Ultra-superoovulation for the CRISPR-Cas9-mediated production of gene-knockout, single-amino-acid-substituted, and floxed mice

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ABSTRACT

Current advances in producing genetically modified mice using genome-editing technologies have indicated the need for improvement of limiting factors including zygote collection for microinjection and their cryopreservation. Recently, we developed a novel superovulation technique using inhibin antiserum and equine chorionic gonadotropin to promote follicle growth. This method enabled the increased production of fertilized oocytes via in vitro fertilization compared with the conventional superovulation method. Here, we verify that the ultra-superoovulation technique can be used for the efficient generation of clustered regularly interspaced short palindromic repeats (CRISPR)-CRISPR-associated protein 9 (Cas9)-mediated knockout mice by microinjection of plasmid vector or ribonucleoprotein into zygotes. We also investigated whether single-amino-acid-substituted mice and conditional knockout mice could be generated. Founder mice bearing base substitutions were generated more efficiently by co-microinjection of Cas9 protein, a guide RNA and single-stranded oligodeoxynucleotide (ssODN) than by plasmid microinjection with ssODN. The conditional allele was successfully introduced by the one-step insertion of an ssODN designed to carry an exon flanked by two loxP sequences and homology arms using a double-cut CRISPR-Cas9 strategy. Our study presents a useful method for the CRISPR-Cas9-based generation of genetically modified mice from the viewpoints of animal welfare and work efficiency.

KEY WORDS: Knockout, Floxed, CRISPR-Cas9, Zygote, Ultra-superoovulation, Inhibin antiserum and equine chorionic gonadotropin (IASe)

INTRODUCTION

Reproductive engineering techniques are essential for the efficient production and maintenance of genetically modified (GM) mice. Fertilized oocytes generated by mating or in vitro fertilization (IVF) are used for microinjection with or without cryopreservation to generate genome-edited mice. We have previously shown that IVF and freeze-thawing techniques efficiently generated GM mice using genome-editing technology (Nakagawa et al., 2014, 2015). The use of freeze-thawed fertilized oocytes allows flexible scheduling by allowing the required number of fertilized oocytes to be utilized at any time.

Use of the superovulation technique using equine chorionic gonadotropin (eCG) and human chorionic gonadotropin (hCG) is important to collect as many oocytes as possible from female mice. To further increase the yield of oocytes, we recently developed a new ultra-superoovulation technique using inhibin antiserum (IAS) and eCG (IASe) to promote follicle growth (Takeo and Nakagata, 2015). We have shown that approximately threefold numbers of oocytes could be collected from C57BL/6 female mice at 4 weeks of age by the administration of IASe compared with the conventional administration of IAS or eCG alone. We and other investigators have confirmed that fertilized oocytes created by IVF via the conventional superovulation method were applicable for creating genome-edited mice (Nakagawa et al., 2014, 2015; Li et al., 2014; Miura et al., 2015; Tsuchiya et al., 2015; Kaneko and Mashimo, 2015; Nakao et al., 2016; Sakurai et al., 2016). However, it has not yet been elucidated whether the ultra-superoovulation technique can be applied for the generation of genome-edited mice.

Recently, various genome-edited mice have been quickly generated using the microinjection technique and the clustered regularly interspaced short palindromic repeats (CRISPR)-CRISPR-associated protein 9 (Cas9) system (Singh et al., 2015). The CRISPR-Cas9 system induces DNA double-strand breaks (DSBs) by forming a complex of Cas9 and single guide RNA (gRNA) that targets a desired genomic locus. The DSBs are mainly repaired by error-prone non-homologous end-joining, which introduces random insertions and deletions of nucleotides around the target site, resulting in targeted gene disruption. Alternatively, DSBs can also be repaired using homology-directed repair (HDR) by co-injection of donor DNA harboring homology arms. Using genome editing technology and this repair system, precisely defined modifications can be efficiently introduced into the targeted locus. The CRISPR-Cas9 system and single-stranded oligodeoxynucleotide (ssODN) carrying objective arrangements with small modifications, including point mutations, multiple base substitutions, and insertion of tag sequences such as loxP, have been often used for the generation of genome-edited mice (Wang et al., 2013, 2015; Yang et al., 2013; Mizuno et al., 2014; Inui et al., 2014; Han et al., 2015; Hashimoto and Takemoto, 2015; Qin et al., 2015; Kaneko and Mashimo, 2015).

