Next-generation mammalian genetics toward organism-level systems biology

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Organism-level systems biology in mammals aims to identify, analyze, control, and design molecular and cellular networks executing various biological functions in mammals. In particular, system-level identification and analysis of molecular and cellular networks can be accelerated by next-generation mammalian genetics. Mammalian genetics without crossing, where all production and phenotyping studies of genome-edited animals are completed within a single generation drastically reduce the time, space, and effort of conducting the systems research. Next-generation mammalian genetics is based on recent technological advancements in genome editing and developmental engineering. The process begins with introduction of double-strand breaks into genomic DNA by using site-specific endonucleases, which results in highly efficient genome editing in mammalian zygotes or embryonic stem cells. By using nuclease-mediated genome editing in zygotes, or ~100% embryonic stem cell-derived mouse technology, whole-body knock-out and knock-in mice can be produced within a single generation. These emerging technologies allow us to produce multiple knock-out or knock-in strains in high-throughput manner. In this review, we discuss the basic concepts and related technologies as well as current challenges and future opportunities for next-generation mammalian genetics in organism-level systems biology.

INTRODUCTION

Systems Biology is a natural extension of molecular and cellular biology,1–3 which consists of multi-stage processes beginning with a (1) comprehensive identification and (2) quantitative analysis of individual system components and their networked interaction, which leads to the ability to (3) control existing systems toward the desired state and (4) design new systems based on an understanding of the underlying structural and dynamical principles. After identification of key genes by classical forward and reverse genetics, systems biology in mammals has been further accelerated by a series of genome projects, especially at the molecular-to-cellular levels, where in vitro cell culture systems allow system-level identification, analysis, control, and design of molecular networks. On the other hand, organism-level systems biology in mammals still remains an important challenge in biology.4

In order to identify and analyze molecular networks and/or cellular circuits in organisms, gene knock-out (KO) or knock-in (KI) are powerful technologies often used in mammalian reverse genetics. However, this classical genetics requires several generations of crosses to produce mutant animals of sufficient quality and quantity for phenotype analysis. The time consuming conventional methods for producing KO or KI mice usually involve targeting-vector construction (2 weeks to a few months; depending on the complexity of constructs), the introduction of target mutations into embryonic stem cells (ESCs) by homologous recombination (a few weeks), and the injection of the mutant ESCs into wild-type blastocysts to produce chimera mice (~3 weeks). If the mutant ESCs contribute to the germ-line of the newborn chimera mice, their next-generation offspring will possess a heterozygous mutation (~3 months). Further crossings of the offspring (several months to years; at least 3 months per generation) will produce mice with completely homozygous KO or KI mutations on an inbred genomic background, which is required for reliable phenotype analysis. Thus, conventional methods require substantial amounts of time, space, and effort to knock out or knock in even a single gene. Therefore, to comprehensively identify and quantitatively analyze molecular networks and/or cellular circuits in organisms in an efficient manner will require next-generation genetics, i.e., genetic alterations without crossing. In this review, we discuss the basic concepts and related technologies as well as current challenges and future opportunities for next-generation mammalian genetics in organism-level systems biology.

CONVENTIONAL MAMMALIAN GENETICS

Mammalian genetics (particularly in mice) has been widely exploited in order to investigate complex and dynamic biological processes executed by molecular networks and/or cellular circuits in organisms. Forward genetics (germline mutagenesis and gene-trap) and reverse genetics (targeted KO or KI) are available in mouse genetics as in other model organisms such as yeast, nematode and fly. Especially, developmental engineering based on the establishment of cultured ESCs was often used to generate KO and/or KI mice.5–7 Various genetic tools can be also introduced by transgenic (Tg) mice techniques.8

However, the production of genome-edited mice has been generally low-throughput, and needed huge time and effort in the
**a Conventional mammalian genetics**

| Original strain | Genome editing in zygote | F0 founder | Crossing | Animal for analysis |
|-----------------|--------------------------|------------|----------|---------------------|
| Hybrid strain   | Injection into pronuclei of one-cell zygote | Founder Tg | (Founder selection) (Backcross and expansion) | Established Tg (congenic) |
|                 |                          |            |          | 3 months ~ / generation |

| Original strain | Genome editing in ESC | F0 chimera | Crossing | Animal for analysis |
|-----------------|-----------------------|------------|----------|---------------------|
| 129 strain      | ESC injection into blastocyst | Heterozygous mutant | Inbred strain for backcross | Heterozygous mutant (congenic) |
|                 | Targeting (dependent on spontaneous HDR) | Heterozygous mutant (congenic) | Inbred strain for backcross | Homozygous mutant (congenic) |
|                 |                        |            |          | 1 month ~ 3 months ~ / generation |

