANOVA followed by Dunnett's multiple comparisons test.

(B) Genome editing efficiency was measured by T7 endonuclease I (T7EI) assay after 48 h of transfection or induction in (A) at the human *DMD* gene locus. DNA cleavage percentages are represented as means \pm SD (n = 3 experiments). One-way ANOVA followed by Dunnett's multiple comparisons test.

(C) Cas9 protein (50 ng) was incubated for 15 min in vitro with total cellular RNA (1-4,000 ng) extracted from HEK293T cells before (red line) or after (green line) the addition of sgRNA (12.5 ng). DNA cleavage activity was assessed by the in vitro cleavage assay using TapeStation 2200. DNA cleavage percentages are represented as means \pm SD (n = 3 experiments). Two-way ANOVA followed by Sidak's multiple comparisons test. (D) To identify whether the inhibitory effect is dependent on oligomeric RNA and can be reversed, Cas9 protein was preincubated with cellular RNA for 15 min at room temperature. For one group, no RNase A was added (blue bars). For another group, RNase A was added to digest the inhibitory RNA and then RNase inhibitor so as not to interfere with sqRNA, which was added af-

terward (green bars). Anti-DMD sgRNA was used for target DNA cleavage, and anti-HLA sgRNA was used as a non-cleaving, antagonist sgRNA. In vitro DNA cleavage percentage values are presented as means \pm SD (n = 3, technical triplicate).

(E) RNA pull-down assay confirmed that cellular RNA interferes with sgRNA binding to Cas9. Recombinant Cas9 protein with a hemagglutinin (HA)-tag was treated with or without cellular RNA before the addition of sgRNA, and the amount of sgRNA was measured by qRT-PCR as input (blue bars). Then, Cas9 protein was precipitated by anti-HA tag antibody, RNA components bound to Cas9 protein were eluted, and the amount of sgRNA in the elution was measured by qRT-PCR (orange bars). sgRNA amounts relative to no cellular RNA samples are represented as means \pm SD (n = 3, technical triplicate).

See also Figure S1.

the amount of sgRNA binding to Cas9 by over 60% (Figure 1E). Taken together, these results supported our hypothesis that when Cas9 protein encounters cellular RNA prior to sgRNA, cellular RNA inhibits sgRNA binding to Cas9 for optimal cleavage activity. Our findings also indicate that direct transfection of Cas9 RNP



precomplexed with sgRNA is a preferable method to avoid inhibition by cellular RNA.

Improving RNP-Mediated ssODN Knockin Efficiency in iPSCs

Next, because iPSCs have more utility than HEK293T cells for disease modeling and regenerative medicine applications, we sought to optimize the RNP-mediated genome editing strategy in iPSCs in combination with ssODN donor templates to induce precise single-nucleotide editing. Previously, the ssODN knockin efficiency of several transfection methods, including Lipofectamine CRISPRMAX, Neon, and 4D-Nucleofector electroporation, has been reported (Li et al., 2016; Okamoto et al., 2019; Ran et al., 2013; Takayama et al., 2017). However, as there are limited reports using the MaxCyte electroporation platform for iPSC engineering, we optimized transfection conditions for CRISPR-Cas9 genome editing and ssODN knockin efficiency in human iPSCs.

We examined the electroporation efficiency of GFP mRNA in healthy donor-derived 1383D2 iPSCs transfected at various electroporation energy settings (E3, E6, and E8) using MaxCyte. One day after electroporation, fluorescence microscope imaging of electroporated iPSCs showed that nearly all of the cells were GFP positive, with higher GFP intensity corresponding with higher electroporation energy (Figure S2A). In line with these results, flow cytometry analysis confirmed that almost 100% of the cells were GFP positive and had an increasing GFP mean fluorescence intensity with increasing electroporation energy (Figure S2B). These results indicated that the MaxCyte electroporator can effectively transfect GFP mRNA into iPSCs. Next, we examined the indel percentage of 1383D2 iPSCs after electroporation of the Cas9/sgRNA complex targeting the human DMD (dystrophin) gene locus using the MaxCyte electroporator. T7E1 analysis revealed that increasing the electroporation energy also increased the percentage of indels up to 36% (Figure S2C).