Here, we report the CRISPR-Cas9-mediated generation of various GM mice, such as gene-disrupted mice, single-amino-
Results

Generation of gene-knockout mice using freeze-thawed fertilized oocytes created by IVF using an ultra-superovulation method

To examine whether freeze-thawed fertilized oocytes created by IVF using an ultra-superovulation method can be applied to the CRISPR-Cas9-mediated generation of knockout mice, the birth and mutation rates of these oocytes injected with CRISPR-Cas9 plasmid vectors were compared with oocytes previously created by IVF using the conventional superovulation method.

First, we generated fertilized oocytes by IVF using sexually matured C57BL/6 female and male mice. Female mice were treated by the ultra-superovulation method using IASe to promote follicle growth and hCG to induce ovulation. Consistent with our previous report, we collected greater numbers of oocytes compared with the conventional superovulation method (Table S1). The fertilized oocytes were cryopreserved as the stock for microinjection. Next, previously constructed and validated CRISPR-Cas9 nuclease vectors or a FokI-dCas9 vector to target the interleukin-11 (Il11) gene (Fig. 2) (Nakagawa et al., 2015) were microinjected into the pronucleus of freeze-thawed fertilized oocytes generated using ultra-superovulated sexually mature mice. Then, the surviving oocytes were transferred to pseudopregnant mice and the resultant pups were obtained. Tail lysates from pups were used for polymerase chain reaction (PCR) to amplify the targeted sequence of Il11. Each PCR product was analyzed by direct sequencing. In the previous report, we investigated the birth rate and the mutation rate of zygotes injected with these vectors. Microinjection of Cas9 nuclease vectors resulted in relatively low birth rate and high mutation rate, whereas microinjection of FokI-dCas9 vector resulted in moderate birth and mutation rates (Nakagawa et al., 2015). The birth and mutation rates of zygotes generated using the ultra-superovulation method were similar to those created by the conventional method (Table 1). To examine birth and mutation rates using zygotes from sexually immature female mice, from which greater numbers of fertilized oocytes can be collected compared with mature females, we performed microinjection and transfer using the same method. Although we confirmed that gene-knockout mice could also be created using ultra-superovulated sexually immature female mice, the birth rate was lower than using oocytes from mature females (Table 1). To decrease the toxicity, we subsequently tested microinjection of Cas9 protein and synthesized gRNA_B instead of the plasmid vector (Nuclease_B) into the zygotes generated from immature female mice. This alteration resulted in the successful production of mutant pups at the Nuclease_B locus, although no pups were born by plasmid microinjection, suggesting that the birth rate of genome-edited mice can be improved by replacing CRISPR-Cas9 vectors with Cas9 ribonucleoproteins (RNPs) (Table 1). Collectively, our ultra-superovulation technique was confirmed to be applicable to the production of knockout mice using genome-editing technology.

Generation of single-amino-acid-substituted mice

To investigate further whether gene knock-in mice can be generated using freeze-thawed zygotes created via ultra-superovulation and IVF, as well as the production of gene-knockout mice, we aimed to generate founder mice harboring a three-base substitution in the secreted phosphoprotein 1 (Spp1) gene. The three-base substitution in Spp1 gene was designed to encode thrombin cleavage-

METHODS & TECHNIQUES


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incompetent mutant osteopontin protein (Nishimichi et al., 2011). To knock-in these substitutions precisely, we designed a gRNA on the target locus as well as the ssODN carrying the intended substitutions (Fig. 3A).