**b Next-generation mammalian genetics**

| Original strain | Genome editing in zygote / ESC | F0 founder | Animal for analysis |
|-----------------|--------------------------------|------------|---------------------|
| Triple-CRISPR   | CRISPR with sgRNAs             | Single gene | Animal for analysis |
|                 | Inbred strain for analysis     | Injection into pronuclei of cytoplasm | Bi-allelic KO mouse (One-step generation) |
|                 |                                |            | 3 months ~          |

| Original strain | Genome editing in ESC | F0 chimera | Animal for analysis |
|-----------------|-----------------------|------------|---------------------|
| ES mouse        | ESC injection into 8-cell embryos | KI ES mouse | 1 month ~ 3 months ~ |
|                 | Site-specific nuclease | ESC of inbred strain (3i25 + LIF) | 1 month ~ 3 months ~ |

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conventional ways (Fig. 1a). For example, a Tg mouse strain is produced by pronuclear injection of a DNA fragment harboring a transgene, which is randomly integrated. Therefore, non-specific expressions of the transgene are usually observed in the resultant strain and the F0 founders must be selected and further expanded for use as the strains for the subsequent research. In case of gene targeting in ESCs, a chimera mouse (mouse having both ESC and host embryo-derived cells) is first produced by injection of the ESCs into blastocyst-stage embryos. If the injected ESCs by chance contribute to germ-line cells, the resultant F0 chimera can transfer the introduced mutation to the next F1 generation. Therefore, the homozygous mutants can be obtained, in principle, at least in the third (F3) generation, which takes 9 months after ESC injection. However, these procedures are not robust and it usually takes longer because of low targeting rates in ESCs, low germ-line transmission rate in chimera, or unexpected infertility of the created mutant strain. Furthermore, because a mixed genetic background can cause phenotypical alterations which make the experimental results difficult to interpret, the generated strains additionally need to be backcrossed to a ‘standard’ inbred strain such as C57BL/6 (hereafter denoted as B6) several times. This labor-intensive step is practically required in most cases because F1 hybrid strains or 129 strain-derived ESCs are commonly used in Tg zygote production or targeting in ESCs, respectively due to their higher viability or efficient germ-line transmission in F0 chimera.

Despite the limitations of conventional mammalian genetics, systematic, large-scale mouse genetics projects have been performed. For example, ethyl-nitrosourea mutagenesis in mice was exploited to screen mammalian circadian clock genes and for systematic gene function studies. The gene-trap strategy has more recently been applied to such forward-genetics approaches, and >100,000 of trapped ESC lines have been established and kept in international organizations (e.g., International gene trap consortium or IGTC, http://www.genetraps.org). Other systematic international efforts to collect, prepare and maintain mutant mice and ESCs have also been performed, such as the International Knockout Mouse Consortium/International Mouse Phenotype Consortium (http://www.mousephenotype.org). Multiple Cre Tg/KI strains have also been established by individual researchers, institutes and international consortiums. However, to carry out organism-level systems biology, these large-scale efforts should be scaled down to the single-laboratory scale or even to the single-researcher scale. To address this technological challenge, next-generation mammalian genetics without crossing is proposed here to allow completion of KO or KI mouse production and phenotyping analysis within the F0 generation (Fig. 1b). This can be realized by the application of highly efficient genome editing by site-specific nucleases for one-step generation of whole-body genome-edited inbred animals within a single generation. Recently, there has been rapid progress in next-generation mammalian genetics, as introduced below, which will form an essential platform for organism-level systems biology.

**CURRENT TECHNOLOGIES FOR EFFICIENT GENOME EDITING BY SITE-SPECIFIC ENDONUCLEASES**

Double-strand breaks (DSBs) lead to several DNA repair pathways, such as (1) homology-directed repair (HDR) where a homologous DNA sequence is used for recombination or annealing, (2) non-homologous end joining (NHEJ), where the broken ends are directly reconnected with a frequent insertion or deletion of a random number of bases (denoted as “indels”), (3) microhomology-mediated end joining (MMEJ), where a small microhomology fragment is used for the end-joining. Induced DSB and the following repair processes enable highly efficient genome editing (both KO or KI) at the broken locus. Although the genome editing induced by DSBs was previously investigated by introduction of I-SceI (a mitochondrial endonuclease from *S. cerevisiae*) into cultured mammalian cells, which stimulated extrachromosomal homologous recombination, DSB-induced genome editing has been practically used in recent years. We review such efficient genome editing methods using site-specific nucleases, which can accelerate the production of KO and KI mice via relatively simple steps and thus help realize next-generation mammalian genetics (Fig. 2).

