Re-Evaluating One-step Generation of Mice Carrying Conditional Alleles by CRISPR-Cas9-Mediated Genome Editing Technology

Authors:

Channabasavaiah Gurumurthy^{1,2}*, Rolen Quadros¹, John Adams Jr³, Pilar Alcaide⁴, Shinya Ayabe⁵, Johnathan Ballard³, Surinder K. Batra⁶, Marie-Claude Beauchamp⁷, Kathleen A Becker⁸, Guillaume Bernas⁹, David Brough¹⁰, Francisco Carrillo-Salinas⁴, Ruby Dawson¹¹, Victoria DeMambro⁸, Jinke D'Hont^{12,13}, Katharine Dibb¹⁴, James D. Eudy¹⁵, Lin Gan¹⁶, Jing Gao⁴⁴, Amy Gonzales³, Anyonya Guntur⁸, Huiping Guo³, Donald W. Harms¹, Anne Harrington⁸, Kathryn E. Hentges¹⁷, Neil Humphreys¹⁸, Shiho Imai¹⁹, Hideshi Ishii²⁰, Mizuho Iwama⁵, Eric Jonasch²¹, Michelle Karolak⁹, Bernard Keavney²², Nay-Chi Khin⁴⁴, Masamitsu Konno²³, Yuko Kotani²⁴, Yayoi Kunihiro²⁴, Imayavaramban Lakshmanan⁶, Catherine Larochelle²⁵, Catherine B. Lawrence¹⁰, Lin Li²⁶, Volkhard Lindner⁸, Xian-De Liu²¹, Gloria Lopez-Castejon²⁷, Andrew Loudon²⁸, Jenna Lowe⁴⁴, Loydie Jerome-Majeweska⁷, Taiji Matsusaka¹⁹, Hiromi Miura^{29,30}, Yoshiki Miyasaka²⁴, Benjamin Morpurgo³, Katherine Motyl⁸, Yo-ichi Nabeshima³¹, Koji Nakade⁵, Toshiaki Nakashiba⁵, Kenichi Nakashima⁵, Yuichi Obata⁵, Sanae Ogiwara³², Mariette Ouellet⁹, Leif Oxburgh⁸, Sandra Piltz¹¹, Ilka Pinz⁸, Moorthy P. Ponnusamy⁶, David Ray³³, Ronald J. Redder¹⁵, Clifford J Rosen⁸, Nikki Ross⁴⁴, Mark T. Ruhe³⁴, Larisa Ryzhova⁸, Ane M. Salvador⁴, Radislav Sedlacek³⁵, Karan Sharma³⁶, Chad Smith²¹, Katrien Staes^{12,13}, Lora Starrs⁴⁴, Fumihiro Sugiyama³⁷, Satoru Takahashi³⁷, Tomohiro Tanaka³⁸, Andrew Trafford¹⁴, Yoshihiro Uno²⁴, Leen Vanhoutte^{12,13}, Frederique Vanrockeghem^{12,13}, Brandon J. Willis³⁴, Christian S. Wright³⁹, Yuko Yamauchi²⁴, Xin Yi³⁹, Kazuto Yoshimi²⁴, Xuesong Zhang²¹, Yu Zhang²⁶, Masato Ohtsuka^{29,30}, Satyabrata Das⁴⁰, Daniel J. Garry^{40,41}, Tino Hochepied^{12,13}, Paul Thomas¹¹, Jan Parker-Thornburg²¹, Antony D Adamson¹⁸, Atsushi Yoshiki⁵, Jean-Francois Schmouth⁹, Andrei Golovko³, William R. Thompson³⁹, KC. Kent Lloyd^{34,42}, Joshua A. Wood³⁴, Mitra Cowan⁴³, Tomoji Mashimo²⁴, Seiya Mizuno³⁷, Hao Zhu²⁶, Petr Kasparek³⁵, Lucy Liaw⁸, Joseph M. Miano¹⁶, and Gaetan Burgio⁴⁴*.

- 1. Mouse Genome Engineering Core Facility, Vice Chancellor for Research Office, University of Nebraska Medical Center, Omaha, NE, USA
- 2. Developmental Neuroscience, Munroe Meyer Institute for Genetics and Rehabilitation, University of Nebraska Medical Center, Omaha, NE, USA
- 3. Texas A&M Institute for Genomic Medicine (TIGM), Texas A&M University, College Station, TX 77843, USA
- 4. Department of Immunology, Tufts University School of Medicine, Boston, USA
- 5. RIKEN BioResource Research Center, Tsukuba, Ibaraki 305-0074, Japan

- 6. Department of Biochemistry and Molecular Biology, University of Nebraska Medical Center, Omaha, NE, USA
- 7. Departments of Anatomy and Cell Biology, Human Genetics and Pediatrics, Research Institute McGill University Health Center (RI-MUHC), Montreal, Canada
- 8. Maine Medical Center Research Institute (MMCRI), Scarborough, ME, USA.
- 9. Transgenesis and Animal Modeling Core Facility, Centre de Recherche du Centre Hospitalier Universitaire de Montreal (CRCHUM), Montreal, Canada
- 10. Division of Neuroscience and Experimental Psychology, School of Biological Sciences, Faculty of Biology, Medicine and Health, Manchester Academic Health Science Centre, University of Manchester, AV Hill Building, Oxford Road, Manchester, M13 9PT, U.K
- 11. South Australian Health & Medical Research Institute and Department of Medicine, University of Adelaide, Australia
- 12. Transgenic mouse core facility, VIB Center for Inflammation Research, Ghent, Belgium
- 13. Department of Biomedical Molecular Biology, Ghent University, Ghent, Belgium
- 14. Unit of Cardiac Physiology, School of Medical Sciences, Manchester Academic Health Science Center, University of Manchester, UK
- 15. High-Throughput DNA Sequencing and Genotyping Core Facility, Vice Chancellor for Research Office, University of Nebraska Medical Center, Omaha, USA
- 16. University of Rochester Medical Center, Rochester, NY 14642, USA
- 17. Division of Evolution and Genomic Sciences, School of Biological Sciences, Faculty of Biology, Medicine and Health, Manchester Academic Health Science Centre, University of Manchester, UK
- 18. Transgenic Unit core facility, Faculty of Biology, Medicine and Health, University of Manchester, Manchester, UK
- 19. Department of Basic Medicine, Division of Basic Medical Science and Molecular Medicine, School of Medicine, Tokai University, 143, Shimokasuya, Isehara, Kanagawa 259-1193, Japan
- 20. Department of Medical Data Science, Osaka University Graduate School of Medicine, Japan
- 21. The University of Texas, MD Anderson Cancer Center, Houston, TX, USA.
- 22. Division of Cardiovascular Sciences, School of Medical Sciences, Faculty of Biology, Medicine and Health, The University of Manchester AND Manchester Heart Centre, Manchester University NHS Foundation Trust, Manchester Academic Health Science Centre, Manchester, UK
- 23. Department of Frontier Science for Cancer and Chemotherapy, Osaka University Graduate School of Medicine, Osaka, Japan

- 24. The Institute of Experimental Animal Sciences, Osaka University Graduate School of Medicine, Osaka, Japan
- 25. Centre de Recherche du Centre Hospitalier Universitaire de Montreal (CRCHUM), Montreal, Canada
- 26. Children's Research Institute Mouse Genome Engineering Core, University of Texas Southwestern Medical Center, Dallas, TX 75390, USA.
- 27. Manchester Collaborative Centre for Inflammation Research (MCCIR), School of Biological Sciences, Faculty of Biology, Medicine and Health, The University of Manchester, Manchester, UK.
- 28. Centre for Biological Timing, School of Medical Sciences, Faculty of Biology, Medicine and Health, University of Manchester, Manchester, UK
- 29. Center for Matrix Biology and Medicine, Graduate School of Medicine, Tokai University, Isehara, Kanagawa, 259-1193, Japan
- 30. Department of Molecular Life Science, Division of Basic Medical Science and Molecular Medicine, School of Medicine, Tokai University, 143, Shimokasuya, Isehara, Kanagawa 259-1193, Japan
- 31. Laboratory of Molecular Life Science, Foundation for Biomedical Research and Innovation, at Kobe, Japan
- 32. Department of Laboratory Animal Science, Support Center for Medical Research and Education, Tokai University, 143, Shimokasuya, Isehara, Kanagawa 259-1193, Japan
- 33. Oxford Centre for Diabetes, Endocrinology and Metabolism, University of Oxford, OX37LE, UK
- 34. Mouse Biology Program, University of California, Davis, USA
- 35. Laboratory of Transgenic Models of Diseases and Czech Centre for Phenogenomics, Institute of Molecular Genetics of the Czech Academy of Sciences, Czech Republic
- 36. College of Osteopathic Medicine, Marian University, Indianapolis, IN 46222, USA
- 37. Laboratory Animal Resource Center, University of Tsukuba, Japan
- 38. Department of Gastroenterology and Metabolism, Nagoya City University Graduate School of Medical Sciences, Nagoya, Japan.
- 39. School of Health and Human Sciences, Department of Physical Therapy, Indiana University, Indianapolis, IN 46202, USA
- 40. Lillehei Heart Institute Regenerative Medicine and Sciences Program, University of Minnesota, Minneapolis, MN, USA.

- 41. Paul and Sheila Wellstone Muscular Dystrophy Center, University of Minnesota, Minneapolis, MN, USA
- 42. Dept. of Surgery, School of Medicine, University of California, Davis, Davis, USA
- 43. McGill Integrated Core for Animal Modeling (MICAM), Montreal, Canada
- 44. Department of Immunology and Infectious Disease, the John Curtin School of Medical Research, the Australian National University, Canberra, Australia.

^{*} Corresponding authors: cgurumurthy@unmc.edu and gaetan.burgio@anu.edu.au

Abstract:

CRISPR-Cas9 gene editing technology has considerably facilitated the generation of mouse knockout alleles, relieving many of the cumbersome and time-consuming steps of traditional mouse embryonic stem cell technology. However, the generation of conditional knockout alleles remains an important challenge. An earlier study reported up to 16% efficiency in generating conditional knockout alleles in mice using 2 single guide RNAs (sgRNA) and 2 single-stranded oligonucleotides (ssODN) (2sgRNA-2ssODN). We re-evaluated this method from a large data set generated from a consortium consisting of 17 transgenic core facilities or laboratories or programs across the world. The dataset constituted 17,887 microinjected or electroporated zygotes and 1,718 live born mice, of which only 15 (0.87%) mice harbored 2 correct LoxP insertions in cis configuration indicating a very low efficiency of the method. To determine the factors required to successfully generate conditional alleles using the 2sgRNA-2ssODN approach, we performed a generalized linear regression model. We show that factors such as the concentration of the sgRNA, Cas9 protein or the distance between the placement of LoxP insertions were not predictive for the success of this technique. The major predictor affecting the method's success was the probability of simultaneously inserting intact proximal and distal LoxP sequences, without the loss of the DNA segment between the two sgRNA cleavage sites. Our analysis of a large data set indicates that the 2sgRNA-2ssODN method generates a large number of undesired alleles (>99%), and a very small number of desired alleles (<1%) requiring, on average 1,192 zygotes.

Introduction:

Many inherited diseases are caused by defective genes. A better understanding of the mechanisms of these defects is critical to obtaining precise diagnoses and finding new therapeutics. Gene inactivation through knockout alleles in model organisms such as flies, worms, zebrafish and mice provides invaluable insights into mechanisms of gene function and disease [1]. However important challenges remain to successfully analyze the phenotypic impact of knockout genes in adult model organisms as over 30% of the genes in mice are essential for development and cause embryo lethality or neonatal subviability when deleted [2]. To overcome lethal phenotypes in gene-knockout models, conditional knockout (cKO) strategies have emerged [3]. cKO models usually involve insertion of LoxP sites in introns flanking critical exon/s or (less commonly) in intergenic regions or flanking regulatory regions such as promoters and enhancers. When crossed with a *Cre* recombinase expressing driver mouse, the *Cre* enzyme recognizes LoxP sequences and removes the intervening sequence. This leads to functional inactivation of the targeted gene in only the cells where the Cre is expressed and capable of targeting the DNA [3]. Generating a cKO mouse previously required the use of embryonic stem (ES) cell-based homologous recombination in combination with embryo manipulation, microinjection (MI), and assisted reproduction technologies (ART) [4]. These techniques were established in the 1980s and are still being used as gold-standard methods. Based on this technology, large-scale efforts such as the KnockOut Mouse Project (KOMP) [5] and the European Conditional Mouse Mutagenesis (EUCOMM) Program [6] have designed thousands of gene targeted constructs in ES cells for over 90% of coding genes. Using the ES cell clones, about 25% of mouse genes have been converted into cKO mice, all readily available and accessible in public repositories [7].

The recent emergence of genome editing technologies such as ZFN, TALENS and CRISPR-

Cas9 enables an improvement in efficiency of gene targeting and has considerably facilitated the

generation of genetically-engineered animal models based on homology directed repair of donor

constructs in mouse zygotes [8]. Endonucleases, particularly Class 2 CRISPR systems, generate

a precise double strand break (DSB) in the DNA under a chimeric single guide RNA (sgRNA)

[9]. The DSB leads to error-prone, non-homologous end joining (NHEJ) repair or the precise

homology-directed repair (HDR) under the guidance of a repair template [8]. In an earlier study,

a high success rate (16%) of targeting LoxP sites in cis was reported by using 2 sgRNAs and 2

single-stranded oligonucleotides (ssODN) containing LoxP sites (2sgRNA-2ssODN) flanking a

targeted critical exon (Figure 1) [10].