We initially tested the microinjection of CRISPR-Cas9 nuclease vector and ssODN, followed by single blastocyst assays for the validation of gene knock-in. The surviving zygotes were cultured for 3.5 days, and restriction fragment length polymorphism (RFLP) and direct sequencing analyses were conducted. We confirmed correct three-base substitution without any insertion and deletion (indel) mutations in one blastocyst (Fig. S1, Table S2). To generate the founder mice, we subsequently performed microinjection of CRISPR-Cas9 nuclease vector and ssODN, and then surviving zygotes were transferred to pseudopregnant mice. We used the oocytes collected from immature or mature female mice treated with ultra-superovulation for IVF. The birth rates were low and no pups with the three-base substitution were detected by RFLP analysis using BsmFI (Table 2). Therefore, to improve the birth rate and the knock-in rate, Cas9 RNPs were injected with ssODN into zygotes instead of the CRISPR-Cas9 nuclease vector, similar to the method used for Il11 targeting. By using Cas9 RNPs, both the birth rate and the knock-in rate were significantly improved (Table 2). The precise knock-in of the three-base substitution was confirmed in three pups by RFLP analysis and direct sequencing analyses. We confirmed the cleavage activity of these two gRNAs. We microinjected the left gRNA in rates by using Cas9 RNPs.

**Table 1. Generation of Il11 mutants using freeze-thawed fertilized oocytes created via IVF**

<table>
<thead>
<tr>
<th>Superovulation</th>
<th>Age in weeks</th>
<th>Reagent</th>
<th>Injected</th>
<th>Survived (%)</th>
<th>Transferred</th>
<th>Pups (%)</th>
<th>Mutants (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional method*</td>
<td>8–12</td>
<td>5 ng/µl nuclease_A plasmid</td>
<td>32</td>
<td>25 (78.1)</td>
<td>25</td>
<td>3 (12.0)</td>
<td>3 (100)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 ng/µl nuclease_B plasmid</td>
<td>34</td>
<td>22 (64.7)</td>
<td>22</td>
<td>2 (9.1)</td>
<td>2 (100)</td>
</tr>
<tr>
<td>Ultra-superoxovulation method</td>
<td>10–13</td>
<td>5 ng/µl FokI-dCas9 plasmid</td>
<td>39</td>
<td>35 (89.7)</td>
<td>34</td>
<td>6 (17.6)</td>
<td>2 (33.3)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 ng/µl nuclease_A plasmid</td>
<td>90</td>
<td>80 (88.9)</td>
<td>40</td>
<td>4 (10.0)</td>
<td>2 (50.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 ng/µl nuclease_B plasmid</td>
<td>133</td>
<td>127 (95.6)</td>
<td>85</td>
<td>6 (7.1)</td>
<td>5 (83.3)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 ng/µl FokI-dCas9 plasmid</td>
<td>49</td>
<td>43 (87.8)</td>
<td>43</td>
<td>7 (16.3)</td>
<td>5 (71.4)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 ng/µl nuclease_A plasmid</td>
<td>112</td>
<td>104 (92.8)</td>
<td>53</td>
<td>3 (5.7)</td>
<td>3 (100)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 ng/µl nuclease_B plasmid</td>
<td>64</td>
<td>62 (96.9)</td>
<td>62</td>
<td>0 (0)</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 ng/µl FokI-dCas9 plasmid</td>
<td>68</td>
<td>63 (92.6)</td>
<td>49</td>
<td>2 (4.1)</td>
<td>1 (50.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 µM Cas9 protein and 40 ng/µl gRNA_B</td>
<td>42</td>
<td>38 (90.5)</td>
<td>38</td>
<td>4 (10.5)</td>
<td>4 (100)</td>
</tr>
</tbody>
</table>

*Results using the conventional method were from a previous report (Nakagawa et al., 2015).