Site-specific endonucleases used in modern genetics

Three major classes of site-specific endonucleases have been used for genome-editing, transcription activator-like effector nuclease (TALEN) and clustered regularly interspaced short palindromic repeats (CRISPR)-associated Protein9 (Cas9). ZFN and TALEN are categorized into customizable endonucleases because they are composed of a customizable sequence-specific DNA-binding domain fused to a nonspecific DNA catalytic domain of FokI endonuclease. On the other hand, Cas9 is a RNA-guided endonuclease and recruited to specific DNA sequences by a short RNA guide molecule that recognizes target DNA via base-pairing.

The DNA-binding domain of ZFN contains 3–6 arrays of Cys2-His2 zinc finger motifs. The individual zinc finger recognizes 3 bp in the major groove of DNA. Selected zinc-finger modules that recognize nearly all of the 64 possible nucleotide triplets have been developed. Therefore, an assembled custom array of six zinc fingers can be constructed to recognize a unique 18 bp sequence. This length of target sequence covers a possible 68 billion unique DNA sequences.

TALEN contains another type of customizable DNA-binding domain, which is composed of a tandem 33–35-amino acid repeat (TALE repeat) derived from the plant pathogenic bacteria genus *Xanthomonas*. The individual TALE repeat recognizes a single base pair via two hypervariable amino acid residues inside the repeat (repeat-variable di-residues or RVDs). The four most common RVDs (HD, NG, NI, and NN) are known to recognize each of the four nucleotides (C, T, A, and G). Therefore, the tandem TALE-repeat is usually constructed with approximately 18 TALE repeats of different base pair-binding specificities, under consideration of its limitation that TALE-binding sites should start with a T base. The TALE repeat domain generally gives similar DNA-binding specificity and more flexibly when compared with ZFNs.
DSB induction by site-specific endonucleases

**ZFN**
- FokI
- Genome DNA

**TALEN**
- Indel insertion

**CRISPR/Cas9**
- Large fragment deletion
- Indels

Null mutation (with NHEJ)
- Indel insertion

Fragment insertion

**Large fragment insertion (with HDR)**
- Targeting vector (with long homology arm)
- Inserted fragment

**Large fragment insertion (with NHEJ)**
- Donor plasmid (digested in vivo)
- Donor fragment (PCR products, double-strand ODN)
- Indels
- Inserted fragment

**Small fragment insertion (with HDR)**
- ssODN
- Inserted sequence (Intended mutation, protein tag, LoxP etc.)

**Large fragment insertion (with MMEJ)**
- Donor plasmid or fragment (with microhomology sequences)
- Inserted fragment

**Introduction of null mutation**

DSBs induced by site-specific endonucleases activate an internal DSB-repair pathway, which is exploited for efficient genome editing. Among them, NHEJ-dependent indel insertion is the simplest and the most effective method for gene KO. In the CRISPR/Cas9 system, the DSB is repaired by the NHEJ pathway, which can cause a nonsense or null mutation in the targeted ORF. Long deletions can also be repaired by NHEJ pathway, small deletion or insertion of nucleotides (indels) occurred at the joint site, which cause a nonsense or frame-shift mutations in multiple DSBs. This means that two ZFN or TALEN molecules must bind on both right and left sides of the target site with an appropriate orientation and spacing. Therefore, the dimer recognizes 2-fold longer sequence at the target site than single ZFN or TALEN molecules. This molecular property gives higher specificity and reduced off-target effect.

Unlike the former molecules, Cas9 is an RNA-guided DNA endonuclease derived from the type II bacterial adaptive immune system CRISPR, and is recruited to specific target sequences by two short RNA molecules, which anneals with the target sequence, and the trans-activating crRNA (tracrRNA) which is partially complementary to the crRNA and anneals to the crRNA. This two-component RNA system was further simplified to synthetic single-guide RNA (sgRNA) consisting of a fusion of crRNA and tracrRNA. The target sequence in the CRISPR/Cas9 system can be readily designed by simply redesigning a part (around 20 bp) of the crRNA or sgRNA. This simplicity is in contrast to the much more burdensome procedures in ZFN and TALEN vector construction. This simplicity endows the CRISPR/Cas9 system with a significant advantage for use as a site-specific endonuclease for various genome editing purposes, including multiple gene KO, or even genome-wide gene perturbations.