Next, we selected MaxCyte condition E8 to examine the efficiency of ssODN-mediated knockin into Duchenne muscular dystrophy patient iPSCs (CiRA00111) with exon 44 deletion in the *DMD* gene. To evaluate HDR frequency, we electroporated various amounts of ssODN (DMD1) together with CRISPR RNP to disrupt the splicing acceptor and induce exon 45 skipping in the *DMD* gene. The ssODN knockin efficiency was checked by using the restriction enzyme AgeI, which is present only if ssODN-mediated HDR occurs (Figure S3A). As shown in Figure S3B, ssODN knockin was observed with as little as 1.5 µg ssODN and plateaued from 5 µg with over 60% knockin efficiency, whereas NHEJ indel efficiency was maintained regardless of ssODN amount (Figure S3C).

Optimization of ssODN-Mediated Knockin to Alter Two Nucleotides in iPSCs

We next examined the ssODN-mediated knockin efficiency by targeting the *ILF3* gene, also known as *NF110* or *NFAR2*. The *ILF3* gene encodes a dsRNA-binding protein involved in cellular host defense and is known to be phosphorylated by dsRNA-dependent protein kinase (Harashima et al., 2010). To introduce an amino acid substitution at S691C or G694A, we designed two 100-mer ssODNs to be targeted with the same sgRNA (NF110-Ex17-SA2). To assess the successful induction of the point mutations, we incorporated an additional silent mutation to introduce a blocking mutation and to generate a restriction enzyme site (BstUI or PstI), respectively (Figure S3D). With 13.3 µg ssODN, we found that MaxCyte gave better knockin efficiency than 4D-Nucleofector in iPSCs (1383D2) for both S691C (Figure S3E) and G694A (Figure S3F) substitution experiments.

One-Step Homozygous Correction of the *DYSF* Gene in Miyoshi Myopathy Patient iPSCs

To demonstrate efficient and precise genome editing, we compared the editing of a disease-associated single-nucleotide mutation with the 4D-Nucleofector and MaxCyte instruments side by side. In this experiment, we utilized the iPSC line CiRA00396 derived from a patient with Miyoshi myopathy (a form of dysferlinopathy), who has a homozygous recessive nonsense mutation of c.C3166T (p.Arg1056Ter) at exon 29 in the Dysferlin (DYSF) gene (Liu et al., 1998). Miyoshi myopathy patient iPSCs were electroporated with 6.7 or 13.3 µg ssODN, together with the Cas9/sgRNA complex. After 3 to 7 days, the iPSCs were harvested and analyzed for knockin efficiency by Hpy99I restriction enzyme digestion, which was introduced into the sgRNA targeted site upon successful ssODN knockin (Figure 2A). Although 4D-Nucleofector-mediated electroporation showed a high knockin efficiency of 46% with 6.7 µg ssODN, higher amounts of ssODN resulted in overall lower knockin efficiency. On the other hand, the MaxCyte-mediated electroporation resulted in 67% and 73% knockin efficiency at 6.7 and 13.3 µg ssODN, respectively (Figure 2B).

To obtain single edited clones, we subcloned iPSCs electroporated with CRISPR RNP and 13.3 μ g ssODN using the MaxCyte instrument. Single cells were sorted and expanded, and 45 clones were examined for ssODN knockin efficiency by Sanger sequencing. Astonishingly, 73.3% (n = 33) of the clones were homozygous for the ssODN knockin, while 20% (n = 9) were heterozygous knockin and 6.7% (n = 3) showed no evidence of ssODN knockin (Figure 2C). For the heterozygous clones (n = 9) with one corrected allele by the ssODN knockin, eight also contained indels on the other allele, and the last clone contained an unedited allele. Among the three clones without ssODN knockin, two contained indels in both





Figure 2. Correction of a Homozygous Mutation in Miyoshi Myopathy Patient iPSCs

(A) A patient with Miyoshi myopathy (dysferlinopathy) has a homozygous nonsense mutation (c.C3166T, p.Arg1056Ter, in isoform 1) at exon 29 of the *DYSF* gene on chromosome 2, which results in the loss of dysferlin protein expression. Through ssODNmediated knockin genome editing, we attempted to correct the nonsense mutation. Successful correction can be detected by the appearance of the Hpy99I restriction enzyme site (CGWCG, W = A or T).