**Design and validation of two loxP insertions**

We designed a novel strategy for the one-step generation of floxed mice by utilizing double-cut CRISPR-Cas9 and a single ssODN donor. To perform a proof-of-concept examination of this strategy, we constructed an all-in-one CRISPR-Cas9 vector expressing two gRNAs to target the both regions outside a 55-bp exon in osteopontin protein (Okazaki et al., 2010) (Fig. 4A). The ssODN was designed to carry the 55-bp exon flanked by the loxP sequences and 30-bp homology arms.

To confirm the cleavage activity of the all-in-one CRISPR-Cas9 vector and the integration rate of ssODN, the vector and ssODN were microinjected into the pronucleus of zygotes. The resultant 115 surviving oocytes were cultured for 3.5 days. We observed 33 blastocysts, and each blastocyst was used for single blastocyst assays. To roughly genotype the blastocysts, three kinds of PCR were performed (Fig. 4A): PCR1 to amplify the whole targeted locus, PCR2 to detect the integration of the left loxP, and PCR3 to detect the integration of the right loxP. Additionally, we performed RFLP analysis to confirm the integration of the left and right loxP sequences in the PCR1 products using PsiI and XmnI, respectively (Fig. 4A). The result of PCR1 suggested that one blastocyst (#19) had a floxed allele (Fig. S2A). Consistent with this result, blastocyst #19 was also positive in PCR2 and PsiI-RFLP (Figs S2B, S3A). In PCR3 and XmnI-RFLP, however, four (#11, #14, #25, and #27) and six (#1, #11, #14, #19, #21, and #27) blastocysts were estimated to be positive, respectively (Fig. S2C, Fig. S3B). These discordances might be caused by various patterns of partial integrations, such as the integration of only one loxP sequence and/or integration with indel mutations, as well as the difficulty of performing single blastocyst assays because of the small amount of template genome available. To ensure the exact genotype, the six PCR1 products from blastocysts #1, #11, #14, #19, #21, and #27, were analyzed by direct sequencing. Consequently, we confirmed the nearly correct integration of ssODN in one blastocyst (#19) (Fig. S4), correct integration of the right loxP in four blastocysts (#11, #14, #21, and #27) and nearly correct integration of the right loxP in one blastocyst (#1). No loxP insertion or indel mutation was observed at the left target site in the six blastocysts except for #19, suggesting that the left gRNA has a lower cleavage activity compared with the right gRNA.

**Generation of floxed mice**

Based on the results of Il11 and Spp1 targeting, we chose microinjection of Cas9 RNP instead of CRISPR-Cas9 vector to generate the founder mice. According to the validation using blastocysts, we increased the concentration of the left gRNA compared with that of the right gRNA to balance the different cleavage activity of these two gRNAs. We microinjected the left and right gRNAs together with the Cas9 protein and ssODN into zygotes generated via the ultra-superovulation method, under several concentration conditions. We then transferred the surviving zygotes to pseudopregnant mice. As summarized in Table 3, higher concentrations of ssODN resulted in a reduction of the birth rate. Tail lysates of all pups were analyzed using the PCRs and two RFLP analyses similar to the blastocyst assay. The results of these assays suggested as follows: five pups, including one pup injected with 2 µM, two with 1.5 µM, and two with 1 µM Cas9, had the left loxP but did not have the right loxP, because they were positive for both the PCR2 and PsiI-RFLP analyses, while they were negative for both the PCR3 and XmnI-RFLP analyses (four pups) or for the XmnI-RFLP analysis (one pup). Another five pups, including one injected with 2 µM, two with 1.5 µM, and two with 1 µM Cas9, had the left loxP but did not have the right loxP, because they were positive for both the PCR3 and XmnI-RFLP analyses.
analyses, while they were negative for both the PCR2 and PsiI-RFLP analyses (four pups) or for the PsiI-RFLP analysis (one pup). Among 37 pups analyzed, three pups, including two injected with 1.5 µM and one pup with 1 µM Cas9, were positive for all analyses, which were identified as potential floxed mice (Table 3). The PCR products from these pups were cloned and analyzed by DNA sequencing. Of these, one pup was identified with a precisely floxed allele, and the other two pups carried nearly correct integrations (Fig. 4B).