We sought to investigate the efficiency of the 2sgRNA-2ssODN method for the generation of

cKO alleles. We describe here for the first time a global community effort from a consortium of

over a dozen laboratories, transgenic core facilities and programs across the world to evaluate the

efficiency of generating cKO alleles using the 2sgRNA-2ssODN approach. We surveyed over 50

loci and over 17,000 microinjected or electroporated zygotes using this method, which enabled

robust statistical power to evaluate the efficiency of the technique. In contrast to the earlier report

[10], we find this method does not efficiently produce cKO alleles. Rather, it generally results in

a series of undesired editing events at the cleavage sites which occur nearly 100 fold higher rate

7

than the precise insertion, in *cis*, of the two *LoxP* sites.

Material and methods:

Ethical statement:

All experiments were approved from the respective Institutional Animal Care and Use Committees in the USA and Ethics Committees in Australia, Belgium, Czech Republic, Japan, Spain and UK according to guidelines or code of practice from the National Institute of Health in the USA, the National Health and Medical Research Council (NHMRC) in Australia, Animals (Scientific Procedures) Act 1986 in UK or MEXT (Ministry of Education, culture, sports, Science and Technology), MHLW (The Ministry of Health, Labor and Welfare) in Japan, the central commission for Animal Welfare (CCAW) in Czech Republic, the Canadian Council on Animal Care (CCAC) in Canada, the National Ethics Code from the Royal Belgian (Flemish) Academy of Medicine in Belgium, and the European code of Conduct for Research Integrity from All European Academies.

Mecp2 gene targeting using CRISPR-Cas9:

Mecp2 left single chimeric guide RNAs (sgRNA) 5'-CCCAAGGATACAGTATCCTA-3' and *Mecp2* right sgRNA 5'-AGGAGTGAGGTCTAGTACTT-3' target sites were designed as described in Yang et al [10]. Ultramer Oligonucleotides (Integrated DNA Technologies, Coralville, IA) were designed with sequences to T7 promoter for *in vitro* transcription, DNA target region, and chimeric RNA sequence. Complimentary oligos for each target sequence were annealed at 95°C for 5 mins and the temperature was reduced 0.20°C/second to 16°C using a PCR machine (BioRad T100) before use as template for sgRNA synthesis. sgRNAs were synthesized with the HiScribeTM T7 Quick High Yield RNA Synthesis Kit (New England Biolabs). Cas9 mRNA was obtained from Life Technologies or in-vitro transcribed from a Chimeric pX330-U6-Chimmeric-BB-CBh-hSpCas9 expression plasmid obtained from Addgene repository (Plasmid 42230; donation from Zhang laboratory).

SgRNA design:

SgRNAs were designed using available online tools such as CRISPOR, Chop-Chop or CCTop [11, 12]. SgRNAs were cloned into pX330 and *in vitro* transcribed[13-15], or synthesized and annealed [16]. Cas9 mRNA or protein was purchased, *in vitro* transcribed or purified in house. Cas9 protein was complexed with thesgRNA or crRNA and the tracrRNA [17] and then mixed with the ssODN prior to microinjection. Concentrations and site of injection for Cas9 protein or mRNA, sgRNA, and template repairs for each locus are indicated in Supplementary Table 1.

Mouse husbandry and zygote microinjection and electroporation:

Mice were purchased from various sources and maintained under specific pathogen-free conditions. Mice were maintained under 12/12 hr light cycle and food and water were provided *ad libitum*. Three to five week-old females were superovulated by intraperitoneal injection of Pregnant Mare Serum Gonadotropin (5IU) followed by intraperitoneal injection of Human Chorionic Gonadotropin hormone (5IU) 48 hours later. Superovulated females were mated with 8 to 20 week-old stud males. The mated females were euthanized the following day and the zygotes were collected from their oviducts. Cytoplasmic or pronuclear injections were performed under an inverted microscope, associated micromanipulators, and a microinjection apparatus. Electroporation of the embryos were performed with an electroporation device using a cuvette or 1mm plate electrodes with the following parameters: 30 V square wave pulses with 100 ms interval using a BioRad electroporator device or 4 poring pulses (40 V, 3.5 ms, interval 50 ms, 10% voltage decay + polarity) followed by 5 transfer pulses (5 V, 50 ms, interval 50 ms, 40% voltage decay, alternating + and – polarity) using a NEPA21 electroporator device. Microinjected or electroporated zygotes were either surgically transferred into the ampulla of pseudo-pregnant

females or cultured overnight at 37°C and then surgically transferred at the 2-cell stage of development.

Genotyping:

DNA extraction was performed on ear punch or tail tip from mouse pups over 15 days using a DNA extraction kit according to the manufacturer instructions. Primers were designed to amplify the regions encompassing the integrated *LoxP* sequence. PCR was performed using Taq polymerase under standard PCR conditions. The PCR products were then purified with ExoSAP-IT1 or a PCR Clean-Up System kit according to the manufacturer's instructions. Sanger sequencing was performed in core facilities. To identify *LoxP* insertions, as a general practice at all centers, the two target sites were amplified individually to look for increase in the amplicon size, which occurs if *LoxP* sites are inserted successfully. If the *LoxP* insertion was not observed in this first set of PCR analyses, the samples were declared negative, and in many such cases the samples were not analyzed further (as the end goal of the project, ie., generation of floxed allele was not met). In some cases, such samples were also sequenced to assess *indels* to understand if the guides were successful in cleaving the target site. In some cases, the entire regions encompassing both the guide cleavage sites were amplified to assess for deletions between the cleavage sites.

Statistics:

To determine the statistical differences between proportions or means, we performed a Fisher Exact test or a Kruskal Wallis sum rank test. A Generalized linear model calculation was performed with success of the 2sgRNA-2ssODN method as a response. Predictive variables were: efficiency of the sgRNA, probability of *LoxP* insertions in 5' and 3 (5' *LoxP* and

3'_LoxP), simultaneous insertion of the 2 LoxP sites (interaction between 5'_Loxp and 3'_LoxP) Cas9 mRNA, protein, plasmid and ssODN concentrations and distance between distal and proximal target sites. Variance for each predictor was determined from the diagonal of the variance-covariance matrix. Effect sizes and type II error were determined using Cohen effect size d statistics and power calculation. All statistical analyses were performed using Rstudio v1.1.423. Results were considered statistically significant at p<0.05.

Results:

Mecp2 gene targeting in blastocysts:

To assess the efficiency of the technique and compare to previously published results [10], we reproduced an experiment on *Mecp2* gene, essential for DNA methylation during development using the same sgRNAs and ssODNs as previously described in the original report [10]. Three independent centers at the Australian National University in Australia (ANU), University of Nebraska Medical Center in the USA (UNMC) and the Czech Centre for Phenogenomics in Czech Republic (IMG) performed these experiments on C57BL/6N inbred strain of mice. We evaluated the success rate of the 2sgRNA-2ssODN method in blastocysts for *Mecp2* (Table 1). Using a concentration mix of 20 ng/µl of Cas9 mRNA, 20 ng/µl of in-vitro transcribed sgRNA, and 10 ng/µl of ssODN, we observed no successful targeting (i.e., correct insertion of 2 *LoxP* sites in *cis* configuration) even though both sgRNAs cleaved target DNA as indicated by the presence of *indels* or integration of a *LoxP* site at the desired location, which varied from 13% to 34% (Table 1).

	Zygotes injected	Blastocysts genotyped	Correctly targeted	Incorrectly targeted at the 5' site (%)	Incorrectly targeted at the 3' site (%)
Australian National University (ANU) Australia	106	51	0	11 <i>indels</i> and 6 <i>LoxP</i> correctly inserted (33%)	6 <i>indels</i> and 1 <i>LoxP</i> correctly inserted (13%)
University of Nebraska Medical Center (UNMC) USA	80	70	0	14 <i>indels</i> and 1 <i>LoxP</i> correctly inserted (34%)	21 indels (30%)
Czech Centre for Phenogenomics, Czech republic (BIOCEV/IMG)	40	28	0	8 <i>indels</i> and 1 <i>LoxP</i> correctly inserted (32%)	5 indels (18%)

Table 1: Summary of the edited blastocysts for *Mecp2* gene from three different centers.

Interestingly we noted the occasional presence of mutations within *LoxP* sites indicating illegitimate repair events at the target site. The frequency of successful targeting of two *LoxP* sites *in cis* was previously reported to be 16% [10], which we failed to achieve. One possible explanation is the mouse genetic background influences the likelihood of ssODN integration. This variance could also be explained by an inherently low probability to successfully replace 2 genomic loci in *cis*, the lack of efficiency of the sgRNA, or the relatively low sample size.

A global survey of the generation of conditional alleles using 2sgRNA-2ssODN method.

To better understand how to successfully generate conditional alleles using the 2sgRNA–2ssODN approach and to assess its efficiency, we evaluated this method on 56 additional genes and two intergenic regions of the mouse genome from a consortium of 17 institutions across Australia, Belgium, Japan, USA, UK, Czech Republic and Canada. A majority of attempts were

performed on a C57BL/6J background (39) whereas 18 projects used C57BL/6N background and 3 additional ones used a hybrid mouse background (B6C3HF1, B6SJLF1, FVBCD1F1). We assessed whether the mouse background strain would have an impact over the success of the method using Fisher Exact test statistics. We failed to find such evidence in our data (Fisher exact test, p = 0.74). Out of the 56 targeted loci (49 microinjected and 7 electroporated), 21 were ranked as essential genes based on early embryonic or postnatal lethality of the homozygous knockout mice according to mouse genome database http://www.informatics.jax.org [18]. Different knockout mice from 18 out of 56 targeted loci were described viable to adulthood as homozygous mice and 17 loci were unknown. Together this indicates the repartition between putative essential and non-essential targeted gene was in equal frequency (Fisher exact test, p = 0.76). The distance between sgRNA varied from 250 bp to 1.1 Mb with a median of 2 Kb. Single exons to entire genes or regulatory genomic regions (Supplementary Table 1) were floxed. We investigated whether the distance between sgRNA is critical for the likelihood of success of the 2sgRNA-2ssODN method. We failed to find such evidence in our data set (Kruskal Wallis rank sum test, chi-squared = 32, p=0.42), although the sample size was too low to form a conclusion (Cohen's effect size d = 0.40 with power 1-beta = 0.27). Among the microinjected zygotes in 53 Loci, significantly higher number of zygotes were microinjected in the pronucleus alone (26/53) than the cytoplasm alone (10/53) or pronucleus and cytoplasm (17/53) (Fischer exact test p = 0.004), which is consistent with the current practice in most mouse transgenic core facilities (Figure 2A). Various forms of CRISPR reagents (sgRNA, Cas9 and ssODN), were microinjected or electroporated to generate the models (Supplementary Table 1). Consistent with the general practice in mouse transgenic facilities from 2013 to 2016 using CRISPR-Cas9 gene editing technology, the majority of the reagents were delivered in 59 Loci (49 unique loci

microinjected, 3 different designs for one loci and 7 electroporated Loci) in the form of in-vitro transcribed mRNA (35/59) at various concentrations varying from 10 ng/µl to 100 ng/µl of Cas9 mRNA (Figure 2B) and from 10 ng/µl to 50 ng/µl sgRNA. ssODN were delivered at a concentration varying from 10 ng/µl to 200 ng/µl. In 18 instances, Cas9 was delivered as protein with a concentration varying from 10 ng/µl to 75 ng/µl. Interestingly for 6 loci, Cas9 and sgRNAs were delivered in the form of a chimeric sgRNA-SpCas9 plasmid (pX330) at a concentration of 5 ng/µl. We sought to determine whether the forms of reagent delivery such as plasmid, ribonucleoprotein (RNP) or mRNA would have an effect on the overall efficiency in targeting using the 2sgRNA-2ssODN method. We failed to find such evidence (Fisher exact test p = 1). We therefore hypothesized that the success in generating floxed alleles using the 2sgRNA-2ssODN approach may depend on factors such as: (i) sgRNA efficiency, (ii) simultaneity in LoxP insertion, or; (iii) the concentration of the Cas9, sgRNA and ssODN reagents. To get insight on these possibilities, we further analyzed data from the 56 loci (Supplementary Table 2, 4 and 5). Note that the offspring for 54 loci were analyzed post-natal stage (Supplementary Tables 2 and 4) whereas 2 loci were analyzed at the blastocyst stage (Supplementary Table 4). Out of 17,887 (17,557 microinjected and 330 electroporated; see details below) zygotes, 12,764 (71.4%) zygotes were surgically transferred into recipient females. The recipient females gave birth to 1.718 pups (9.6% the microinjected/electroporated zygotes). As a general practice, at all centers, the mice were first analyzed by PCR to observe the putative insertion of the LoxP sites at both the sites; the animals were declared negative if genotyping did not reveal the presence of the desired allele. In some cases the loci were further analyzed to assess guide cleaving activity. Of the 1,684 founder mice, 659 (39%) showed some type of editing (indels and/or substitutions), 235 (14%) and 144 (9%)

mice harbored a single LoxP insertion or deletions between the two cleavage sites, respectively (Figure 2C). The mice for 25 (of the 56) loci were further assessed for additional events including large deletions (Figure 2C). Of the 487 founder mice analyzed (from those 25 loci), 41%, 11% and 2.7% samples contained *indels*, single *LoxP* insertions or large deletions respectively (Figure 2D). From the 1,684 animals analyzed, only 15 mice (0.87%) were correctly targeted with intact LoxP sites in the cis configuration (Supplementary Table 5). Out of 56 loci only 11 loci were successfully targeted (19.6%). The average number of zygotes needed to generate 1 correctly targeted animal was 1,192. The essentiality of the genes had no impact on the likelihood of success of the 2sgRNA-2ssODN technique (4/23 success in targeting for embryonic or postnatal lethality versus 5/18 for viable homozygous mice and 2/15 for unknown embryonic or postnatal lethality, Fisher exact test p = 0.27). We also noted from our data, among the 56 loci analyzed 14% loci showed deletions between two target sites for Cas9 cleavage. We also noted a relatively high occurrence of single LoxP insertions for > 20% of the mice genotyped (from all the loci) and few instances of trans LoxP insertions (on different alleles, reducing the probability for correct insertion of the LoxP sites) (Figure 3). We therefore hypothesized that success of this approach depends on the combined efficiency of the sgRNA and the likelihood of LoxP insertion on both sites to enable two in cis HDR events to occur simultaneously. To assess this postulate, we performed a generalized linear regression analysis to model the relationship between Cas9, sgRNA concentration, sgRNA cleavage efficiency, distance between LoxP insertions, occurrences of LoxP insertions, and success of the 2sgRNA-2ssODN method. The analyses are summarized in Supplementary Table 3. The efficiency of LoxP insertions at both 5' and 3' sites appears to be the best predictor for the likelihood of success of the 2sgRNA-2ssODN method accounting for over 60% of the total variance.