(B) Quantitative result of restriction enzyme digestion with Hpy99I enzyme to assess the bulk knockin efficiency in Miyoshi myopathy iPSCs (CiRA00396 clone). Digested PCR samples were analyzed by TapeStation. A PCR product from healthy 1383D2 iPSCs was used as a positive control for Hpy99I cleavage. Cleavage percentages are represented as means \pm SD (n = 3, technical triplicate).

(C) After subclone isolation from MaxCyte electroporation of the ssODN template, we genotyped each subclone of Miyoshi myopathy iPSCs by Sanger sequencing. The pie chart shows the proportion of homozygously corrected subclones (green) either with or without indels. The sequences of 45 iPSC clones in total were analyzed.

(D) Cell viability assay was performed 1 day after electroporation with 4D-Nucleofector or MaxCyte by using Cell Counting Kit-8. Cell viability values relative to non-transfected sample are represented as means \pm SD (n = 3, technical triplicate).

See also Figures S2-S4.

alleles and the other was unedited in both alleles. Because CRISPR-Cas9 genomic DNA cleavage may induce large deletions at the target site (Mou et al., 2017; Weisheit et al., 2020), we performed additional PCR analysis of four randomly selected homozygous knockin clones (nos. 2, 10, 12, 32) and one heterozygous knockin clone (no. 6) (Figure S4A). As shown in Figure S4B, we observed no large deletions around the targeted DYSF gene site of the analyzed clones. In addition, to check whether the Max-Cyte genome-edited iPSCs maintained pluripotency, we checked the expression of the pluripotent markers SSEA-4 and TRA-1-60 by flow cytometry. All five subclones maintained expression of these markers comparable to the parental iPSC clone (Figure S4C). These results confirmed that RNP- and ssODN-mediated knockin using MaxCyte electroporation enabled us to obtain homozygously edited clones at very high efficiency. To explain why ssODN

knockin efficiency with MaxCyte was higher than with 4D-Nucleofector, we investigated the relative viable cell number 1 day after electroporation. The results showed that transfection using MaxCyte had better survival compared with 4D-Nucleofector, especially after introduction of the ssODN (Figure 2D).

Exon Skipping in Duchenne Muscular Dystrophy Patient iPSCs

We also analyzed ssODN knockin efficiency by targeting the exon 45 splicing acceptor site of the *DMD* gene in CiRA00111 iPSCs (Figure 3A), which do not express DMD protein due to a deletion of exon 44 and can be reframed by exon 45 skipping (Gee et al., 2020; Li et al., 2015; Ousterout et al., 2015). We transfected precomplexed Cas9/ sgRNA RNP with 6.7 μ g ssODN by using 4D-Nucleofector and MaxCyte. After 3 to 7 days, confluent iPSCs were



Figure 3. Disruption of a Splicing Acceptor on the *DMD* Gene

(A) To induce therapeutic exon skipping by disrupting a splicing acceptor site on exon 45 of the *DMD* gene, we designed an sgRNA and ssODN template to introduce two nucleotide alterations. Successful knockin can be evaluated by the appearance of the AgeI restriction enzyme site (A|CCGGT).

(B) We electroporated precomplexed Cas9/ sgRNA together with ssODN to edit the *DMD* gene locus in DMD patient iPSCs by using either 4D-Nucleofector or MaxCyte. Three to five days after electroporation, we evaluated the NHEJ-mediated indel frequency by T7EI assay. Data are represented as means \pm SD (n = 3, technical triplicate).

(C) From the same genomic DNA sample in (B), we assessed ssODN-mediated HDR efficiency by Sanger sequencing and Sequencher software. Data are represented as means \pm SD (n = 3, technical triplicate).