**DISCUSSION**

We have previously shown that reproductive engineering techniques were useful for creating knockout mice using a genome-editing technique (Nakagawa et al., 2014, 2015). In this study, we showed that approximately two and three times the number of fertilized oocytes could be generated from sexually matured and immature C57BL/6 female mice, respectively, by IVF using our ultrsuperoovulation method, consistent with a previous report (Takeo and Nakagata, 2015). To clarify its applicability in the generation of genome-edited mice, we demonstrated the successful production of gene-disrupted, single-amino-acid-substituted, and floxed mice by microinjecting the CRISPR-Cas9 plasmid vectors or RNPs with or without ssODN donors.

We initially aimed to generate various genome-edited mice by microinjecting of an all-in-one CRISPR-Cas9 plasmid vector in the C57BL/6 strain because of its simple handling. However, the birth rates of Cas9 nuclease plasmid-injected zygotes were low, similar to other reports using inbred strains (Li et al., 2013; Horii et al., 2014), when compared with B6D2F1 hybrid mice (Mashiko et al., 2013). To solve this problem, we demonstrated that the use of Cas9 protein and synthesized gRNA boosted the birth rate, leading to the successful generation of various genome-edited mice.

Recent achievements using a long double-stranded plasmid donor together with CRISPR-Cas9 enabled the one-step generation of floxed mice (Lee and Lloyd, 2014). In our study, a long ssODN was...
used instead of the plasmid donor to avoid the labor of donor construction and lower the risk of random integration, and the successful generation of the intended mouse was demonstrated. Unfortunately, three pups containing precise or slightly mutated floxed alleles died within about 2 weeks after birth. This might be explained because they were mosaic founders containing the conditional allele with some other alleles, such as alleles harboring short indel mutations and exon-skipped alleles, which might be

Table 2. Generation of Spp1-modified mice

<table>
<thead>
<tr>
<th>Age in weeks</th>
<th>CRISPR-Cas9</th>
<th>ssODN</th>
<th>Injected</th>
<th>Survived (%)</th>
<th>Transferred</th>
<th>Pups (%)</th>
<th>Precise knock-in</th>
<th>Knock-in with indel mutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>4–5</td>
<td>3 ng/µl plasmid</td>
<td>10 ng/µl</td>
<td>76</td>
<td>70 (92.1)</td>
<td>70</td>
<td>2 (2.9)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>3 ng/µl plasmid</td>
<td>10 ng/µl</td>
<td>117</td>
<td>112 (95.7)</td>
<td>112</td>
<td>0 (0)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>5</td>
<td>5 ng/µl plasmid</td>
<td>10 ng/µl</td>
<td>35</td>
<td>31 (88.6)</td>
<td>31</td>
<td>1 (3.2)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>0.5 µM Cas9 protein and 20 ng/µl gRNA</td>
<td>10 ng/µl</td>
<td>160</td>
<td>151 (94.4)</td>
<td>113</td>
<td>8 (7.1)</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>0.75 µM Cas9 protein and 30 ng/µl gRNA</td>
<td>10 ng/µl</td>
<td>78</td>
<td>69 (88.5)</td>
<td>69</td>
<td>8 (11.6)</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>11</td>
<td>0.5 µM Cas9 protein and 20 ng/µl gRNA</td>
<td>10 ng/µl</td>
<td>155</td>
<td>148 (95.5)</td>
<td>111</td>
<td>13 (11.7)</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>10</td>
<td>0.75 µM Cas9 protein and 30 ng/µl gRNA</td>
<td>10 ng/µl</td>
<td>41</td>
<td>39 (95.1)</td>
<td>39</td>
<td>5 (12.8)</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

Fertilized oocytes were created by IVF using the ultra-superovulation method (IASe-hCG).