Many studies have tried to increase the flexibility and decrease any off-target effect of the CRISPR/Cas9 system for practical use. The DNA cleavage activity of Cas9 molecules is dependent on the presence of a short (around 2–6 nucleotides) protospacer adjacent motif (PAM), which is located beside the complementary sequence of crRNA/sgRNA-targeted region. PAM sequence varies according to the CRISPR-based systems and organisms, and restricts the flexibility of the target sequence. However, the sequence dependency of the Cas9 molecule can be artificially modified and such PAM engineering can expand the target range of the system. Other modification of Cas9 molecules contribute to off-target suppression. Cas9 induces DSBs at approximately three bases upstream of the PAM by two endonuclease domains, a RuvC-like endonuclease domain (RuvC domain) and a HNH-like endonuclease domain (HNH domain), which are located at the amino terminus and the mid-region of the Cas9, respectively. The RuvC domain cleaves the non-complementary strand while the HNH domain cleaves the crRNA-complementary strand. Inactivation of these endonuclease domains via point mutations can convert Cas9 endonuclease into a DNA “nickase” that creates a single-stranded break, which reduces off-target activity by 50-fold to 1500-fold in cell lines and zygotes without sacrificing on-target cleavage efficiency. Others have tried FokI-dCas9 fusion protein as a dimer to improve target specificity by their recognition of distinct sites. Use of truncated sgRNA can also suppress undesired off-target activity by >5000-fold without sacrificing on-target genome editing efficiency, possibly by decreasing the sgRNA-DNA interface.

Dimerization of the FokI endonuclease catalytic domain is essential for cleavage of DNA by ZFN and TALEN. This means that two ZFN or TALEN molecules must bind on both right and left sides of the target site with an appropriate orientation and spacing. Therefore, the dimer recognizes 2-fold longer sequence at the target site than single ZFN or TALEN molecules. This molecular property gives higher specificity and reduced off-target effect.
In a recent study, the improvement in KO efficiency with the use of three sgRNAs (triple-CRISPR) was examined in depth. Based on simulation, the average KO efficiency expected with single sgRNA was around 60% and triple-CRISPR would increase the rate to over 80%. The actual rate reached to over 95% due to long deletions between CRISPR targeted sites. A set of triple sgRNAs which cover ~80% of all genes in the mouse genome has been created as an open database (http://crispr.riken.jp/). Potential off-target effects can be also excluded by using the second set of triple sgRNAs that covers ~70% of all mouse genes.62

Introduction of DNA fragments
Targeted insertion or KI of a DNA fragment with mutated sequence, short functional sequence (restriction enzyme site, recombinase recognition site, or protein tag etc.), or functional expression cassette can be also facilitated via HDR, NHEJ and MMEJ by co-transfer of linear or circular donor vector, PCR fragment or single-stranded oligo DNA nucleotide (ssODN) together with the site-specific endonucleases (Fig. 2).

Homologous recombination (HR)-dependent targeting is mediated by a form of HDR. This pathway has been widely used for a large fragment insertion or KI both in cultured cells and zygotes by using a donor targeting vector with long homology arms.43, 44, 63–69 The targeting rate is relatively low but efficiently enriched by antibiotic drug selection in the culture. A shorter functional sequence or small mutation can be more simply introduced by using ssODN.51, 55, 59, 66, 70–75

NHEJ-mediated fragment insertion/KI is easier and more efficient than the HR pathway, because the NHEJ-repair reaction is thought to predominate over the HR reaction for DSB repair.76, 77 In the NHEJ-mediated insertion, both the donor plasmid and the target genome loci are digested simultaneously. And then, the digested donor plasmid is integrated into the digested genome loci. A PCR fragment or double-stranded ODN can be also applied as an integrated fragment without digestion. This pathway works not only in cultured mammalian cells (including ESCs) but also in zebrafish, and does not necessarily require antibiotic selection.56, 78–81

In addition, there is no need to prepare a targeting vector with long homology arms, which is generally a time-consuming process. On the other hand, it is of note that the direction of the inserted fragment is not controllable, and indels are usually introduced at the junction site. Therefore, the method is inappropriate for some KI purposes, such as in-frame KI of an exogenous ORF into an endogenous gene.