(D) A cell counting assay was performed 1 day after electroporation by 4D-Nucleofector or MaxCyte by using Cell Counting Kit-8. Relative cell viability values are represented as means \pm SD (n = 3, technical triplicate).

harvested, NHEJ-mediated indel efficiency was evaluated by T7E1 assay, and HDR-mediated precise base alteration was assessed by restriction fragment-length polymorphism analysis. T7E1 assay results showed that NHEJ-mediated cleavage activity was comparable between 4D-Nucleofector and MaxCyte (Figure 3B); however, MaxCyte showed more consistent HDR-mediated knockin efficiency (Figure 3C). When we measured cell survival 1 day after electroporation, results similar to those in Figure 2D were observed by targeting the *DMD* gene locus (Figure 3D). The addition of ssODN caused high cell toxicity, but MaxCyte electroporation attenuated this cytotoxicity in iPSCs.

High Efficiency of a *loxP* Insertion in iPSCs without Antibiotic Selection

The Cre-*loxP* system has been widely utilized for custom deletion or targeted integration in the genetics field,

especially for mouse genome manipulation (Miura et al., 2018). However, insertion of a *loxP* sequence in human iPSCs has been challenging because of very low efficiency; hence, antibiotic selection has been necessary (Chen et al., 2015; Yang et al., 2017). To elucidate whether loxP knockin by an ssODN template without antibiotic selection is possible in iPSCs, we sequentially introduced two *loxP* sites into the dystrophin gene on chromosome X to generate a floxed allele (Figure 4A). First, we transfected RNP with the 134 bp ssODN, which contains a 34 bp loxP site and 50 bp homology arms on each side, to target the exon 45 region in male 1383D2 iPSCs using MaxCyte. The first ssODN-mediated loxP knockin was checked by PCR and restriction digestion with XmnI (Figure S5A). We could obtain >20% loxP insertion efficiency (Figure S5B). As above, we performed subcloning by single-cell sorting and examined the ssODN knockin efficiency by Sanger





of the 342 kb region between exon 44 and exon 55 in the dystrophin gene. See also Figure S5.

sequencing. Of the 20 clones we analyzed, eight (40%) showed insertion of the intact loxP sequence at the intended site, eight (40%) were accompanied by indels or NHEJ only, and four (20%) were unedited (Figures 4B and S5C). We selected one subclone (no. 10) into which the first loxP was inserted (Figure S5D) and inserted a second loxP site after exon 55 of the dystrophin gene (342 kb distal to the first loxP). Of the 14 clones analyzed, two (2/14 =14.3%) contained the intact *loxP* site successfully inserted (Figures 4C and S5E). We selected the two clones (nos. 10-5 and 10-7) containing both loxP sites and transfected them with a Cre expression plasmid to excise the 342 kb region (Figure 4D). Bulk PCR was performed to amplify the junction region, and it was analyzed by Sanger sequencing to confirm precise excision between the two loxP sites (Figure 4E). Of the total 11 clones analyzed, four showed removal of the 342 kb region flanked by the two *loxP* sites (Figure S5F), indicating the function of the two *loxP* sites. These results demonstrate that our electroporation method of precomplexed RNP and ssODN could be applied to introduce a 34 bp loxP site without the need for antibiotic selection.

Figure 4. Insertion of Two *loxP* Sequences to Induce a Large Deletion in iPSCs

(A) Schematic of the *loxP* insertion sites in the dystrophin gene on chromosome X. Target sequence of sgRNAs and *loxP* insertion sites are indicated.

(B) After MaxCyte electroporation of Cas9 RNP, sgRNA#1, and ssODN (DMD1+*loxP*) into male iPSCs (1383D2), 20 subclones were isolated and genotyped by Sanger sequence using a reverse primer. The intact *loxP* sequence is indicated by the green marker and Cre-binding sequences in the *loxP* site are indicated by underlining.

(C) A second *loxP* site was inserted into clone 10 from (B) by MaxCyte electroporation of Cas9 RNP, sgRNA in55-3, and ssODN (DMD-in55-g3+*loxP*). Of the 14 clones analyzed by Sanger sequencing, two (nos. 10-5, 10-7) showed successful insertion of the second *loxP* site.