Fig. 4. Generation of floxed mice at the Gpcpd1 locus. (A) Schematic illustration to generate a conditional allele at the Gpcpd1 locus. A 55-bp exon was targeted by two gRNA, which were designed to cut both ends of the exon. A single ssODN was designed to carry the two loxP sequences. Bold arrows indicate primers for PCR. Thin arrows indicate the recognition sites of restriction enzymes for the RFLP analysis. (B) Sequencing analysis of subcloned PCR1 products from three pups that were positive for all analyses. The intended floxed allele is shown at the top (Floxed) with loxP sequences (enclosed in black boxes). The wild-type allele is shown at the bottom. Dots indicate the same bases as the floxed sequence.
disruptive in Gpcpd1 functions. To overcome this, the knock-in efficiency of ssODN must be further increased by using chemically modified ssODN (Renaud et al., 2016) and/or HDR-enhancing molecules such as Scr7 (Maruyama et al., 2015; Chu et al., 2015) and RS-1 (Song et al., 2016), or the two-step integration of ssODNs should be performed. Another possible reason for the lethality is abnormal pre-mRNA splicing caused by the loxP insertions in the proximity of the target exon. This problem is difficult to solve using chemically synthesized ssODN as a donor, but several methods to synthesize longer single-stranded DNA (ssDNA) were reported recently (Miura et al., 2015; Yoshimi et al., 2016). By incorporating these methods into our strategy, the length limit of chemically synthesized ssODN should be resolved. In summary, our reproductive engineering techniques, especially the ultra-superoovulation method, and the improved methods of genome editing will help improve animal welfare and work efficiency when producing GM mice.

MATERIALS AND METHODS
Plasmid construction, gRNA synthesis, and preparation of Cas9 protein and ssODN
All-in-one CRISPR-Cas9 plasmids for the Il11 gene were constructed as previously described (Nakagawa et al., 2015). All-in-one CRISPR-Cas9 plasmids for Spp1 and Gpcpd1 genes were constructed using the Multiplex CRISPR/Cas9 Assembly System Kit (Addgene, Cambridge, MA, USA, Kit #1000000055) according to a previous report (Sakuma et al., 2014). Briefly, sense and antisense oligonucleotides for the gRNA templates were annealed and inserted into the pX330-based vectors contained in the kit. Subsequently, gRNA cassettes were tandemly assembled using the Golden Gate assembly method. In vitro transcribed gRNAs were prepared according to a previous report (Aida et al., 2015). Briefly, template DNA fragments were generated using PCR amplification from the CRISPR-Cas9 vectors described above using primers containing a T7 promoter sequence. Subsequently, the gRNAs were synthesized using a MEGASHortscript T7 Kit (Life Technologies, Carlsbad, CA, USA), and then purified with a MEGAClear Kit (Life Technologies). The ssODNs were synthesized by IDT (Corvalle, IA, USA). The sequences of oligonucleotides for gRNA templates, primers, and ssODNs are listed in Table S3. The recombinant Cas9 protein was obtained from New England Biolabs Japan (Cas9 Nuclease NLS, Streptococcus pyogenes; Tokyo, Japan).