However, this predictor was not significant in our linear regression model. Additional predictors such as sgRNA efficiency or efficiency in 5'- or 3'- insertion of LoxP explained approximately 35% of the total variance but none of these predictors were significant in our model. The concentration of Cas9 mRNA accounted less than 0.1% of the total variance but was statistically significant (p <0.01) in the generalized linear regression model as a predictor for the success of the 2sgRNA-2ssODN approach. However, success of the 2sgRNA-2ssODN approach was not significantly correlated with an increase of Cas9 mRNA concentration (r^2 Pearson = 0.27, p=0.08). From our analysis, the sample size of the successful LoxP insertions in cis was too small to definitively rule out any others predictors (Cohen's effect size d = 0.4, power 1-beta = 0.41). Together, these results suggest that the presence of two simultaneous HDR events is the best predictor to generating two floxed alleles in cis.

Recently, electroporation of zygotes has been developed as an efficient method for generating knockout, point mutations, tagged, or conditional alleles [19-25]. From our consortium, 3 laboratories and programs surveyed the likelihood of success of the method. For 7 loci surveyed, we noted success in inserting a single *LoxP* allele (Supplementary Table 4) from analysis of blastocysts or live mice for 2 out of the 7 loci. In contrast we noted a relatively high frequency of large deletions and *indels* (up to 39% of large deletions) indicating successful editing. However, none of the loci showed two *LoxP* sites inserted in *cis* in the offspring, suggesting that the delivery of CRISPR reagents by electroporation does not make a statistical difference in obtaining a desired outcome from the 2sgRNA-2ssODN floxing approach, although the large numbers of embryos that can be manipulated allows for the recovery of the very small number of those that are correctly targeted.

Discussion

CRISPR-Cas9 technology has greatly facilitated the generation of mouse lines containing knockout or knockin alleles. However, the generation of conditional alleles remains a challenge using traditional ES cells and CRISPR-Cas9 gene editing technologies. An earlier paper demonstrated 16% efficiency with 2 chimeric sgRNAs and 2 single-stranded oligonucleotides to produce conditional alleles in mice [10].

To evaluate the efficiency of this 2sgRNA-2ssODN method, three laboratories replicated the experiments described in the initial report on *Mecp2* (10) using the same methods to generate the sgRNA and Cas9 and microinjected the mouse zygotes at similar reagent concentrations. Although we observed single *LoxP* site insertions and *indels* at the cleavage sites, the method was unsuccessful in generating two *LoxP* sites in *cis*. A previous report attempting to replicate the findings of Yang et al [10], found an efficiency of floxing *Mecp2* varying from 2% to 8% with the 2sgRNA-2ssODN approach [26]. We surmise the lack of efficiency in targeting *Mecp2* here was due to a low concentration of reagents delivered by microinjection, a notion corroborated by previous work from Horii and colleagues [26]. Of note, it was reported that up to 6% targeting efficiency was achieved using 25 ng/μl of Cas9, 6 ng/μl of sgRNA and 100 ng/μl of ssODN but toxic to embryonic development; these concentrations are 2-fold higher than those described in Yang *et al*, 2013 [10].

What determines the success of the 2sgRNA-2ssODN method?

To better understand the critical factors predicting the likelihood of success with the 2sgRNA-2ssODN approach, we surveyed 56 unique loci in mice zygote. We noted that the efficiency of simultaneous insertion of the two *LoxP* sites simultaneously was the best predictor of success

using this approach. We also noted a low success rate in generating a conditional allele across all centers (< 1%), varying from 0 to 50% (median = 0%) for individual loci. These results are comparable with previous reports demonstrating an important disparity in success rate varying from 0% to 7% of mice harboring two LoxP sites insertions in cis whether delivered by microinjection [26-29] or by electroporation [26]. We and others also have noted the large number of deletions at the target sites following DNA cleavage [28]. Our results on a larger number of loci suggest the efficiency in generating a successful cKO with the 2sgRNA-2ssODN method is lower than previously described [10]. One hypothesis for this discrepancy in success rate might relate to strain-specific differences. We analyzed this variable and did not find any significant differences among strains, whether the donor strain was a F1 cross, inbred, or outbred mouse line as a donor strain. Another possibility to improve the efficiency of the method is to avoid recombination between the target sites by placing the LoxP sites hundreds of kb apart. This was reported previously for a success rate varying from 0% to 18% for 6 loci [30]. We did not find such evidence in our data, although our sample size is too small to formally rule out this hypothesis. A recent report found the successful use of sequential introduction of the LoxP sites to improve efficiency and avoid recombination between alleles [26]. Indeed, a 3 to 10 fold improvement in successful targeting was observed, though it should be noted that such an approach requires a more protracted period of time to completion [26]. Additional work has demonstrated over 5 fold improvement in targeting using a long ssODN [22, 29, 31] or doublestranded donor DNA [28].

In conclusion, we find the 2sgRNA-2ssODN method to be inherently biased for *indels* or substitutions at the DSB, deletion between the guide cleavage sites, or *trans* insertion of the LoxP sites. Even though the overall success rate is very low — \sim 1,200 zygotes were needed to

generate 1 correctly targeted animal — it is possible to generate floxed alleles using CRISPR-Cas9 gene editing technology with 2sgRNA-2ssODN. The method, however, requires two inefficient simultaneous HDR events leading to correct insertion of both *LoxP* sites in the *cis* configuration, an outcome we find occurs very infrequently (<1%).

This work was supported by the National Collaborative Research **Acknowledgements:** Infrastructure (NCRIS) via the Australian Phenomics Network (APN) (to Gaetan Burgio and Paul Thomas), by an Institutional Development Award (PI: Shelley Smith) P20GM103471 (to CBG, RMQ, DWH, JDE and RR), by NIGMS 1P30GM110768-01 and P30CA036727 (as part of support to University of Nebraska Mouse Genome Engineering and DNA Sequencing Cores), the British Heart Foundation (FS12-57), FS12/57/29717, CH/13/2/30154 and the program grant RG/15/12/31616 (to Kathryn Hentges and Bernard Keavney), the Wellcome Trust grants 107849/Z/15/Z and 105610/Z/14/Z, the Medical Research Council, MR/N029992/1 (to DB and CBL), the National BioResource Project of Ministry of Education, Culture, Sports, Science and Technology/Japan Agency for Medical Research and Development (MEXT/AMED), Japan, the Canadian Institutes of Health Research, MOP#142452 (MCB and LJM). LJM is a member of the Research Centre of the McGill University Health Centre which is supported in part by FQRS. Dr William Thompson was supported by the Indiana Clinical and Translational Sciences Institute, funded in part by grant #UL1 TR001108 from the National Institute of Health (NIH), National Center for Advancing Translational Sciences, Clinical and Translational Sciences Award. D Kent Lloyd is supported from the NIH (UM1OD023221), and work contributed by staff from the UC Davis Mouse Biology Program (MBP) is supported by a grant from the American College of Laboratory Animal Medicine. The work contributed from Xiande Liu, Chad Smith, Eric Jonasch.

Xuesong Zhang and Jan Parker-Thornburg is supported from the NIH under the award number P30CA16672. R Sedlacek was supported by LM2015040 (Czech Centre for Phenogenomics), CZ.1.05/1.1.00/02.0109 (BIOCEV), and CZ.1.05/2.1.00/19.0395 by the Ministry of Education, Youth and Sports (MEYS) and by Academy of Sciences of the Czech Republic (RVO 68378050). David Ray was supported from a Wellcome Trust Investigator (107849/Z/15/Z) and the Medical Research Council (MR/P011853/1 and MR/P023576/) grants. Andrew Loudon was supported from a Wellcome Trust Investigator (107849/Z/15/Z), Biotechnology and Biological Sciences Research Council (BB/N015584/1), Medical Research Council (MR/P023576/1). The work contributed from Gloria Lopez-Castejon is supported from the Wellcome Trust (104192/Z/14/Z) and the Royal Society. Pilar Alcaide was supported from the NIH (HL 123658).

Literature Cited:

- 1. Lee, D. and D.W. Threadgill, *Investigating gene function using mouse models*. Curr Opin Genet Dev, 2004. **14**(3): p. 246-52.
- 2. Dickinson, M.E., et al., *High-throughput discovery of novel developmental phenotypes*. Nature, 2016. **537**(7621): p. 508-514.
- 3. Skarnes, W.C., et al., A conditional knockout resource for the genome-wide study of mouse gene function. Nature, 2011. **474**(7351): p. 337-42.
- 4. Thomas, K.R. and M.R. Capecchi, *Site-directed mutagenesis by gene targeting in mouse embryo-derived stem cells*. Cell, 1987. **51**(3): p. 503-12.
- 5. Austin, C.P., et al., *The knockout mouse project*. Nat Genet, 2004. **36**(9): p. 921-4.
- 6. Friedel, R.H., et al., EUCOMM--the European conditional mouse mutagenesis program.

 Brief Funct Genomic Proteomic, 2007. **6**(3): p. 180-5.

- 7. Ayadi, A., et al., Mouse large-scale phenotyping initiatives: overview of the European Mouse Disease Clinic (EUMODIC) and of the Wellcome Trust Sanger Institute Mouse Genetics Project. Mamm Genome, 2012. 23(9-10): p. 600-10.
- 8. Wang, H., et al., *One-step generation of mice carrying mutations in multiple genes by CRISPR/Cas-mediated genome engineering.* Cell, 2013. **153**(4): p. 910-8.
- 9. Jinek, M., et al., A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity. Science, 2012. **337**(6096): p. 816-21.
- 10. Yang, H., et al., One-step generation of mice carrying reporter and conditional alleles by CRISPR/Cas-mediated genome engineering. Cell, 2013. **154**(6): p. 1370-9.
- 11. Haeussler, M., et al., Evaluation of off-target and on-target scoring algorithms and integration into the guide RNA selection tool CRISPOR. Genome Biol, 2016. **17**(1): p. 148.
- 12. Stemmer, M., et al., CCTop: An Intuitive, Flexible and Reliable CRISPR/Cas9 Target Prediction Tool. PLoS One, 2015. **10**(4): p. e0124633.
- 13. Cong, L., et al., *Multiplex genome engineering using CRISPR/Cas systems*. Science, 2013. **339**(6121): p. 819-23.
- 14. Harms, D.W., et al., *Mouse Genome Editing Using the CRISPR/Cas System*. Curr Protoc Hum Genet, 2014. **83**: p. 15.7.1-27.
- 15. Yang, H., Wang, and R. Jaenisch, Generating genetically modified mice using CRISPR/Cas-mediated genome engineering. Nat Protoc, 2014. 9(8): p. 1956-68.
- 16. Bassett, A.R., et al., *Highly efficient targeted mutagenesis of Drosophila with the CRISPR/Cas9 system.* Cell Rep, 2013. **4**(1): p. 220-8.

- 17. Aida, T., et al., Cloning-free CRISPR/Cas system facilitates functional cassette knock-in in mice. Genome Biol, 2015. **16**: p. 87.
- 18. Blake, J.A., et al., *The Mouse Genome Database: integration of and access to knowledge about the laboratory mouse.* Nucleic Acids Res, 2014. **42**(Database issue): p. D810-7.
- 19. Kaneko, T., et al., Simple knockout by electroporation of engineered endonucleases into intact rat embryos. Sci Rep, 2014. 4: p. 6382.
- 20. Chen, S., et al., *Highly Efficient Mouse Genome Editing by CRISPR Ribonucleoprotein Electroporation of Zygotes*. J Biol Chem, 2016. **291**(28): p. 14457-67.
- 21. Hashimoto, M. and T. Takemoto, *Electroporation enables the efficient mRNA delivery* into the mouse zygotes and facilitates CRISPR/Cas9-based genome editing. Sci Rep, 2015. 5: p. 11315.
- 22. Miyasaka, Y., et al., *CLICK: one-step generation of conditional knockout mice.* BMC Genomics, 2018. **19**(1): p. 318.
- 23. Ohtsuka, M., et al., *i-GONAD: a robust method for in situ germline genome engineering using CRISPR nucleases.* Genome Biol, 2018. **19**(1): p. 25.
- 24. Qin, W., et al., Efficient CRISPR/Cas9-Mediated Genome Editing in Mice by Zygote Electroporation of Nuclease. Genetics, 2015. **200**(2): p. 423-30.
- 25. Teixeira, M., et al., Electroporation of mice zygotes with dual guide RNA/Cas9 complexes for simple and efficient cloning-free genome editing. Sci Rep, 2018. **8**(1): p. 474.
- 26. Horii, T., et al., Efficient generation of conditional knockout mice via sequential introduction of lox sites. Sci Rep, 2017. 7(1): p. 7891.

- 27. Bishop, K.A., et al., CRISPR/Cas9-Mediated Insertion of loxP Sites in the Mouse Dock7

 Gene Provides an Effective Alternative to Use of Targeted Embryonic Stem Cells. G3

 (Bethesda), 2016. 6(7): p. 2051-61.
- 28. Kueh, A.J., et al., An update on using CRISPR/Cas9 in the one-cell stage mouse embryo for generating complex mutant alleles. Cell Death Differ, 2017. **24**(10): p. 1821-1822.
- 29. Lanza, D.G., et al., Comparative analysis of single-stranded DNA donors to generate conditional null mouse alleles. BMC Biol, 2018. **16**(1): p. 69.
- 30. Pritchard, C.E.J., L.J. Kroese, and I.J. Huijbers, *Direct Generation of Conditional Alleles Using CRISPR/Cas9 in Mouse Zygotes*. Methods Mol Biol, 2017. **1642**: p. 21-35.
- 31. Quadros, R.M., et al., Easi-CRISPR: a robust method for one-step generation of mice carrying conditional and insertion alleles using long ssDNA donors and CRISPR ribonucleoproteins. Genome Biol, 2017. **18**(1): p. 92.