MMEJ-mediated editing provides more simplified KI strategy with precise direction and junction sequence. Instead of the conventionally used long homology arms for HR-mediated KI, this pathway uses only extremely short microhomology sequences (10–40 bp) for the precise fragment insertion. MMEJ-mediated KI also works in mammalian cells, and the inserted fragment can be supplied as an in vivo digested plasmid or a PCR fragment.82, 83

Fig. 3 One-step generation of genome-edited mice. a An example of triple-CRISPR KO littermates (before weaning) in B6 strain. Tyr gene coding tyrosinase (an enzyme involving black coat color) was knocked out by the triple-CRISPR method.62 All littermates had white coat color, indicating ~100% bi-allelic KO rate of the targeted gene. b An example of B6 ES mouse littermates (before weaning) by 3i + LIF culture and 8-cell injection.141 All littermates had black coat color, indicating efficient generation of ~100% ESC-derived mouse. c An ES mouse embryo (E13.5) derived from an H2B-EGFP KI ESC clone.170 Only the embryo (but not the extraembryonic tissues) expresses EGFP, suggesting the unique contribution of ESC-derived cells. Entire section of the ES mouse is shown in the right panel. All animal experiments here were approved by the Institutional Animal Care and Use Committee of RIKEN Kobe Branch, and all of the animal care was in accordance with the Institutional Guidelines.
Therefore, the editing pathway potentially overcomes problems in HR-mediated or NHEJ-mediated KI.

CURRENT TECHNOLOGIES FOR ONE-STEP PRODUCTION OF GENOME-EDITED MICE

Direct genome editing in one-cell zygotes
The compelling advantages of the site-specific endonucleases in efficient genome-editing has been examined in recent years. In particular, zygotic genome editing enables one-step production of genome-edited animals, skipping the in vitro targeting step in ESCs. Introduction of components into one-cell zygotes are relatively simple and easy, particularly for the CRISPR/Cas9 system, which just requires cytoplasmic microinjection or electroporation.

KO animals can be generated in a one-step manner by exploiting endonuclease-mediated DSBs followed by NHEJ with indel insertion in zygotes. In an earlier study, ZFN was tested in rat zygotes, where up to 75% of live-born F0 founders were harboring mutations. TALEN has been similarly tested, while CRISPR/Cas9 was mainly used in the most recent studies, since 2013. Others reported F0 phenotyping of CRISPR/Cas9 KO animals, suggesting the potential of this method for use in next-generation genetics schemes. Several modifications of the CRISPR/Cas9 system have been also introduced to improve the efficiency and specificity of targeted mutations in a genome. However, two problems have remained: (1) first-generation mice often contain a mosaic of wild-type and KO cells, and (2) the rate of whole-body bi-allelic mutant mice generated is relatively low (usually ~60–80% at best). Therefore, the highly efficient (>90%) production of whole-body bi-allelic KO in a single generation remained a fundamental challenge for next-generation mammalian genetics. To realize this vision of next-generation mammalian genetics, the triple-CRISPR method significantly improved bi-allelic modification efficiency and further elicited almost perfect (~100%) whole-body bi-allelic KO mice. KO animals can be generated in a one-step manner, at the rate of approximately 20% in a triple KO experiment. The other modern genome-editing methods discussed above may potentially be applicable as well.

ESC-derived mouse (ES mouse)
Although one-step generation of KI mice would be ideal for next-generation mammalian genetics, a number of issue have yet to be overcome, specifically inefficient editing and live-birth rate (particularly with a large fragment KI) and undesirable indel insertion and mosaicism in resultant animals. Alternatively, two-step generation of KI mice by almost completely ESC-derived mouse ("ES mouse") is an attractive alternative at present. The advantages of using ESCs are in its selectivity of sex, easier storage and flexibility for more complex genome editing (e.g. multi-gene KO and KI) in in vitro culture. Furthermore, genome editing in ESCs is becoming easier by the site-specific nuclease method, which just requires cytoplasmic microinjection or electroporation.