(D) Schematic of the Cre recombination to remove a 342 kb region on the dystrophin gene flanked by the two *loxP* sites.

(E) After transfection of the tamoxifeninducible Cre expression plasmid vector into clone 10-5 or 10-7, genomic PCR was performed to amplify the junction of the two *loxP* sites. Sanger sequencing of the PCR product confirmed successful deletion

ssODN-Mediated One-Step Biallelic Modification to Convert HLA-C1 to HLA-C2

To facilitate HLA matching for allogeneic transplantation of iPSC derivatives, the use of HLA homozygous iPSC stocks has been investigated (Sugita et al., 2020). However, HLA homozygous cells may be susceptible to attack by HLA-C1/C2 heterozygous recipient NK cells due to KIR mismatching (Ichise et al., 2017), since NK cells express KIR 2DL2/2DL3 and KIR 2DL1 receptors, which suppress HLA-C1 and HLA-C2 groups, respectively (Figure 5A). The difference between HLA-C1 (i.e., HLA-Cw1, Cw7, Cw8, Cw9) and HLA-C2 (i.e., HLA-Cw2, Cw4, Cw5, Cw6) is determined by two amino acids at the 77th and 80th positions of HLA-C protein; the HLA-C1 group has serine at position 77 and asparagine at position 80, while the HLA-C2 group has asparagine and lysine instead (Moesta et al., 2008). To avoid the HLA-C/KIR mismatch, we sought to generate HLA-C1/C2-heterozygous HLA-C alleles from HLA-C1/C1 homozygous iPSCs (Figure 5A).

To accomplish this, we chose a clinical-grade HLA homozygous iPSC line, Ff-XT28s05, which has homozygous HLA-C1 alleles (HLA-Cw7). We first designed two ssODNs;



Figure 5. Simultaneous **Biallelic** Replacement to Generate an HLA-C1/C2 Haplotype in HLA-C1/C1 Homozygous iPSCs

(A) NK cells in HLA-C1/C2 heterozygous donors express KIR 2DL3 and KIR 2DL1 receptors, which are suppressor receptors of HLA-C1 and HLA-C2 groups, respectively. However, when cells from HLA-C1/C1 homozygous iPSCs are transplanted, these NK cells pose a risk of attacking the transplanted iPSC derivatives because of the absent inhibitory signal from the HLA-C2/ KIR 2DL1 axis. Therefore, we replaced one allele of HLA-C1 with a C2-like allele to suppress KIR 2DL1.

(B) HLA-C1 and C2 groups differ by two amino acids at positions 77 and 80. We designed two ssODNs, one containing two nucleotide mutations to enable conversion from HLA-C1 to C2, and the other containing a silent mutation to prevent Cas9 from recutting the HLA-C1 allele.

(C) Ff-XT28s05 iPSCs were established from

a healthy HLA-C1 homozygous donor. After electroporation of precomplexed Cas9/sqRNA with the two ssODNs by MaxCyte, 29 subclones were analyzed, and one (no. 25) showed biallelic knockin patterns by Sanger sequencing (R = A or G, Y = C or T, S = C or G). (D) To sequence individual alleles separately, we performed TA cloning of the PCR amplicon from clone 25 and analyzed by Sanger sequencing.

one, ssODN2, contains two mutations to enable changing C1 to C2, while the other, ssODN1, contains a silent mutation to prevent Cas9 from recutting the targeted HLA-C1 locus (Figure 5B). Cas9 and sgRNA with two ssODNs were electroporated, and the knockin efficiency of each allele was 24% and 28%, respectively. Of the 29 subclones we analyzed, we could establish a clone (no. 25) that had all three intended mutations in both alleles, which we confirmed by bulk Sanger sequencing (Figure 5C) and TA cloning (Figure 5D).