Figure Legends:

Figure 1: Schematic of 2sgRNA-2ssODN CRISPR method of creating conditional knockout

alleles. (A) Wild type locus showing exons 3, 4 and 5 of a hypothetical gene where exon 4 is

chosen as a target exon for inserting *LoxP* sites. The guide 1 and the guide 2 target introns 3 and

4 respectively. (B) CRISPR components showing 2sgRNA-2ssODN donors and a Cas9 source.

(C) Delivery of CRISPR components into zygotes via microinjection (n=17,557) or

electroporation (n=330). (D) The conditional knockout (cKO) allele showing target exon (#4)

with flanking *LoxP* sites.

Figure 2: Quantitative assessment of the 2sgRNA-2ssODN methods (A) Doughnut graph

indicating the methods of zygote injections (pronuclear, cytoplasmic or both) of the CRISPR

reagents. Numbers indicate the percentage of the total zygotes microinjected or electroporated.

(B) Doughnut graph indicating the form of delivery of the CRISPR reagents (mRNA, protein or

plasmid) in the zygotes. Numbers indicate percentages. (C) Flow chart indicating the number of

successful edited alleles and correct LoxP insertions out of the number of live born pups from

microinjected and transferred zygotes. Numbers indicate absolute numbers. (D) Doughnut chart

indicating the editing types observed amongst the live born pups genotyped on a sub sample

from 25 loci. Numbers indicate absolute values.

Figure 3: Desired and undesired outcomes of the 2sgRNA-2ssODN CRISPR method of

creating conditional knockout alleles. (A). Desired outcome showing a floxed allele and its

occurrence is <1%. (B) to (F): various undesired outcomes including only one LoxP site insertion

(B), only *indels* created at one or both sites (C), combination of *LoxP* insertion and *indels* (D), deletion between the two cleavage sites (E) and no *indel* or no insertion events (F).

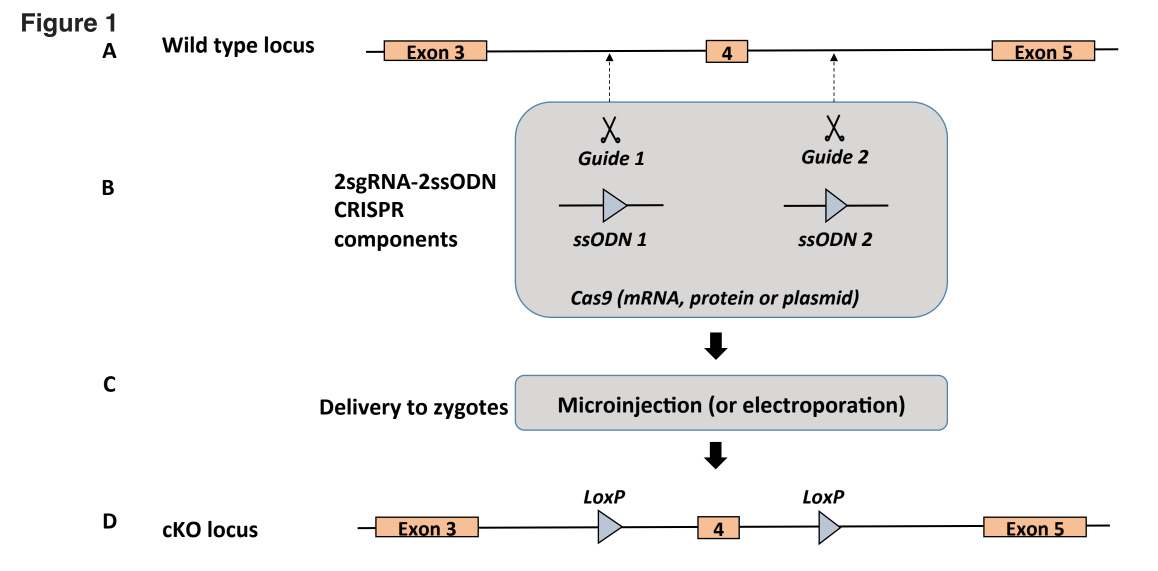
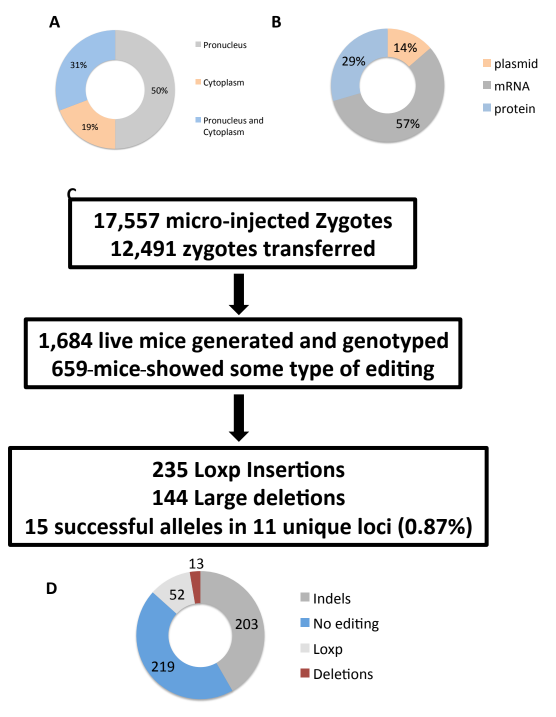


Figure 1: Schematic of 2sgRNA-2ssODN CRISPR method of creating conditional knockout alleles. (A) Wild type locus showing exons 3, 4 and 5 of a hypothetical gene where exon 4 is chosen as a target exon for inserting *LoxP* sites. The guide 1 and the guide 2 target introns 3 and 4 respectively. (B) CRISPR components showing 2sgRNA-2ssODN donors and a Cas9 source. (C) Delivery of CRISPR components into one-cell stage zygotes via microinjection (n=17,867) or electroporation (n=330). (D) The conditional knockout (cKO) allele showing target exon (#4) with flanking *LoxP* sites.

Figure 2

Figure 2: Quantitative assessment of the 2sgRNA-2ssODN methods (A)

Doughnut graph indicating the methods of zygote injections (pronuclear, cytoplasmic or both) of the CRISPR reagents. Numbers indicate the percentage of the total zygotes microinjected or electroporated. (B) Doughnut graph indicating the form of delivery of the CRISPR reagents (mRNA, protein or plasmid) in the zygotes. Numbers indicate percentages. (C) Flow chart indicating the number of successful edited alleles and correct LoxP insertions out of the number of live born pups from microinjected and transferred zygotes. Numbers indicate absolute numbers. (D) Doughnut chart indicating the editing types observed amongst the live born pups genotyped on a sub sample from 24 loci. Numbers indicate absolute values.



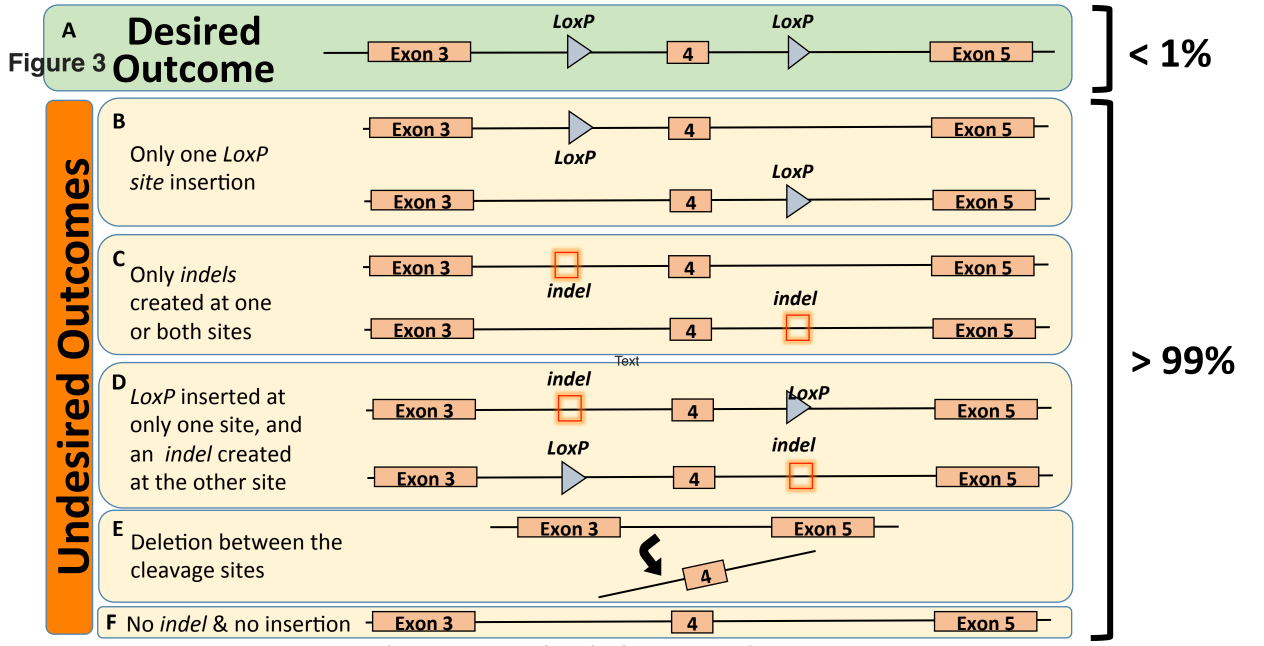


Figure 3: Desired and undesired outcomes of the 2sgRNA-2ssODN CRISPR method of creating conditional knockout alleles. (A). Desired outcomes showing a floxed allele and its occurrence is <1%. (B) to (F): various undesired outcomes including only one *LoxP* site insertion (B), only *indels* created at one or both sites (C), combination of *LoxP* insertion and *indels* (D), deletion between the two cleavage sites (E) and no *indel* or no insertion events (F).