Validity of ES mouse production and phenotyping analysis within a single generation was first proposed and tested by using the tetraploid complementation method. However, several possible drawbacks of the method are known. First, substantial contamination of host cells was often observed in chimera mice produced by this method, which can cause developmental abnormalities. Second, the genetic heterozygosity of both tetraploid embryos and ESCs seems to be crucial for survival of the resultant ES mice, which means that the use of inbred ESCs does not seem possible and further backcrossing is required. In addition, preparing hundreds of tetraploid embryos every time does not seem practical for routine generation of many ES mice. For these reasons, few reports have used tetraploid complementation in a large-scale phenotyping assay of ES mice. Furthermore, it is striking that ES mice from inbred strains, such as B6 and BALB/C were efficiently generated at the live-birth rates of 11–29% (ES mice/embryos transferred), which was comparable to the 129 strain (9–15%) and hybrid ES mice (~6–40%) and significantly higher than in blastocyst injection in any case (all ~0%). Therefore, F0 phenotyping of inbred ES mice generated with 8-cell injection/aggregation was considered plausible in the study. Possible drawbacks would be the ES mouse production rate might depend on the ‘quality’ of cultured ESCs. A further synthesized dual-crRNA:tracrRNA into pronuclei. The use of Cas9 protein also reduces mosaicism when introduced with proper timing in early zygotes. Another study reported the generation of two KI newborns out of 123 injected embryos, where one was heterozygous KI while another was homozygous KI. ssODN-mediated KI, in which the cut sites of the targeted genome and the inserted fragment are ligated with 80-bp ssODNs homologous to the two cut ends (Fig. 2), was also shown to integrate up to ~200 kb of bacterial artificial chromosome (BAC) into the targeted locus. Finally, it was recently reported that HDR accuracy is dramatically increased by incorporating silent CRISPR/Cas9-blocking mutations in sgRNA or Cas9-targeted sequence (CORRECT method) although testing this in zygotes still remains to be done. Please note however, that the reproducibility of these recent studies should be further examined because some of the conclusions are based on small number of experiments. Further improvements in the targeting rate and specificity will be needed for one-step generation of KI mice.
optimized method to stably maintain ESCs in their naive pluripotent state was thus required.

Secretory regulatory factors and their downstream mechanisms for maintaining ESC's naive pluripotency have been well studied (recently reviewed in Huang et al., and others). Historically, the leukemia inhibitory factor–signal transducer and activator of transcription 3 (LIF/Stat3) pathway has been found to be indispensable for maintaining pluripotency and self-renewal ability of ESC. More recently, additional pathways essential for ESC maintenance or differentiation were uncovered. One is the Wnt–β-catenin pathway which supports ESC propagation and maintenance of the pluripotent naive state, and is antagonized by glycogen synthase kinase-3 (Gsk3)–Tcf3 (also known as Tcf71). The other is fibroblast growth factor 4–mitogen-activated protein kinase (Mek)–mitogen-activated protein kinase (Mapk or Erk) pathway which leads ESC differentiation and thus its inhibition suppresses ESC differentiation. Therefore, naive pluripotent ESCs can be stably maintained by shielding the cells from these differentiation triggers, and by addition of inhibitors (3i: SU5402 for FGF receptor, PD184352 for Mek, CHIR99021 for Gsk3, or 2i: PD0325901 for Mek and CHIR99021 for Gsk3) and in serum-free media containing serum) (Fig. 3b, c). In addition, ~100% ESC-derived ESC chimera. To overcome these problems, B6 ESCs were suggested to have problems of maintenance from even inbred mouse strains (including B6) or rat, and increased germ-line transmission of B6-derived ESC chimeras. The stable establishment and maintenance of B6 ESCs in ES mouse production is particularly critical for the next-generation mammalian genetics without crossing. B6 ESCs were suggested to have problems of maintenance, less efficient chimera formation and germ-line transmission, and genomic instability in standard culture conditions. To overcome these problems, B6 ESCs were established and maintained in serum-free 3i/2i + LIF medium, which demonstrates significantly higher success rate (67% vs. 3% in media containing serum) (Fig. 3b, c). In addition, ~100% ESC-derived mice from the B6-3i ESCs can be stably generated with eight-cell injection, even after many passages and traditional homologous-recombined targeting, at a production rate of 30–100% (ES mice/live-born mice). Therefore, 3i/2i + LIF culture and eight-cell injection/aggregation of ESCs enables the efficient one-step generation of ES mice, and subsequent F0 phenotyping can be performed once the genome-edited ES mice are created. Indeed, production and data acquisition of a novel ES mouse strain expressing a bright fluorescent protein was completed within a few months. CRI/Spr/Cas9-mediated knock-in in 2i + LIF-cultured ESCs followed by eight-cell injection have also been performed. These examples support the potential of ES mouse schemes. Note that ICR/CD-1 host embryos can be used for the ES mouse production and there is no need for maintaining a specific Tg colony for host embryos. Furthermore, experimental procedures (such as an operation for implantation) are similar to, or even less than conventional chimera mouse production.