Animals
C57BL/6Jcrl strain was purchased from CLEA Japan (Tokyo, Japan). After breeding, C57BL/6J female mice were used as oocyte donors at 4–5 or 10–13 weeks of age. C57BL/6J male mice were used as sperm donors over 12 weeks of age for IVF. C57BL/6 female mice were intraperitoneally administered with L: 20 ng/µl and R: 20 ng/µl of 2 µM L: 50 ng/µl and R: 30 ng/µl of 1.5 µM L: 20 ng/µl and R: 20 ng/µl of 1 µM L: 25 ng/µl and R: 15 ng/µl of ssODN. To detect the targeted gRNA, PCR were carried out using KOD FX (Toyobo, Osaka, Japan) with the Spp1 F and R primers listed in Table S3 under the following conditions: 95°C for 1 min, followed by 39 cycles of 98°C for 10 s, 61°C for 30 s, and 68°C for 30 s. Each PCR product was analyzed by direct sequencing using an ABI 3130 Genetic Analyzer (Life Technologies). To detect the targeted Gpcpd1 gene, PCR1 was performed with the F and R primers listed in Table S3 under the following conditions: 95°C for 1 min, followed by 39 cycles of 98°C for 10 s, 61°C for 30 s, and 68°C for 45 s. The PCR1 products were subjected to automatic electrophoresis using MultiNA (Shimadzu Corporation, Kyoto, Japan). The PCR products were purified and analyzed by RFLP analyses using BsmFI (New England Biolabs Japan), then the PCR products identified as positive were analyzed by direct sequencing using an ABI 3130 Genetic Analyzer (Life Technologies) with a BigDye Terminator v1.1 Cycle Sequencing Kit (Life Technologies). To detect the targeted Il11 gene, PCR1 was performed with the F and R primers listed in Table S3 under the following conditions: 95°C for 1 min, followed by 39 cycles of 98°C for 10 s, 61°C for 30 s, and 68°C for 45 s. PCR2 or PCR3 were performed with the F and L-lox primers or R-lox and R primers listed in Table S3 under the following conditions: 95°C for 1 min, followed by 39 cycles of 98°C for 10 s, 57°C for 30 s, and 68°C for 25 s. Each PCR product was subjected to automatic electrophoresis using MultiNA. The PCR1 products were purified and analyzed by RFLP analyses using PsaI (New England Biolabs Japan) or XmnI (New England Biolabs Japan), then the PCR1 products identified as positive in all analyses were analyzed by direct sequencing.

Analysis of pups
Tail lysates of pups were prepared by an alkaline lysis method and PCR was performed using KOD FX with each primer set. For the analysis of Il11 mutants, the IL11 F and R primers listed in Table S3 were used according to a previous report (Nakagawa et al., 2015). Each PCR product was analyzed by direct sequencing. For the analysis of Spp1 gene, PCR1 were carried out according to the previous report and their names were added to the report.
with 37 cycles, and then the products were analyzed as described in the same way as the single blastocyst assay. For the analysis of Gpcpa1 gene, three kinds of PCR were carried out with 37 cycles, and then the products were analyzed as described in the same way as the single blastocyst assay. The PCR1 products harboring floxed mutations were subcloned into a pTA2 vector using Target Clone -Plus- (Toyobo). Each subcloned vector was PCR1 products harboring floxed mutations were subcloned into a pTA2 vector and analyzed as described in the same way as the single blastocyst assay. The kinds of PCR were carried out with 37 cycles, and then the products were analyzed as described in the same way as the single blastocyst assay.

Acknowledgements

We thank Ms Kazuko Wamakatsu and the other lab members for their technical assistance.

Competing interests

The authors declare no competing or financial interests.

Author contributions

Y.N designed the study, carried out the mouse experiments and analyses, and wrote the manuscript. T.S. designed the study, constructed CRISPR-Cas9 vectors, synthesized gRNAs, and wrote the manuscript. N.N., Y.Y., N.Y., and T.T. provided instructions. N.N and T.Y supervised the work. All authors read and approved the manuscript.

Funding

This work was supported by a Japan Society for the Promotion of Science (JSPS) Grants-in-Aid for Scientific Research B [grant number 15H04606 to N.N.] and Grants-in-Aid for Young Scientists B [grant number 16K18478 to T.S.].

Supplementary information

Supplementary information available online at http://bio.biologists.org/lookup/doi/10.1242/bio.019349.supplemental

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