Laboratory (name)	Gene	5' guide RNA	3' guide RNA	5" donor sequence	3' donor sequence	Cas9 concentration (in ng/μl)	SgRNA concentration (in ng/µl)	ssODN concentration (in ng/μl)	Exons floxed	distance betwee targets
	Lcmt1	5'-ACAAAAGTTTACCCTCACGC-3'	5'-AAGAGGTTTTGTATCTATCC-3'	S- ATCCATCTGGGCTGCACAATAATACCTTGTCTCAAAAGAA CCAAAACACCAACAATAACTTGGTATAGCATACATATAC GAAGTTATTGAGGGTAAACTTTTGTGCCGTTTTATAGGT TTAGTAACAACTGATAGGCTTTTAAGTA-3 TTAGTAACAACTGATAGGCTTTTAAGTA-3	CTCCTCCCCACTATCCCACTATAACTTCGTATAGCATACAT	mRNA (IVT)=100 ng/μl	(IVT) 20 ng/μl	100 ng/μl	Exon 3	8-900 bp
ixperimental Animal Division,	Trp53	5'-AGGACCCAAGATGGCCGTGA-3'	GCCCCGGATTAGGTCCCCAG	GCCTGAATAAAAGACGGAAGAGCTGCCCCATTCCTGCTT CTCTGGAAATGGTGTCCCTCAATAACTTCGTATAGCATAC ATTATACGAAGTTATGGCCATCCTTGGGTCCTGACTTCTTCT CAAAGGAGCCTGGCCGACTTCTTGGATACTTGTA-3'	TATAACTTCGTATAATGTATGCTATACGAAGTTATGGGGAC	Protein = 100 ng/μl	synthetic 75 ng/μl	25 ng/μl	Exon 2 to 9	4kb
RIKEN BRC, Japan	Trf	5'-GTACTGCTTGTGTCCCCGGG-3'	5'-GTGAGGCACTCGGGGTATGT-3'	5- AACAGAGTGGGCAAAGTCTAGGGCAAGCCTTGTTTAGC AAGAAGTACTGCTTGTGTCCCCATAACTTCGTATAGCATA CATTATACGAAGTATGGGAGGGTAACTGAAGAGCAGGG GTTAACTTATTCCTGTCTAGAAACCCTGCTTAGTACATGA	GAAGGGACCTACAATAACTTCGTATAGCATACATTATACG AAGTTATTACCCCGAGTGCCTCACAGTAGACCATGCTCA	mRNA (IVT)=100 ng/μl	(IVT) 20 ng/μl	100 ng/μl	Exon 5	5-600 bp
	Ppme1	5'-AATCCTTATCTAGGGGATTC-3'	5'-ACTAGCTAATGAATTCATGC-3'	CTCAAGTGTCATGTTGGTGTAATTCCTGAGCCAAAATAGT ACTTCCAACTATAACTTCGTATAGCATACATTATACGAAGT TATTTCTGGCTCTAAACTAAA	GGGTATGTATCTTTTATAGTCTTCTGTTTGGAGAAATATTT GAATATAAGTAGCCTTGAATAACTTCGTATAGCATACATTA	шкла (тv т)=100 пg µг	(1V 1) 20 ng/µ	100 lig jii	Exon 4 and 5	1.1 kb
	Trmt6	5'-GACTGAGATCTCCGATAGGAAGG-3'	5'-CCAGCGTTATGCTCTCAACATCC-3'	CTGTCGTGTGCTTGTCTTTGAAGTTGCTCTAAGAGACTG AGATCTCCGATGAATTCATAACTTCGTATAATGTATGCTAT ACGAAGTTATAGGAAGGCTAATGCCTGACCCTTGGCAGT	GGACCAGCGTGGATCCATAACTTCGTATAATGTATGCTATA				Exon 2 to 8	4.8 kb
	Slc7a14	5'-CATCAGCAGGACTATACCCCAGG-3'	5'-TAGCCATGGATGTATTGGTCAGG-3'	GGGCCATGGATGGTCCGAGGCACCAGGACCTGTTCATCA GCAGGACTATAGCTAGCATAACTTCGTATAATGTATGCTAT ACGAAGTTATCCCCAGGCATCTTCTCTCTGCAACCCCACA	ATGGATGTATTATAACTTCGTATAATGTATGCTATACGAAG				Exon 4 to 7	15.6 kb
	Tmem163	5'-AACACTCATCCGATACTGCCAGG-3'	5'-CCCAATACCATGCACCACGCTGG-3'	CCTCTAACTCAGAGAGCTTCTGCTACCCTGGGGAAACAC TCATCCGATACATAACTTCGTATAATGTATGCTATACGAAG TTATTGCCAGGGATGCCAGAATTTTCGATAATTTTGTTTTG TTTTTAACGTGCA-3'	ATGTCCAGCGTATAACTTCGTATAATGTATGCTATACGAAG				Exon 2 to 7	173.5 kb
Osaka University, Japan	Wtap	5'-TGGATTATCACTACAAGCCTGGG-3'	5'-CCAAGGTACTAATAGCATGCAGG-3'	ATCCTCACCACAGGGCACTTTGATGCCCTCCCTCCCTCC TTCCCCAGGCATAACTTCGTATAATGTATGCATATCGAAGT TATTTGTAGTGATAATCCATTTTCCAGTTTCTTTAACTTCC TGAGTGCCTCAG-3*	GTACTAATAGCATAACTTCGTATAATGTATGCTATACGAAG	mRNA (IVT)=20 ng/μl	(IVT) 25 ng/μl	50 ng/μl	Exon 3 to 7	15 kb
	Mettl3	5'-AGGTGATCTAGAGCTAACGCTGG-3'	5'-CCCAAGGATCGTCAGCTATTCTG-3'	5°- TTGGCAAAACAGCAAGTGCTGCCATGTGAATGAAAGGT GATCTAGAGCTAATAACTTCGTATAATGTATGCTATACGAA GTTATACGCTGGTCAGAGACCCTGCTTGAAGTGAAAGAT GTGTGTGCTAGCGATG-3'	TGCCCAAGGATAACTTCGTATAATGTATGCTATACGAAGTT				Exon2 to 10	4.6 kb
	Mettl14	5'-TTTCATTCACCATTGGCGACAGG-3'	5'-CAGAATAGCTGACGATCCTTGGG-3'	5'- CTITICTGTGTTACTGCTTCTGATGCCAACTTGTATTTCATT CACCATTGGATAACTTCGTATAATGTATGCTATACGAAGTT ATCGACAGGGAATCATCCTAATTTCAAATTTAACCAAATG CGTAAAATACTT-3'	TCTGCACGCTATAACTTCGTATAATGTATGCTATACGAAGT				Exon2 to 9	11.7 kb
	Klotho	5'-GGCCACAGGATTGTGCGATGTGG-3'	5'-CTTCCCTTTGGTGTTACGGCTGG-3'	5: GTGGACGCGGGGAGTGGGCGACGCGGGGACATCTCAGG ATGGAGGCCACAGGATTGTGCGATAACTTCGTATAATGTA TACTATACGAAGTTATATGTGGAATAGTCTGCTCCCTGAG CTGGCTGCAGCAGGTGCTTGTTCTCCGACGTCCCTA-3'	GCACTTCCCTTTGGTGTTACATAACTTCGTATAATGTATAC TATACGAAGTTATGGCTGGGGTCCCTGCTCAGGAAGTTA				Exon1 to 3	30.6 kb
Tokai University, Japan	Arhgef16	5'-CGACATCAGAGGTCGGCCGA-3'	5'-CTCACTTTGCGGTTACTAAC-3'	5: CATAGCTCAGGATCACCCACTGGCCTCACATACCAGGCG GCCATGACCTAGAGAACCCTCATAACTTCGTATAATGTAT GCTATACGAAGTTATGTCGACCCGACC	CTTGCTCACTTTGCGGTTACGAATTCATAACTTCGTATAAT GTATGCTATACGAAGTTATGATGCTGTAGCCTGCAGACAC	mRNA (IVT)=20 ng/µl	(IVT) 100 ng/μl	100 ng/μl	Exon 7 and 8	1 kb
	Cd226	5'- AATGTCTTTCTAACTAGATC-3'	S-GTTTACGACATTATTCGTTG-3	5- TAACATTGAAGGAGTTCTGTAAATAAGTATTTGTATATGTG TTCAATGTCTTTCTAACTACTCGGAGTAACTTCGTATAGCA TACATTATACGAAGTTATGATCAGGGATCCCGTAGACTGT AAAATACCTCTCTATTATGGTCATACATACCTCCTGGT-3	ş: Cacttigcactcagaaatgacagcaattaagaaaaaaatg Agatiggittacgacattaticataactitggiataatgatg Ctatacgaagttatgicgacgitgaggaaaaattagatca Aaaatctagcaatggaattgggaaaagaatacaataaca T.y'				Exon2 and 3	2.8 kb
	Igfl	5'-CGAAAGGAGCCGAGGATAGG-3'	5'-TCTAAAGAGCCGAAAATGCA-3'	5- ACTATTGTTCCATTTTATGGGTGAGGAAGCAAGTTCACAT GGTCGAAAGGAGCCGAGGATATAACTTCGTATAGCATACA TTATACGAAGTTATCTCGAAGGGGGTTGAGCTCACATCA ACTGTTGTGATTTTTACCTGATTTTTCAATCTTCTGCTC-	CCTCTAAAGAGCCGAAAATATAACTTCGTATAGCATACAT TATACGAAGTTATCTGCAGGCATGGATGTTGTAGAGATCC				Exon 4	1.4 kb

Laboratory Animal Resource	Pik3cb	5'-ACTTGATCATCGCAGGAGTT-3'	5'-GGTACTAGCTGCCGGCACTT-3'	5. CAACCCCGAGTCTGGTTTTGTTTTGTTTTGTTTTAGCCAT CCAAACCTGATCATCGCAGGAATAACTTCGTATAGCATAC. TITATAGAAGTTATGATTCGTTGGGAGAAGCATAGGTGG TGCCTATCTGAGGGAGGAGCCAGGGAGCATGGGTG TGCTATCTGAGGGAGGAGCAGGGAGCATGGTTG TATATCGAAGTTATGATTCGTTTGGTTTTCACTTGGTCT TGCCTATCTGAGGGAGCAGGAGCAGGAGTGTGTT T,3' E,3' E,3'	2 3	i∩⊫Spg/ul	10 ng/μl	Exon22 and 23	3.8 kb
Center, University of Tsukuba	Gata3	5'-GACAAATCCCAATATAGCTG-3'	5'-GGAAGCCAGAAGTTGCTATC-3'	GTAAAATAATACAATTCATAACAAAAATAATAAAAAGCTA ATAGCAATAGAAATCTAACTAGGATAGTGTGTGTGTATGTA	C A Γ	o) sign	10 ду	Ebhancer	7 kb
	Gen111	5'-AGAGCCCTCACCTATCCTAT-3'	5'-TCAAATTCCTGGCACACCGA-3'	\$. GTGGGAGAGGGTGCAGAGGAGGAGGGCGTGTTTCCCA GTGGGAGAGGGCTCACCTATCCATAACTTCGTATAGCATA GCACCAGAGCCCTCACCTATCCATAACTTCGTATAGCATA CATTATACGAAGTTATGAATTCTATTGGAGACTAAACATTT TTATATCGAAGTTATGATATCCATAGGAGGTTTAGAGTTAG AC.3 CATCAGACCCTCGCCCCTCAGCACCTACGCTCTCAGGCCACCTTACACTTCAGGAGGAACACTAAGGAGGTTAGAGTTAGATACCCACCTGGCCCTCTAGACCTGCTCTAGACCTGCTTACACCTGCCCCTCTAGACCTGCTCTAGACCTGCCTTACACCTGCCCCTAGACCTGCTCTAGACCTGCTCTACACCTGCCCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTCTAGACCTGCTAGACCTGCTAGACCTGCTAGACCTGCTAGACCTGCTAGACCTGCACACACA	A.			Exon 46 to 53	4.8 kb
	St3gal5	5'-TAAGTACAGTCAGCCAACTT-3'	5'-TATGTGTGGGCTGCCGTCAC-3'	5'. TCCTTGTGTTTTCTATAGATCTGACCACTATGCATACTGCA GGTAAGTACAGCCAAATAACTTCGTATAGCATACAT GGCCAAGTGTCCAAACTGCATGCTATAACCTCAGATCCAA GGTAAGTACAGTCAGACCAAATAACTTCGTATAGCATACAT TATACGAAGTTATGAATTCCTTAGGGCCCATCTTCTGGTG ACAGTTTGCCTCAGTGGCATAATGAAAGTAATGGCCTTG AGAAAGGAGGTTTGGAAAGGTGGAGGGGGGGGGG	G G			Exon 4	8-900 bp
	Irf3	5'-CATGAGTTTGTCAGCACCGT-3'	5'-GATGTTTCCTCAGCTGCTT-3'	5. CAAGTAGATAACTAGAATGATGGCTTAAAAAAAAAAAA CGGGCAGTGAGGGAGGACGGTGGGCTCGCAGCTCAGGAG CCTTGAAGGTGGCACATGAGTTTGTCAGCAGC TGGCAGGGCAG	C			Exon 3 to 6	1.9 kb
TIGM, USA	Phactr1	5-CCACCCCGACTTGGTGTGCCAA-3'	5'-TAACGTGTATATCCGTACTAGG-3'	5'. ATAACCCTGACCCACGCAGTGTAATCGGTCATGAGTGGA CCCACCAGCCCACGCACCACACACTCCCCCAATAACTTC GTATAATGTATGCTATACGAAGTTATGAATTCCCCGACCT GGTGTGCCAAAGTCACACTCCATAACCCACTGAGTCCAC AAAAAAAACTGCTCAATGACAG-3' CATGTGTGGTTAGTCATCCATGAACGACTTGCACACACTGAGTCGAC CATGTGTGCTAACAACATTCCCTAACACACTGATCCACACACTGAGTCGAC CATGTGTGGTTAGTCATCGTGA-3' CATGTGTGGTTAGTCATCGTGA-3'	C r	(IVT) 100 ng/μl	100 ng/μl	Exon 7	1.6 kb
110.11, 0.01	Actrt2	5-CCCAAGTTGAGTGCCAGGTCTC-3'	5'-CCACCTGAATGCTACTGTTGAA-3'	5'- TATGAACCCACCTITCCAAGGTICCCTICATCATCAGGA TATGAACCCACCTTTCCAAGGTICCCTICATCATCAGGAG AAAGGGAAACTGTGTCCAGGCTCAGCCCCCATAACTTCG CCCCCATGGGCCATCACACAGCACGAGCTACCAATAACTT TATAATGTATGCTATACGAAGTTATGAATTCAAGTTGAGTG CCAGGTCTCAGAGGCTACCAAGTCCCTCTGTGTGTGTGAG CCAGTGTCAAGTTGCATCTCAAGATGCTCTAGGTGCC CCCATAGTAGGTGGCTCCTCC-3' AAGGCTGGGCCCATTCTCCCG-3'	3	(111) 100 ng pi	100 ng pi	Entire Gene	2.6 kb
	ApoA1	5'-CCTTGTAACCAGCACCGACCAC-3'	5'-CTAGCCGAGTTTCCAGGTGGGG-3'	CCCCAGTACTGGTCAGACAGCACCCAAAACAAACAAA CACAAAACAAA	G			Exon 4	1.6 kb
University of Adelaide, Australia	Depdc5	5'-GATAGGGATACTGGTCTTCT-3'		5. CCTTGGGGACAAGAGTTTTACCTACTCACAGTCTTCATA TTTGATAGGGATACTGGTGAATTCATAACTTCGTATAAC GTTAGCGAGAGTTATTCAGAGTCTACCAAGCTAT ATCTGTCTTATAAAACACACACACACCACCTATATACA.3' TIGCT3 TGGCT3 TGGCTAGTAGAGAGTTAGAGAAGAAAAAAAAAAAAAAAA	MRNA (IVT)=100 ng/μl Γ (Scr7 1 mM)	(IVT) 20 ng/μl	100 ng/μl	Exon 4	2.1 kb
University of Rochester, USA	Akap12	5'-AGGATTTCAGACACGAA <i>TCA-3'</i>	5'-AAGACTAGAGAACGGCGGTC-3'	2- CAGGCCTCAGGCTCAAGCTAACCCTCCAACTTCAGCCAC CGGATGCCCCACCGGCCATCCCGCAGGCTGCCAGCCCA TGCAGCATCAGGATTTCAGACAATAACTTCGTATAATGTA TGCTATACGAAGTTATCGAATCATAACACCCGGCTCACCT AGCATACATTTAACGAAGTTATCGGAGTTATCGGCGTCAACCAGAT TGAC-3' TGAC-3' CAGCCCCCCACAAA-3' CAGCCCCCACAAA-3'	MRNA (IVT)=50 ng/μl	(IVT) 100 ng/μl	100 ng/μl	Exon 1	1.5 kb
	Cav3 (design 1)	5-TGCCTACAAAAAGGGTCCTT TGG-3'		ACAGITEATIGTGGCTGAAAACACATGTTGGAGGAAGGA ACAGITEATIGTGGCTGAAAACACATGTTGGAGGAAGGAA ATTCACCTCAIGGCAGGGAAGTGAAAGAGAAAGAAGAAA TGCACCTAGGTGCACACCAAGATGCAACACACAGCACGTGTGCAAGAAGGACACTTTGGGTCACCAGTAGATGCCTACAA TTCCAAATTTCCAAGTTCAGCAACACAAGCACGTGGTGCAGGAAGGCACCACAGCACGTGTTGCAGGTAGCACCACAGGCACTTTTCCAAATTCCAAGTCCAACACAACAAATGAGCACCACCTAGGACTCCAGCTTCTAGCAGCACTCTACCATCCAGCACTCTACACCACACACA	G G A	(IVT) 50 ng/μl	50 ms/ml	Exon 2	1.9 kb
	Cav3 (design 2)	5'-ACATGCCAAACCTACCCATC TGG-3'		AGCTECTTCCTCCCCCCCCCCCGAGTCCCTTACTCTA TCTCCCATACAAGGAGATTATTTTCAAATGACTTCTGCTT TACCTAACCTCTTTGGTTGACCAATTCCATTTAAAGAGG TCCCTGAGGCAGGTGAGAGTACCGAGGAGGAGACAATCCATCC	G G G	(141) 30 пд µі	50 ng/μl	Exon 2	2.2 kb
	Cav3 (design 3)	5'-CTTGGGATCCCACCGCAGTT AGG-3'		5. AGGGGTAGATCTCACTGAGAGTCATCACCTTCGCTGAGG GATGCAGCATAGCCTTGGGATCCCACCGCAGAATTCATAA GGAAAACACGGGCACCCTGCTCCCCTCAGCCTCCAGCCTCTCCTTATAGCATACATTATACAGAAGTTATGTTAGGGTAGC CTTCGTTATAGCATACATTATACGAAGTTATGTTAGGGTAGC AGGAAGACCTGGGATGAGTCATGAGTGCTGGATTGATCAC TGGTCCTGGTCCAGGCTAGAGCTGAGAGTCTACAC CGGCGCCCCCTCACCTCA	A mRNA (IVT)=25 ng/μl	(IVT) 60 ng/μl	50 ng/μl	Exon 2	6-700 bp