To further improve the efficiency of ES mouse production, better culture methods and ESC quality control will be needed. Genomic instability, in particular, should be avoided during culture because chromosomal aneuploidy can cause embryonic death. Telomere extension seems important for maintaining normal karyotype of ESCs, and frequent activation of telomere maintenance factor 53cn4 restores and maintains the ESC’s potency in long-term culture. Aneuploidy detection in cell culture populations is also important for ESC’s quality control. This can be performed by not only a conventional karyotyping but also by a droplet digital PCR-based screening. Furthermore, additional chemical treatments can possibly ameliorate ESC culture conditions. So far, a variety of chemicals including ROCK inhibitor, PKC inhibitor, ERK/p38 inhibitor, HDAC inhibitors [e.g., trichostatin A, sodium butyrate or valproic acid or Vitamin C] may potentially contribute to improved potency of ESCs. Therefore, applications of these chemicals and routine quality control may help accelerate next-generation genetics based on ES mouse production.

**PRACTICE OF NEXT-GENERATION MAMMALIAN GENETICS**

As discussed above, high-throughput KO or Ki mouse production is pivotal for accelerating system-level identification, and analysis of molecular networks and cellular circuits in organisms. Given that various genetic tools, such as optogenetics and chemogenetic tools, are developing rapidly in recent years, high-throughput genome-edited mouse production is required for their in vivo implementation. Next-generation mammalian genetics potentially enables a single laboratory or a single researcher to generate, maintain and analyze multiple genome-edited strains rather than institutes or consortiums for production, deposit and distribution of various strains and ESCs. Because sgRNAs for targeted sites can be readily designed and prepared, the CRISPR/Cas9 system, in particular, makes such large-scale genetics feasible. Indeed, a recent study generated 31 novel CRISPR-KO mice lacking tests-expressing genes. Since whole-body bi-allelic KO rates were not sufficiently high for the next-generation scheme, the authors performed the crossing of selected F0 founders based on sequence and PCR screening data. In order to realize almost perfect (~100%) whole-body bi-allelic KO rate for next-generation mammalian genetics, we recently performed triple-CRISPR-based large-scale reverse genetics for sleep research. To identify genes involving neural electrophysiological activities during sleep or wake, we first developed an average neuron model in silico and found that genes involved in intracellular Ca2+ regulation (Ca2+ channels, Ca2+-dependent channels, Ca2+-pumps or Ca2+-dependent enzymes) are important for electrophysiological slow-wave-oscillation patterns during sleep. To further assess the roles of these genes in vivo, we next produced KO mice for 33 genes with the triple-CRISPR methods and eventually identified 8 genes important for regulating sleep duration.

ES mouse technology can deal with more complex genome editing, which would be difficult, if not impossible, with the conventional crossing-based genetics. For example, we also produced ~20 KO-rescue ES mice in order to perform system-level analysis of circadian clock-gene circuits in organisms. In this experiment, a 3i + LIF-cultured ESC clone derived from a double-KO mouse lacking two core clock genes (Cry1 and Cry2) was established. A rescue expression cassette of the wild-type or mutated Cry1 gene was then homologously knocked-in in the ESC clone (thus five alleles had been edited). Double-KO ES mice and KO-rescue ES mice were then generated and used for F0 phenotyping to measure in vivo 24 h rhythmicity. As explicitly indicated by these examples, next-generation mammalian genetics enables large-scale organism-level experiments within reasonable time, space and labor.

Next-generation genetics is also important for improving animal welfare and 3R principles, particularly contributing to “reduction” of animal use. In our triple-CRISPR experiments, the yields of bi-allelic KO mice lacking tyrosinase gene (judged by the white coat color) were 17% on average, and 36% in the best case, of the injected and transferred B6 zygotes. Therefore, between 60 (average) and 30 (best case) of host embryos would be enough for generating a sufficient number (around 10) of bi-allelic KO mice. The rate of bi-allelic tyrosinase KO mice among the F0 littersmates was 97.5% on average and 100% at best. Similarly, at least in our ES-mouse experiments of Cry1 rescue in the Cry1/Cry2 DKO background, the yield of ES mice available for phenotyping was 5.5% on average, and 35% in the best case, of the injected 8-cell embryos. Therefore, between 170 (average) and 30 (best case).
case) of host embryos would be enough for generating a sufficient number (around 10) of ES mice. The rate of ES mice among the F0 littersmates was 43% as average and 91% at the best. Only the littersmates of embryonic lethal, non-KO or non-ES mice were sacrificed and no further animals are needed. The number of animals used is thus much smaller than the conventional methods, in which a similar number of host embryos are used for injection, and only a part of the founders or chimera mice are used for further crossing. In the conventional case, dozens of littersmates are produced and sacrificed during crossing to select mice with an expected genotype. With conventional methods the number needed exponentially increases when a more complicated genetic background (e.g., double KO) is desired, while with next-generation genetics the number of used animals is not dependent on genetic complexity.