DISCUSSION

While investigating various delivery methods (plasmid DNA vector, RNP, and Dox-inducible piggyBac vector) of the CRIPSR-Cas9/sgRNA system, we observed that exogenous plasmid DNA and an integrated piggyBac vector could mediate high levels of Cas9 protein expression in cells, but genome editing efficiency was comparable to or even lower than RNP transfection, which resulted in lower amounts of Cas9 protein in cells. To investigate the underlying mechanistic reasons behind this phenomenon, we found that free Cas9 protein (or apo-Cas9) could be inhibited by cellular RNA for binding with sgRNAs in vitro, which dramatically reduced Cas9's DNA cleavage efficiency. This effect suggests that Cas9 is susceptible to inhibition by intracellular RNAs

when Cas9 protein is expressed. On the other hand, Cas9 protein precomplexed with sgRNA in vitro is less susceptible to inhibition by intracellular RNAs. Hence, precomplexed RNP transfection is preferable for achieving high genome editing efficiency compared with expression-based delivery systems.

Recently, several anti-CRISPR proteins (AcrIIs) have been identified in bacteriophages that specifically inhibit type II CRISPR-Cas9 activities (Fuchsbauer et al., 2019; Harrington et al., 2017; Jiang et al., 2019). For instance, AcrIIA1-AcrIIA6 binds to Cas9 and inhibits DNA binding in an allosteric manner. AcrIIC3 also binds to Neisseria meningitidis Cas9 (NmCas9) for blocking the DNA loading step (Zhu et al., 2019). Other anti-CRISPR proteins, such as AcrIIA11 and AcrIIC1, prevent DNA cleavage. To date, only AcrIIC2 has been identified as an anti-CRISPR to prevent sgRNA complexing with NmCas9 (Zhu et al., 2019). Although it is unclear how intracellular RNA inhibits Cas9 for sgRNA loading at the molecular level, we speculate that cellular RNA binds to cationic Cas9 protein (pI 9.01) non-specifically and causes steric hindrance or physically obstructs sgRNA from getting into close proximity to apo-Cas9 (Lim et al., 2016).

In our experiments, up to 4,000 ng RNA was used per 10-20 µL reaction volume; hence, the maximum RNA concentration was 0.4 mg/µL. On the other hand, the cellular RNA concentration is roughly 10-30 mg/µL based on estimating



the amount of total cellular RNA per human cell (10–30 pg, BNID 111205) and the cellular volume (4 × 10^{-6} µL, BNID 105906). Therefore, the physiological concentration of intracellular RNA is one or two orders greater than the concentration range used in our experiments.

Other possible implications of our findings are the importance of Cas9 protein purification when extracted from a crude lysate (Qiao et al., 2019), as contaminated bacterial RNA may potentially compromise Cas9 genome editing activity or lead to a lot-to-lot variation in enzymatic activity. Notably, as shown in Figure 1D, the inhibition of sgRNA loading can be released by RNase treatment. Recently, RNP electroporation has become a standard genome editing method, owing to its simplicity and high efficiency. Here, we provide a plausible explanation for why precomplexed RNP offers higher genome editing specificity compared with other expression methods.

To further optimize RNP-mediated genome editing, we compared electroporation instruments. Although the Lonza 4D-Nucleofector has been widely used (Skarnes et al., 2019), we demonstrated that the MaxCyte electroporation platform is also a suitable instrument for genome editing applications. When we compared the 4D-Nucleofector and MaxCyte instruments side by side, we found that knockin efficiency with ssODN was higher with the MaxCyte enabled predominantly homozygous knockin efficiencies in as many as 72% of subclones at the *DYSF* gene locus in Miyoshi myopathy patient iPSCs (Figure 2).

To promote the HDR pathway over NHEJ, the availability of the donor template at the vicinity of a DSB site is critical. Transfecting a high concentration of ssODN is advantageous to increase the chance for the ssODN to reach the nucleus and be in close proximity to the Cas9 cleavage site; however, ssODN itself is toxic for cells, as shown in Figures 2D and 3D. Because the HDR DNA repair pathway mainly takes place at S or G2 phase of the cell cycle, we speculate that actively proliferating cells should have a higher HDR rate than damaged arrested cells. The different performances of the two electroporation instruments were accompanied by a difference in cell viability, suggesting this parameter could affect the HDR frequency.