				5'-						
	Cav3 (design 4)	5'-CCTGCTACCCTAACTGCGGT GGG-3'	5-GGCGGCCCTGACTGAGCGGAGGG-3	TGCCTATCAAGGGGTAGATCTCACTGAGAGTCATCACCTT AGACAG CGCTGAGGGATGCAGCATAGCCTTTGGGATCCCACCGAAT CTCAGCC TCATAACTTCGTATAGCATACATTATACGAAGTTATGCAGT GCTTATA TAGGGTAGCAGGAAGACCGTGGAGTAGTCATGATGCCTG GGGTTCC	CCTCCCATGAACCCCGGCGGCCCTGACTGAGCAA TAACTTCGTATAGCATACATTATACGAAGTTATGGA	mRNA (IVT)=15 ng/ μ l	(IVT) 60 ng/μl	50 ng/μl	Exon 2	7-800 bp
Maine Medical Center Research	Cav3 (design 5)	5'-CTTGGGATCCCACCGCAGTT AGG-3'	5'-TCAGGGCCGCCGGGGTTCAT GGG-3'	5'- AGGGGTAGATCTCACTGAGAGTCATCACCTTCGCTGAGG CAGTAC. GATGCAGCATAGCCTTGGGATCCCACCGCAGAATTCATAA TAAGGA. CTTCGTATAGCATACATTATACCAAGTTATGTTAGGGTAGC CGAGGC. GAGGACAGCCGTGGAGTGAGCTCGGATTGAGCC GGAGGC TGGTCCTGGTCCAGGCTAGAGCTCCTACCTG-3' TGGGCTC	AACCCTCCGCTCAGTCAGGGCCGCCGGGGTTATA GTATAATGTATGCTATACGAAGTTATAAGCTTCATG	mRNA (IVT)=25 ng/μl	(IVT) 60 ng/μl	25 ng/μl	Exon 2	6-700 bp
Institute, USA	Icam1	5-TACCTCAATTCACTTAGTCTCGG-3'	5'-AGGCAGGAGTCTCATCCAGCAGG-3'	GGCAGGTACAGCTGTAATTCCAGCCAAAGAAGGAGACT GTTACAG GGCAAGGCTGAGGCTAGCCGAGAATAACTTCCTAITAATG TATGCTATTACGAAGTTATCTGCAGCAGCTAAGTGAATTCAGGT TATGCCTATCAGAGTTACTCGAAAACTCCAAGAC TGCAGCAT TAATCCATGAST	AGAAGGCTCAGGAGGAGGCCATAAAACTCAAGG GCCCACCTCCCTGAGCCTGCTCGAGATAACTTCG CATACATTATACGAAGTTATGGATGAGACTCCTGCC CCCCTGCAGGGCAACAGCTGCTGCTGCTTTTGAA	mRNA (IVT)=25 ng/ μ l	(IVT) 60 ng/μl	50 ng/μl	Exon 3 to 7	3.5 kb
	Dock7 (design 1)#	5'-ACGCTCCACACCCTGACCCATGG-3'			TATTCTTGCAGAGGACCTGGGTTTGGTTCCTAGCT TTCGTATAGCATACATTATACGAAGTTATCACTGGC	nRNA (IVT)=60 ng/µl for pronuclear; 100ng/µl for cytoplasmic	(IVT) 15 ng/μl each for pronuclear; 50 ng/μl each for cytoplasmic	50 ng/μl	Exon 3 to 4	2.1 kb
	Dock7 (design2)#	5'-ACGCTCCACACCCTGACCCATGG-3'	5'-CCAACCATTGAGCTAGCTTAGGG-3'	5. TCCATCTTGGGCCTTTTAACCATGTGCAAAGACTCTCCCT TTCCAGC TTACTTTCCATTACCATGTGCAAGTGGCCATGA TAACTTTGGTATACCATACATTATGCAAGTTATCTCAGGGT TTGGTAG TGGGAGCGTTTTGGGAGCTTTACCTGCCACAGTCTTGGTC ACAGTATTTGCTAGAATGATCCCCAGTAACAAC-3* CCTCCAC CCTCCAC CCTCCAC CCTCCAC TCCAC	AATGTACAGCCTTATTTTTAATATACCCTAAATAAC ATAGCATACATTATACGAAGTTATGGATCCAATGGT	mRNA (IVT)=100 ng/µl for cytoplasmic	(IVT) 50 ng/μl each for cytoplasmic	50 ng/μl	Exon 3 to 7	5.3 kb
	Gpr180	5'-ATCCCTGGCCGGAGCACCAT AGG-3'	5'-ACTAGTCAGGGTTGCTTACA TGG-3'	GGAICTTIACTGGCCTGCTCCTCTTGCCTTGCTCAGCCTG GAGGAG CTTTCCTATGGATCCATAACTTGGTATAGCATACATTATAC CCTCGC GAAGTTATGTGCTCCGGCCAGGGATGGAACCGCCCACA ACGAAG TGGGCTGGGCCCTCCCCCA-3' CAACATT	CCATGTGGATCCATAACTTCGTATAGCATACATTAT	mRNA (IVT)=25 ng/μl	(IVT) 60 ng/μl	50 ng/μl	Exon 3 to 6	11.6 kb
	Noct	5-GAATCAAAGCGCGGCCGCTC AGG-3*		TITATITCHTIGGCAATITCATATACAAATACCATGTATGTTA AGGTCA ACCTAGATCCACCCCCGTCCACCCTGAGGATCCATAACTT ATGTTTC CGTATAGCATACATTATACGAAGGTTACGACCGGCTTTGATATAGACT TITCTATCCCCTGGGGAATGATCCTGTTCATTCGTAAGGAG CAACAGA AGGAAGCTCATTT-3' GGACAC	TGTGGGCCTGATGCACTCACTCGCTCATAACTTCG CATACATTATACGAAGTTATGGATCCACCAGGGGC	mRNA (IVT)=60 ng/μl	(IVT) 30 ng/μl	50 ng/μl	Exon 3	2.8kb
	FoxD1	5'-GCTCACTGGAATATTGAGCG AGG-3'	5'-AGATGGACAAAGCTTGGACT TGG-3'	5- TITCATCAGATTCCACAGGCGGAGACTGGCCCAGTCCAG AGAGCTGGGACACCCTACCAGACTCCCTCGCGGATCCAT AACTTCGTATAGCATACATTATACGAAGTTATTCAATATTC CAGTGAGGACACCTCACACACATTAGAGCAGTTGTCCCCAGT GGGATGACTCCTACAGGCCACTGG-3' AATTTAA	CCGTTACTTTGCAGATAATTCCAAGTATAACTTCG CATACATTATACGAAGTTATGGATCCAAGCTTTGTC	mRNA (IVT)=50 ng/μl	(IVT) 60 ng/μl	50 ng/μl	Exon 1	3kb
MD Anderson, USA	Atg7	S'CCATCCAACAGGGTTGAGCCCCC-3'	5'-GCCACTGGGGCTCGAGACAAAGG-3'	S- AGGCTTAATGATCTTCATAAGGTGCTAGAACATGCAGGTG AGTGTC TCCTATTGGTCAGTGGGTGCCATCCGTCGACATAACTTCG AACTAAA TATAGCATACATTATACCAGGGTTGAGCCC CCAAGCAAGGAGCAGCATGTTTTTCATTACATTGAAGTTT TTATAGCATT-3* TGGATG. S- ACTAAA ACTAAA ACTAAA TATAGCATACATTATACAGTGAACTTT TTATAGCATT-3* TGGATG. TGGATG.	AAAGCCACTGGGGCTCGAGAGTCGACATAACTTC GCATACATTATACGAAGTTATCAAAGGGATTCCTTT GTTTTCCTTTTCTTTATTTAATTAATAGATTTGTTCC	mRNA (IVT)=10 ng/μl	(IVT) 15 ng/μl	10 ng/μl	Exon 16	1.3 kb
	Nr1d1	5'-CAGCTAGGGTCTTAGTTACAAGG-3'	5'-AACTTGTAACACCGCTAGCTCGG-3'	5. TITTITTITGTGTGTGTGTATGAATAACTTGCAGCTGTCC AGGGCT ATCTCCCCAAGTCTACCCATAACTTCGTATAGCATACA. CTGAGG TTATAGGAAGTTATGGTACCAGACCCCTAGCTGTCAGT ATCACTT ACTCCCCAGGAAGCAGCTAGCTCAGGGGTTCCCATGTAT AGGGAAG GAGATG-3' AGTGCA'	GAGATAGAACTTGTAACACCATAACTTCGTATAGC TTATACGAAGTTATGGTACCCACCTCAGGGTCTGT				Exon 2 to 6	3 kb
	Genomic region floxing, BP1-BP2	5'-TGTGTAATTTCAGCAAATGAAGG-3'	5'-GGAGATGTGTTGCAGATCAAGGG-3'	5'- ATTICTACCAACTGGGACCAGCATATGAGCCTATAGGAGC CAATCTCATTCAAATGGCCACCTTTCACTTCCTGCTGCT TGCTACCTTCAATAACTTCGTATAGCATACATTATACGAAG AGTAGT TATATGATATCTTTGCTGAAATTACACATGGGAGTTACTGCT AGTTATC TAT-3' GCAGAG	ITCAATTGTCCCTCCATGAGAGAAGCAGAGGGAG ITGCAGATATAACTTCGTATAGCATACATTATACGA ICCGCGGCAAGGGAAAGTTCATGCGACTGCAGAG				N/A	236 kb
University of Manchester Transgenic Unit, UK	Lrrc8a	5'-GTCTAGTTAGGGACTCCTGGGGG-3'	5'-ACTACCCCATTACCTCTTGGTGG-3'	5- GTCCTTGACTTGCTGTTTACCGCTCTTCCCCACACCAC GAGGGC AGTTATCCACAGGAAGTTACCCATAACCTCCCTCGTGCAC AGTTATCCACAGGAAGTTACCCATAACCTCCATATAACGAAG GCCCCCCCAATAACTTCGTATAGCATACATTATACGAAG GCCCCC TATAGGCACCTAACTAGACCTGCTGTCTCTCC AAGTTAI ATAGCCCTGTCTACACCT-3' GGGCAG	GCCCCTGTGCACCAGCTCTGTGTGTGACTGCAAA CACCAAGAATAACTTCGTATAGCATACATTATACG	Protein = 20 ng/μl	synthetic (RNP) 20 ng/μ	50 ng/μl	Exon 3	3.8 kb
	Usp7	5'-TTATACTTTTGTATGTACGTGGG-3'	5'-TGATAGGCACTTCCATACTAAGG-3'	5- ACAGAAGTITTAAGCTIGAAGGCCTGTCAGCCCTGTGCT GGAACC CTACTGTGCTCCTAGCACAACAGGATGGCCTCTCCCTCCT CAGTCCCACGTAATAACTTCGTATAGCATACATTATACGA AGTTATGGTACCCATACAAAAAGTATAAGTCCTTCTTTTTC1 AGTTATC TCACT-3' GTCCATT	AGCCCCAGCCCTGATTATAAAGAGTGTAATAAAA ACCTTAGTATAACTTCGTATAGCATACATTATACGA IGGTACCATGGAAGTGCCTATCAGCCTGGACTGG				Exon 3	7-800 bp
	Bin1	5'-GCACAAACAGGTAACCTTAGCGG-3'	5'-GACACATGTCCCCAAGCAAAGG-3'	5- GACCTGCCCTCAATCTGCACTGCATGTCCCCTGTCTGCCCT GGGTCCCTGTCTGCTCTGC	CTAGTATCCCAAGTCCGCCATGCCCTCCTGACACA CCAAGCGGATCCATAACTTCGTATAATGTATGCTAT GTTATAACAGGACACACGGGCAGGTTAGGGGTTA				Exon 10	5-600 bp