On the other hand, researchers need to take special care regarding some issues with the use of F0 animals for phenotype studies. In particular, researchers should carefully consider to what extent potential mosaicism (e.g., mutational variations in the triple-CRISPR method, or undetectable contamination of wild-type cells in the ES mouse method) would affect the final results of a scientific study. In our above experiments, the phenotypic variations of F0 mice were comparable with those in wild-type or suitable control animals, suggesting that mutational variations (triple-CRISPR) or undetectable contamination of wild-type cells (ES mouse) do not seem problematic at least in these cases. To further exclude the possibility of artifact phenotypes due to mutational variations or undetectable contamination of wild-type cells, we recommend that researchers independently generate a whole-body bi-allelic KO mice by using a second set of triple-CRISPR for the same gene, or to independently generate a whole-body bi-allelic KI mice using an independent clone of ES cells. Such stringent criteria (the production of independent KO or KI mice to confirm the observed phenotype) is sometimes difficult to fulfill with conventional mouse genetics because it takes a couple of years for another round of production. On the other hand, this step is feasible with next-generation genetics because it only takes a couple of months. In this sense, the quality of scientific studies using next-generation genetics should exceed that of scientific studies using conventional genetics.

Another point that researchers need to consider carefully is experimental design. This includes (1) preparation of appropriate control animals without any obvious defects in the focused phenotype (e.g., ES mice from wild-type ESCs or ESCs without genome editing for a clock gene rescue study9, and tyrosinase triple-CRISPR KO mice for a sleep study29), (2) evaluation of genetic composition of the F0 animals with strict criteria (e.g., detection of a genomic deletion as a proof of efficient triple-CRISPR method92, 163 or a highly sensitive detection of contaminated host embryo-derived cells164). In addition, it is also useful, if necessary, to evaluate undesirable mutations in the coding sequences by exon sequencing164 of triple-CRISPR KO mice. It is also useful to avoid cumulative mutations165 by using ESCs with minimal passage numbers for the production of ES mice. Moreover, these experimental procedures should be described according to a general guideline for animal experiments (e.g., the ARRIVE guidelines166). Further efforts to minimize problems from these issues should be expanded in future studies.

PERSPECTIVES

Next-generation mammalian genetics will facilitate system-level analysis of molecular and cellular circuits in organisms. To further improve the throughput for genome-edited animal production, additional developments of new technologies related to next-generation mammalian genetics will be required. For example, one-step production of whole-body bi-allelic KI mice by more efficient genome editing of zygotes still remains unachieved. For the two-step production of KI mice using ES mouse technology, the preparation of host embryos and surrogate mothers are still labor intensive and pose limitations on its throughput. New technologies which overcome these limitations will further accelerate next-generation mammalian genetics and also reduce the number of experimental animals used to obtain the same information.

Next-generation mammalian genetics together with efficient, quantitative and non-invasive phenotyping methods will provide an attractive platform for investigating the organism-level functions of the molecular and cellular circuits of interests. Genome-wide phenotype analysis has been performed in international KO mouse phenotyping efforts158, 167 to systematically survey the functions of molecular networks in organisms. Although most large-scale organism-level phenotyping projects are usually labor-intensive, the development of more facile alternatives is possible. Recently, whole-body clearing and imaging methods with single-cell resolution have been developed146 and started to provide comprehensive and quantitative experimental data at the cell-to-organism level, further facilitating organism-level systems biology. The development of non-invasive phenotyping will be also an attractive direction. For example, sleep phenotyping represents such a recent attempt where a non-invasive sleep phenotyping system was used instead of a conventional invasive EEG/EMG-based measurement system. In fact, these non-invasive methods have already enabled sleep phenotyping of dozens of triple-CRISPR KO mice.62, 163 Organism-level systems biology is thus coming to fruition with next-generation mammalian genetics, whole-body clearing and imaging with single-cell resolution, and non-invasive and quantitative phenotyping methods. Organism-level systems biology based on such new technologies will accelerate our understanding of complex and dynamic molecular and cellular circuits in the near future.

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COMPETING INTERESTS

The authors declare that they have no competing financial interests.

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