Applying our optimized and robust ssODN knockin approach to engineering human iPSCs, we achieved insertion of a 34 bp *loxP* sequence into a desired locus in up to 40% of cells, which can be used as a hub for site-specific integration. Furthermore, we demonstrated that two *loxP* sites could be inserted into a target gene locus for introducing a floxed allele. Upon the introduction of Cre recombinase, a precise and large deletion of 342 kb could be introduced in iPSCs.

We also investigated the knockin of two unique ssODNs into both alleles, which enabled us to generate an HLA-C1/

C2 heterozygous allele from HLA-C1/C1 homozygous iPSCs. Engineered HLA-C1/C2 iPSCs should be able to suppress NK cells from HLA-C1/C2 donors via the KIR 2DL1 receptor. Taken together, our study would facilitate custom editing of a desired genomic locus with precomplexed RNP and ssODN templates in iPSCs for genetic research and regenerative medicine applications.

EXPERIMENTAL PROCEDURES

Cell Culture

HEK293T cells (ATCC, cat. no. CRL-3216) and CRONUS-HEK293T cells were cultured in DMEM (Nacalai Tesque, cat. no. 08459-64) with 10% FBS (Biosera, cat. no. FB-1365). Feeder-free iPSCs were maintained in StemFit AK03N (Ajinomoto) in a laminin 511 E8 (iMatrix-511 E8, cat. no. 892012, Nippi)-coated six well plate (BD, cat. no. 353046). TrypLE Select (Thermo Fisher Scientific, cat. no. 12563-011) was used for passaging and harvesting cells. 1383D2 iPSCs (donor: healthy Asian male, 36 years of age at the time of donation) were established by Dr. Masato Nakagawa (Nakagawa et al., 2014) and provided by FiT (Facility for iPSC Therapy) at CiRA. DMD-iPSCs (CiRA00111 clone) from a patient with Duchenne muscular dystrophy with exon 44 deletion were described previously (Li et al., 2015). Miyoshi myopathy patient iPSCs (CiRA00396 clone) derived from a patient with a homozygous nonsense mutation (exon 29; c.C3166T, p.Arg1056Ter) were a kind gift from Dr. Hidetoshi Sakurai (CiRA). The protocol to establish the patient-derived iPSC lines was approved by the ethics committee of the Graduate School and Faculty of Medicine, Kyoto University (approval nos. R0091 and G259). Ff-XT28s05 iPSCs were generated from a healthy donor homozygous for the most frequent HLA haplotypes in Japan (A*24:02, B*07:02, C*07:02, DRB1*01:01) under a clinical-grade cell manufacturing protocol at FiT in CiRA, Kyoto University.

Electroporation of RNP by MaxCyte or 4D-Nucleofector

The day before electroporation, iPSCs were seeded onto a six well plate coated with iMatrix-511 at a density of $1.5-2 \times 10^6$ cells per well. On the day of transfection, iPSCs were washed with 2 mL PBS and then incubated with 0.5 mL Accutase or TrypLE Select for 10 min at 37°C. The cells were then detached from the plate by pipetting, transferred to a 15 mL centrifuge tube containing AK03N with 10 μ M ROCK inhibitor (Y-27632, Wako, cat. no 253-00514), and centrifuged for 5 min at 120 × g to harvest the cells by discarding the supernatant.

For MaxCyte electroporation, the cells were resuspended in 1 mL MaxCyte buffer. The cells were counted, washed with 5 mL MaxCyte buffer, and resuspended in MaxCyte buffer at a density of 2.5×10^7 cells/mL. For RNP electroporation, 10 µg recombinant SpCas9 protein (IDT, cat. no. 1081058, or Thermo Fisher, cat. no. A36498) and 2.5 µg IVT sgRNA were incubated for 5 min at room temperature, and then 1.25×10^6 cells with 50 µL MaxCyte buffer and 0.5 to 25 µg ssODN (Fasmac, Table S3) were added and mixed gently before transfer to an OC-100 cuvette. For GFP mRNA electroporation, 5 µg mRNA (TriLink) was used. Electroporation was carried out on the MaxCyte ATX



or STX at the indicated electroporation energy. Immediately after electroporation, the OC-100 cuvettes with iPSCs were incubated at 37°C for 20 min in a humidified incubator to allow for cell membrane recovery. Finally, cells were added to a single well of an iMatrix-511-coated six well plate containing 2 mL prewarmed Stem Fit AK03N medium with 10 μ M Y-27632. Semi-confluent cells (3–7 days after transfection) were harvested for genomic DNA extraction and/or FACS analysis.