	Meox1	5'-TATACCCCAGACCACACTAC-3'	5'-TACATGGGATGGGTTCACAC-3'	5- AGACAAAGAGAACCTTTTTCTTTTTCTGGAGCATTTCTTT CTTTTATACCCCAGACCACACGTCATAACTTCGTATAGCAT ACATTATACGAAGTTATTACTGGGCTGCCAAGATGAGACA CAGACAGACAGACAAACAGACACACACACACAC	GTGTGAGAAGTGTGACATACAGAGTCAGGAAAGATTTGG AGGATACATGGGATGGG				Exon1	1.6 kb
Lillehei Heart Institute Regenerative Medicine and Sciences Program, University of Minnesota, USA.	Meox2	5'-GGTTCCAGCGTAAACACATT-3'		5- CCCTGGCTCTAGGTTTTCTAACAATCTATTCTCAGATACCT AAAAGAGAATAACCTAATATTATAACTTCGTATAGCATA CATTATACGAAAGTTATGGTTTAGCCTGGAACCGTGAAGTC	CATTATACGA AGTTATA ACGTTGGGAGGGGTGTGCCCATT GGGATGCTGCATCCGCTGGGCATGCATGCCTGCGTGCGT-	mRNA (IVT)=30 ng/μl	(IVT) 10 ng/μl	10 ng/μl	Exon 1	1.5 kb
	Vezf1	5'-GAGAACAGTTCATAGGCTCC-3'		5'- GGGGGAGGGAGGGAGAGAGAGAGAGAGAGAGAGAGAGA	CCAGTGTTTAGTTCCAATTGATAACTTCGTATAGCATACAT IATACGAAGTTATGAGAACCGGATATAAAGATCACTGTGC				Exon 2	1 kb
VIB-UGENT IRC Transgenic mouse Core Facility, Belgium	Taok3	5'-GGGTAACTGTGGTGACTTTG TGG-3'	5'-GGAGGCTGAGGCGGAACCAA TGG-3	S'- CAGGACCTCTCAATAGAAGTTGCCTCATACCCTCTTAGGT CAAGGGTAACTGTGGTGACTATAACTTTCGTTATAGCAATACA CTTATACCAAAGTTATTTGTGGCAGAGGCAGGCAGCAGCATGCC CTAGAACGCAGTCAGGAGGGACCAGCAAACCCGGG-3'	GGTAGATCTTGAGCTCCATTGATAACTTCGTATAGCATACA FTATACGAAGTTATGTTCCGCCTCAGCCTCCGAAGGGCCG		(IVT) 20 ng/μl	10 ng/μl	Exon 7	700-800 bp
Molecular Biomechanics Lab, Indiana University, USA	Hspg2	5'-GACACAAGCATTTAGCTGCG-3'	5'- AAGGTGGCCTTGGGCAACCT-3'	5'- CTTAGCATTAGCCTGAGGTCGTGGGGAGCTGATGTCTGT CTTGCCTCCGTTTAGACACAAGCATTAGCTGTCGACATA ACTTCGTATAGCATACATATACGAAGTTATGCGGGGCTTT GCGTGTTCTGAGCAGTGGGATGATTAGCTGCGGGGCTTT TGCGTGTTCTGAGCAGTGGGA-3'	ACACAGCTTATTGGCAAGGTGGCCTTGGGCAAGTCGACA FAACTTCGTATAGCATACATTATACGAAGTTATCCTTGGAA	Protein = 100 ng/μl	synthetic 50 ng/μl	50 ng/μl	Exon 2 to 92	65 kb
	Toel	5'-GTGCCCTAAATTTGGAAAAA-3'	5'-GAAACAAGACCTTTGTGGAA-3'	5'- TTATCTAGTCCACTITTCTCCTGATCAIGAIGCAAAAAAG AAATGATTTCCCAGTCCCACATAAAGGCACTAAAAGACA CAGAIGCCCTTTATAACTTCGTATAGCATACATTATAGGAA GTTATTTCCAAATTTAGGGACTGCCCATTTCAGAGCTTG [3']	GTTTCCCACTGGGGAAGCACCGCCTCATTATCTAACCCCG CCCTGCCCCTTCATAACTTCGTATAATGTATGCTATACGAA	EZ Protein = 6 μM (1,000 ng/μl)	synthetic = 209 ng/μl	435 ng/μl	Exon 2-5	1.1kb
	Tctn1	5'-CGCGTGCTCAGTACTCTGCG-3'	5'-TGCAACAGGGCACCGGGCAG-3'	GCCACCCTGACAGCCACTGTGACTGAGTCTTCAGGACTC GCATCGATCAGGCAGGAACCTCACAGGAGGCGAGAGG GAGAGCTTCCTCGCATAACTTCGTATAGCATACATTATACG AAGTTATAGAGTACTGAGCACGCGATAGACTGGCTCAGT TTAC-3' TTAC-3'	GGGGTCTCTCACTGAATGTAGAAATCATCCCCCTAGCCTA CCCCCGATCCCCGGTTGACAGATGGCTAGCATCCTCTTGT CTCCTCCCTGATAACTTCGTATAATGTATGCTATACGAAG	EZ Protein = 8 μM (1.333	Synthetic = 279 ng/μl	435 ng/μl	Exon2	1.2 kb
Mouse Biology Program (MBP), University of California, Davis, USA	1700069L16Rik	5'-GCTACAGCCGCTGCCCATGG-3'	5'-GTAAATGGCCAGCGCTCAGG-3'	CTGATGGACCACCAAGGCTCATACTGCATACATTCAACTC GGACTGCATCTGCTCCGACTCCCACTGGCACGATTCAG CTCACCACCTCCCATAGCTTCGTATGCATACATTATACGAA GTTATTGGGCAGCGGCTGTAGCCTGTGAGAGAGGGAACG CTGTTATTGGGCAGCGGCTGTAGCCTGTGAGAGAGGGAACG CTGACTGAGACGAACCG	GGTTTCTCTGTGTAGCCCTGGCTGTCCTGGAACTCACTCT GTAGACCAGGCTGGCCTGGAGAACTCAGAAATCCACCTG CCTCTGCCTCCTATAACTTCGTATAATGTATGCTATACGAA	ng/μl)	Symmetre 277 again	133 lig µi	Exon 2	1 kb
	Gene A	5'-GGTGTCTGGCTGCTCCAAAG-3'	5'-GTGGTTTTAGGGGACCTCTG-3'	TGCCAGAGTGCCCCTGAACCCAGGCGACCCCCGCCCGG CCAGCACTTTTGGTCTCTCTGTGGGAGACACTTTTAGGA GGCGCTAGCCGCTATAACTTCGTATAATGTATGCTATAAC AAGTTATTGGAGCAGCCAGAACACCGGGAACTTGAACCTG	GTTCTAGGAGCCAAACTCAACAGCTGTGCTCAACATGTG GGTCTCAATCCCTTTGCTGGGGGGTTGGGGGTTGTCAAAT GACCCTTCCACAGATAACTTCGTATAATGTATGCTATACGA	mRNA (IVT)=100 ng/μl	(IVT) 20 ng/μl	20 ng/μl	Entire gene	1.1 Mb
	Gene B	5'-CTTAGCCAACCCTCACTCGA-3'	5'-ATTAACAAGAGTCCCTGCAA-3'	5- AGGGAATATGATGACCCTATGAATTCTATGCCTTTTCTTAA, AAGTAAGTTATAAAGTGTCATACCTCTGAAGATGTGGCCTC CAAGCCTTCGGCGGCCGCATAACTTCGTATAGCATACATA	FGTATGTATATATATAAAACTAAATGATTAATGAGATTTC CTTTGGCGGCCGCATAACTTCGTATAGCATACATTATACGA		(IVT) 20 ng/μl	20 ng/μl	Exon 4	850 bp
	Arid1b	5'-CCCATTTGCACAGTCTCTAAAGG-3'		5'. TGGAGTCTGGTGAGAAGTGGCCACTCTGAGTAGACTCCA CCTAGTGTGTAAGAATCTTGGTCTTACCCATTTGCAAGTTAT CTAATGGAGTGGCACTGTGATAATGTAATG	CCCCAACCCCAGCCCATGTGAAGACAGGTCTGGCTT AGAATTCATAACTTCGTATAATGTATGCTATACGAAGTTAT GCAGAGGCTCTCAAACTCTGCCTTCCTAGGCACTCTGTT	Protein = 25 ng/μl	synthetic 20 ng/μl	3-20 ng/µl	Exon 7	3-400 bp
Children's Research Institute Mouse Genome Engineering Core, UT Southwestern, USA	E2f9	5'-TAGAGTGATTGGTTCTAGTA-3'	5'-CAGATTCAGATAAGACCGTG-3'	CCTGAAAGCCTTTACCTGATGTCTCATTAGTAGGAATTTA GGAATTTAGAAGTGATCAGTTACTTAGAGTGATTGGTTCT GAATTCATAACTTCGTATAATGTATGCTATACGAAGTTATA GGTATGGAAAGGGAAGGACTGAGCTCTAAGCACTATTATA CAGCCCTCCTTTCTCAAAATAGAATTTAAAAATAAAACCT-C	FGAGCTCATATAGTCACACTATCACAGATTCAGATAAGAC CCATGGATAACTTCGTATAATGTATGCTATACGAAGTTATC GTGAGGGAATTCAAAGCCTCACAAGTCCTGTTAATTTGA	Protein = 25-100 ng/ μ l	synthetic 20-50 ng/μl	2-3 ng/μl	Exon 5	1.3 kb
	Imp3	5-AAGAAGGCCGCGCCCCCATGG-3'	5'-GAGCAGGTTTAGGTCTCAACTGG-3'	GGCACCCTGGCTCTTTAATCCCATCGCCCAGTGCTCCAGG CCAACCTGGATGTGGACCAGAAGAAGGCCGCGCGCGC ACGTCATAACTTCGTATAATGTATGCTATACGAAGGTTATCC CATGGCAGGCTGTTTGTTGGGGGTGTGTCTCTTCCAGAC CTTGGGGGGGCGCCCTTGGAGGGGGTCAGGAGG	TIGGGTTATTGTTGCTGTTTTAAAAGCCTGTCAAGGAAG CTCTCACAGCTACATGTTCTTGGCAGAGCAGGTTTAGGT CTGAATCAGTTCATAACTTCGTATAATGTATGCTATACGAA GTTATCAACTGGCAGCCCGAGTTTCAAAAGCCAATAGTC	Protein = 5-25 ng/μl	synthetic 5-25 ng/μl	2-3 ng/μl	Exon 1	5-600 bp

Centre de Recherche duCentre Hospitalie de l'Universite de Montreal (CRCHUM), Canada	Icam1	5'-GGCTGCCACTCAGTATGAGTTGG-3'		y- ATANATGAATAAAAAAAAAAAAAAGGTGGTGTGTGG CAAAGATGGGCCAACCTGTCT GGTGGACCCCAGAGGTCCAACTGAATTCATAACTTCGTAT ATGACAAAGCCACTGCCATGT AAAGTATCCTATACGAAGTTATCATACTGAGTGGCAGCCTC AGGTATCCATACGAAGCTATCTTTTGTTCTGCGGCCCTG AGACAGGGTTTCCCAGTGTAC GA-3' 3' 3' CAAAGATGGCCAACCAACCACTCTTTTTGTTCTGCGGCCCTG AGACAGGGTTTCCCAGTGTAC GA-3' 3' CAAAGATGAGCCAACCAACTACTTCTTTTGTTCTGCGGCCCTG AGACAGGGTTTCCCAGTGTAC GA-3' CAAAGATGAGCAACAACAACAACAACTACTTTTTGTTCTGCGGCCCTG AGACAGGGTTTCCCAGTGTAC	GCTAGCATAACTTCGTATA FAGTAAAATCTACGTTAGAT Protein = 50	ng/μl synthetic = 12 μM	200 ng/μl	Exon 4 to 7	3 kb
McGill Integrated Core for	Eftud2	5'-ACCTTTCCTACCACGTAGGC-3'	5'-GGAAIGTIGTCIGTAACGGG-3'	5'- AGAAAGGTCTGCGTTCTGGGGACACTATTCAGTAGCCTG GTCAGTCACTCAGCACATCACCACCTGCCTACGGAATT TGTTTTCTTTCTTTATCCCTCCC CATAACTTCGTATAGCATACATTATACGAAGTTATTGGTAG GCAATGACTTAGGAAGTTATGCGAAGTTATCGCACCACTACATTATACGAAGTTATCGCAAGTAAGACCTGCACGCAC	CGATATCATAACTTCGTATA TTACAGACAACATTCCCA mRNA (IVT) =	50 ng/μl (IVT) 25 ng/μl	50 ng/μl	Exon 2	2-300 bp
Animl Modeling (MICAM)	2911112	5'-CCTTACCTTTCCTACCACGT-3'	5'-GAATGTTGTCTGTAACGGGA-3'	AGAAAGGTCTGCGTTCTGGGGACACTATTCAGTAGCCTG AAGAGAGACAAAAGATCTTG, GTCAGTCACTCAGCCCAGTCACCACCTGGCTACGGAATT TGTTTTCTTTTC	CGATATCATAACTTCGTATA TTACAGACAACATTCCCA mRNA (IVT) =	50 ng/μl (IVT) 25 ng/μl	50 ng/μl	Exon 2	2-300 bp
University of Nebraska Medical Center, Omaha, NE, USA	Muc16	5'-GAAACAGCATGAATATAGTC-3'	5'-GGGCTTCTTATTTATCAACC-3'	5'- GITTATIGTAAACATAGGAATACTCAGTGAGAAAIGTTTTA GACTTAGGAAGGGTGACAAG TCCTGAAACAGCATGAATATAATAACTTCGTATAGCATACA ATTCAACCAGGTGAGCTCATA TATACGAAGTTATGAGCCTCGTCTGGGATAAAATTTCTTC ATACGAAGTTAATTGATAATTGATAATTATTATAATTATTATTA	ACTTCGTATAGCATACATT Protein = 10 AGAAGCCCTATTTTTACATT	ng/μl synthetic = 10 ng/μl	10 ng/μl	Exon 2	7.1 kb
	Klk15	5'-TTAGCGTTTCAAGTTGATGC TGG-3'	5'-CTATCAGGGGCCCCAAAGAT GGG-3'	5'- AGGGCAAAGAAGTICCATACCATTCCCTGGAGTTA CAGAATAGAAAATCCTCACCA GCGTTTCAAGTGCTAGCATAACTTCGTATAATGTATGCTAT AGGGGCCCCAATAACTTCGTA ACGAAGTTATTGATCCTGGGAACTCTAATCAGCCCACA TTATGAATTCAGATGGGTCATT GCAAAAGCAAGCTAACTCATCT3' AGGGGCTACAGAGCGCAG-3'	TAGCATACATTATACGAAG Protein = 200	ng/μl synthetic 80 ng/μl	3 μΜ	Exon 2	6-700 bp
Czech Center for Phenogenomics, Czech Republic	Pknox2 (blast only)	5'-CAAAACAACACGTAGTAACC AGG-3'	5'-GCCCACCTTCCCACTTGAGA TGG-3'	5- AAAACAGAATICTCTGITITCTAAGTGTGITTGGCATCCA GTTGGCTGAACTGAGGGTTTG CTCCCTGGTTCAGATAACTTCGTATAATGTATGCTATACG AAGTTATACTACGTGTTTGTTTTGT	ATAGCATACATTATACGAAG Protein = 200	ng/μl synthetic 80 or 120 ng/μl	3 μΜ	Exon 5	6-700 bp
	Dcaf12 (blast only)	5'-GAGGCCGAGGATGCAAGTTC TGG-3'		y: TGITTGTTCAGCACCAGGGTCTCCATATGTGGCTCATGGT GTCCCAGAACTCTAGATAACTTCGTATAATGTATGCTATAC GAAGTTATTTGCATCCTCGGCCTCAGCTTCCTGAATTACA GGATTGCAACCATCCCCCCCCCC	TATAGCATACATTATACGA Protein = 200 CCCTATTTCAACCTCGGAC	ng/μl synthetic 80 ng/μl	3 μΜ	Exon 4	4-600 bp
				SgRNA = Single Guide RNA, ssODN = single sranded Oligonucleotide					
		Note # Data fro	om Bishop, K.A., et al. CRISPR/Cas9-Mediate	Insertion of loxP Sites in the Mouse Dock7 Gene Provides an Effective Alternative to Use of Targeted	Embryonic Stem Cells. G3 (Bethesda), 2016. 6(7)	: p. 2051-61.			

1							ed and non targeted from 49 unique loci		
Laboratory	Gene Lcmt1	Lethality	Zygotes injected 131	Zygotes transferred 112	Live born pups 5	Correctly targeted	Incorrectly targeted in 5' Incorrectly targeted in 3	Mouse strain C57BL/6N	Site of injection
		Embryonic				· ·	Indel = 1		-
iken BioResources Center,	Trp53	None	245	231	25	0	Indels = 9 $LoxP = 2$ Deletions = 8	C57BL/6N	С
Japan	Trf	Postnatal	361	319	18	0	0 $LoxP = 2$ Deletion = 11	C57BL/6N	С
	Ppme1	Postnatal	109	97	1	0	0	C57BL/6N	C
							X 11 0 X D 1		
	Trmt6	Unknown	426	173	16	0	Indels = 8; $LoxP = 1$; Deletions = 5	C57BL/6J	C and N
	Slc7a14	None	282	135	7	0	Indels = 3; $LoxP = 1$ Indels = 2	C57BL/6J	C and N
	Tmem163	Unknown	129	34	3	0	Indels = 3 Indels = 2; $LoxP = 1$	C57BL/6J	C and N
Osaka University, Japan	Wtap	Embryonic	147	91	17	0	Indels = 1 Indels = 17; $LoxP = 2$	C57BL/6J	C and N
	Mettl3	Embryonic	143	84	1	0	Indels = 2 Indels = 2; $LoxP = 1$	C57BL/6J	C and N
	Mettl14	Unknown	269	145	16	0	Indels = 1 Indels = 13	C57BL/6J	C and N
	Klotho	Postnatal	144	93	6	1	Indels = 4 Indels = 5	C57BL/6J	C and N
Tokai University, Japan	Arhgef16	Unknown	221	208	35	0	Indels = 31; $LoxP = 3$ Indels = 4	C57BL/6J	C / C and N
	Cd226	None	237	202	48	1	LoxP = 4; $LoxP$ in Trans = 2; Deletion = 16	C57BL/6J	N
	IgfI	Postnatal	444	383	83	0	LoxP = 8; 7/38 genotyped mice = deletion	C57BL/6J	N
							LoxP = 9; Knock-in but trans or mosaic manner = 6;		
	Pik3cb	Embryonic	459	408	159	0	25/77 genotyped mice: deletion	C57BL/6J	N
							25/77 genotyped finee, defendir		
							25 mice: single LoxP; 3 mice=both Knock-in but trans o		
	Gcn1 1	Unknown	451	395	84	0	mosaic manner; 20 mice= deletion	C57BL/6J	N
University of Tsukuba,									
Japan									
							20		
	St3gal5	None	429	368	89	0	20 mice: single LoxP; 1 mouse: cis knock-in but one Lox sequence is mutated and 1 mouse: both Knock-in but trar		N
	sisguis	None	429	308	89	U	or mosaic manner; 14 mice : deletion	S C3/BL/03	IN.
							or mosaic manner, 11 mice : deterior		
							9 mice: single LoxP; 2 mouse: cis knock-in but one LoxI	,	
	Gata3	Postnatal	441	409	77	0	sequence is mutated; 3 mouse: both Knock-in but trans of		N
							mosaic manner; 22 mice= deletion		
	Irf3	None	1294	651	75	0	LoxP = 2 $LoxP = 2$	C57BL/6N	C and N
TIGM, USA	Phactr1	Unknown	331	187	42	1	LoxP = 3 $LoxP = 5$; Deletions = 6	C57BL/6N	C and N
	Actrt2	Unknown	1013	562	70	3	LoxP = 7 $LoxP = 4$	C57BL/6N	C and N
	ApoA1	None	645	456	68	2	LoxP = 9 $LoxP = 4$	C57BL/6N	C and N
							8 indels with insertion of		
University of Adelaide,	Depdc5	Embryonic	154	128	11	1	single LoxP, 2 mice with 0	C57BL/6J	С
Australia	•						large deletions		
University of Rochester,	Akap12	None	1344	797	109	1	Indels = 9; $LoxP = 3$ $LoxP = 1$	C57BL/6J and (SJL/J	С
USA			1	1	- 47	•		xC57BL/6J)F1	, e

	Cav3 (design 1) Cav3 (design 2)	None	272	257	15	0	Indels = 8	0	C57BL/6J	N
	Cav3 (design 3)	None	107	104	5	0	Indels = 3; $LoxP = 1$	0	C57BL/6J	N
	Cav3 (design 4)	None	26	25	4	0	Indel		C57BL/6J	N
	Cav3 (design 5)	None	136 105	128 99	17 17	0	Indels = 10 ; $LoxP = 3$	0	C57BL/6J	N
Maine Medical Center, USA	Icam1 Dock7 (design 1) #	None None	305	305	47	0	Indels = 19; $LoxP = 8$	Indels = 1 Indels = 19; $LoxP = 10$	C57BL/6J C57BL/6J	N C and N
Maille Medical Celiter, USA	Dock7 (design 1) # Dock7 (design 2) #	None	191	174	20	1	Indels = 10; $LoxP = 6$	Indels = 10; $LoxP = 10$	C57BL/6J	C and iv
	Gpr180	None	203	194	11	0	LoxP = 7	LoxP = 5	C57BL/6J	N
	Noct	None	284	267	11	0	LoxP = 1; 1 mouse has 5' and 3' $LoxP$ in Trans	LoxP = 2	C57BL/6J	N
	FoxD1	Postnatal	247	239	23	0	LoxP = 3; 1 mouse has 5' and 3' $LoxP$ in Trans	LoxP = 1	C57BL/6J	N
MD Anderson, USA	Atg7	Postnatal	59	9	5	0	Indels = 2	0	C57BL/6N	N
	NIrd1	None	52	36	13	0	LoxF	P = 2	C57BL/6J	N
W : 2 04 1 .	Bp1-Bp2	Unknown	243	222	44	0	LoxF	P = 9	C57BL/6J	N
University of Manchester, UK	Lrcc8a	Postnatal	251	238	27	1	LoxF	P = 1	C57BL/6J	N
	Usp7	Embryonic	169	140	21	0	LoxF		C57BL/6J	N
	Bin1	Postnatal	112	92	11	0	LoxF	P = 5	C57BL/6J	N
	Meox I	Embryonic	495	154	9	0	Indel	le = 3	C57BL/6J	С
University of Minnesota,		-								
USA	Meox2	None	465	125	14	0	LoxP = 1	Indels =1; Deletion = 1	C57BL/6J	C
	Vezf1	Embryonic	422	155	11	0	LoxP = 1	Indels = 2	C57BL/6J	С
VIB-UGENT IRC Transgenic mouse Core Facility, Belgium	Taok3	Unknown	422	360	22	0	5' indels = 3; <i>LoxP</i> = 3	Indels = 8; LoxP = 1; Deletion = 6	C57BL/6J	N
Indiana University, USA	Hspg2	Embryonic	899	524	59	0	Indels = 15	5; LoxP = 2	(C57BL/6xC3H/He)F1	N
Mouse Biology Program	Gene A	None	180	115	20	0	C	0	C57BL/6N	C and N
(MBP), University of California, Davis	Gene B	None	100	79	17	0	LoxP = 1	LoxP = 1	C57BL/6N	C and N
Children's Research Institute	Arid1d	Postnatal	167	167	2	1	Indels = 1; $LoxP = 1$	LoxP = 1	C57BL/6J	C and N
Mouse Genome Engineering Core, UT Southwestern,	E219	Unknown	593	593	37	0	LoxP = 1	LoxP = 2; Deletion = 2	C57BL/6J	C and N
USA	Imp3	Unknown	372	372	94	0	Deletio	on = 1	C57BL/6J	C and N
Centre de Recherche du Centre Hospitalié de l'Université de Montréal (CRCHUM), Canada	Icam1	None	150	123	5	0	LoxP = 1	0	C57BL/6J	N
McGill Integrated Core for Animal Modeling (MICAM)	Eftud2	Postnatal	530	389	23	0	Indels = 11; LoxP	P=1; Deletion = 1	C57BL/6N and (FVBxCD1)F1	N
	И. И.	None	68	59	10	1	Indels = 1; $LoxP = 2$;	Indels = 1; $LoxP$ = 3;	C57BL/6J	N
University of Nebraska Medical Center, Omaha, NE, USA)	Muc16	110110					Deletion = 2	Deletion = 1		

Note: # from Bishop, K.A., et al. CRISPR/Cas9-Mediated Insertion of loxP Sites in the Mouse Dock7 Gene Provides an Effective Alternative to Use of Targeted Embryonic Stem Cells. G3 (Bethesda), 2016. 6(7): p. 2051-61.

Supplementary Table 3: Generalized Regression analysis to identify the factors predicting the success of the 2sgRNA-2ssODN methods

	Estimate	Standard Error	% of variance explained	t.value	P.value
Intercept	0.01	0.17	3	-0.06	0.95
Efficiency of 5' SgRNA	0.27	0.23	5.4	1.18	0.24
Efficiency of 3' SgRNA	-0.24	0.22	5.2	-1.08	0.28
5' <i>LoxP</i> insertion	-0.3	0.96	9.4	-0.29	0.77
3' <i>LoxP</i> insertion	-0.06	0.38	15.1	0.19	0.85
5' LoxP X 3' LoxP insertions	5.03	2.45	60.4	2.05	0.051
Cas9 mRNA concentration	0.006	0.002	<0.1	2.8	0.009***
Cas9 protein concentration	0.001	0.003	<0.1	0.32	0.75
SgRNA concentration	-0.0025	0.002	< 0.1	-0.98	0.33
ssODN concentration	0.0006	0.001	<0.1	0.34	0.73
Distance between alleles	<-0.0001	0.003	<0.1	-1.5	0.14

SgRNA = Single Guide RNA, ssODN = single sranded Oligonucleotide *** means p.value < 0.01 * means p.value < 0.05

Dcaf12

_	<u> </u>			_	_	Zygotes	<u> </u>	<u> </u>	
Laboratory	Gene (Name)	Lethality	Zygotes Electroporated	Zygotes transferred	Live born pups genotyped	electroporated and genotyped at blastocyst stage	Correctly targeted	Incorrectly targeted	Mouse strain
McGill, Canada	Eftud2	Postnatal	67	60	0	NA	0	None	C57BL/6N
Mouse Biology Program (MBP),	Toe1	Unknown	50	40	11	NA	0	LoxP = 1 in 5'and 3'; deletions=3	C57BL/6N
University of	Tctn1	Unknown	60	40	3	NA	0	2 deletions	C57BL/6N
California, Davis, USA	1700069L16Rik	Unknown	60	40	11	NA	0	LoxP = 1 in 5'and 3'; deletions= 8	C57BL/6N
	Klk15	Unknown	93	93	9	NA	0	Indels = 6	C57BL/6N
Czech center for phenogenomics,	Pknox2	Unknown	NA	NA	NA	100	0	Indels=36; deletions=39	C57BL/6N
Czech republic	Dcaf12	Unknown	NA	NA	NA	60	0	Indels = 16;	C57BL/6N

NA

34

60

C57BL/6N

deletions =3

NA

273

NA

330

Unknown

Supplemen	tary Table 5: Over	all efficieny of the 2sg	gRNA-2ssODN metho	od of generating the	e cKO alleles
Delievery method	Number of loci	Zygotes processed	Zygotes transferred (% zygotes processed)	Live born pups analyzed (% zygotes transferred)	Correctly targeted (% live born)
Microinjection	49	17,557	12,491	1,684	15
Electroporation	5	330	273	34	0
Total	54	17,887	12,764 (71.4%)	1718 (13.4%)	15 (0.87%)