For 4D-Nucleofector electroporation, the cells were resuspended in StemFit AK03N medium and cell counting was performed. After centrifugation, 3×10^5 cells were gently resuspended in 20 µL P4 Primary Cell Nucleofector Solution from a P4 Primary Cell 4D-Nucleofector Kit (Lonza, cat. no. V4XP-4032). For RNP electroporation, 5 µg recombinant SpCas9 protein (IDT, cat. no. 1081058, or Thermo Fisher, cat. no. A36498) and 1.25 µg IVT sgRNA were incubated for 5 min at room temperature and then added to the cell suspension with or without ssODN (ranging from 0.5 to 25 µg). The cells were gently mixed by pipetting five times and then transferred to a well of a 16 well Nucleocuvette Strip. Immediately after electroporation using the CA-137 protocol of 4D-Nucleofector, the cells were transferred to an iMatrix-511-coated well of a six well plate containing 2 mL prewarmed StemFit AK03N medium with 10 µM Y-27632. Semi-confluent cells (3-7 days post-transfection) were harvested for genomic DNA extraction and/or FACS analysis as described above.

loxP Insertion by Cas9/sgRNA and ssODN

Electroporation

iPSCs (1383D2 clone) were genome edited as described in the Max-Cyte electroporation section. For 1.0×10^6 cells with 50 µL Max-Cyte buffer, 5 µg Cas9 protein (Thermo Fisher, cat. no. A36498), 1.25 µg IVT sgRNA (either DMD-sgRNA1 or DMD-in55-g3), and 6 µg ssODN with loxP sequence (134-mer, either "DMD1+loxPssODN" or "DMD-in55-g3+loxP-ssODN," see Table S4) were electroporated using MaxCyte with Optimization Energy 8. After confirmation of the loxP knockin efficiency in bulk-electroporated cells, single-cell clones were isolated by using a BD FACSAria cell sorter (BD Biosciences) or by limited dilution. For the Cre-loxP excision experiment, floxed subclones (nos. 10-5 and 10-7) were seeded into a 24-well plate (1.5-3.0 × 10⁶ cells/well) and transfected with 500 ng tamoxifen-inducible Cre-expressing plasmid pCAG-IP-MerCreMer (a kind gift from Dr. Makoto Tachibana, Osaka University) with 1.5 µL Lipofectamine Stem Transfection Reagent (Thermo Fisher, cat. no. STEM00015). Four to six hours after the lipofection, the culture medium was replaced with 1 µM 4-hydroxytamoxifen (Sigma-Aldrich, cat. no. H7904) to induce nuclear translocation of Cre-recombinase fused with modified estrogen receptor (Mer). After confirmation of the excision of the floxed site by genomic PCR, two subclones from clone 10-5 and nine subclones from clone 10-7 were established and genotyped by PCR with "DMDexon55(45-55)check_dir5" and "DMDexon45(45-55) check_rev5" primers (Table S1).

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/ 10.1016/j.stemcr.2021.02.013.

AUTHOR CONTRIBUTIONS

Conceptualization, A.K., M.L., H.X., P.G., and A.H.; investigation, A.K., M.L., H.X., T.I., M.O., N.S., Y.K., X.W., and P.G.; writing – original draft, A.K., M.L., P.G., and H.X.; writing – review & editing, P.G. and A.H.; supervision, P.G. and A.H.; funding acquisition, A.H.

CONFLICTS OF INTEREST

P.G. is currently an employee of MaxCyte, Inc. All other authors declare no competing interests.

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