Embryo-Engineered Nonhuman Primate Models: Progress and Gap to Translational Medicine

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Animal models of human diseases are vital in better understanding the mechanism of pathogenesis and essential for evaluating and validating potential therapeutic interventions. As close relatives of humans, nonhuman primates (NHPs) play an increasingly indispensable role in advancing translational medicine research. In this review, we summarized the progress of NHP models generated by embryo engineering, analyzed their unique advantages in mimicking clinical patients, and discussed the remaining gap between basic research of NHP models to translational medicine.

1. Introduction

Animal models of human diseases are the most important foundation for revealing the pathogenesis of diseases and exploring more effective therapeutic strategies. The available approaches to generate animal models of human disease generally include spontaneous animal models [1, 2], induced animal models [3–7], and embryo engineering animal models. Animal models with spontaneous mutation are formed in nature without human intervention. Although they could reflect clinical and pathological characteristics of the disease, they have shortcomings of difficult sources and few types, which greatly limit their applications. There are various methods used to induce animal models, including physical [3], biological [4–6], chemical [7], and/or other pathogenic factors. Among them, chemical and physical methods can generate animal models in a short time but they are difficult to reveal the disease process. By contrast, disease animal models constructed by embryo engineering can not only explore the treatment of diseases but also monitor the dynamic disease progression from an early stage.

Due to the development of assisted reproductive technologies (ARTs), such as in vitro fertilization (IVF) [8], intracytoplasmic sperm injection (ICSI) [9], and embryo culture technology in vitro, mammalian embryo development has been gradually explored. The development of mammalian embryos includes preimplantation and postimplantation development [10, 11]. After the oocyte is fertilized in the ampulla of the fallopian tube, the fertilized egg, known as the zygote, undergoes rapid mitotic division, known as cleavage. A cluster of 16 identical blastomeres is called morula. The blastomeres continue to divide to form blastocyst with an inner cell mass and trophectoderm. After the zona pellucida disappears, the blastocyst implants into the uterus to develop into an individual [12, 13]. During the preimplantation stage, there is a time window for engineering embryos. Several methods can be used during the window, including the sperm carrier method, which modifies the sperm [14], somatic cell nuclear transfer (SCNT), which works at the stage of mature oocytes [15, 16], microinjection of the viral vector or gene editing system, which works at the multiple stages [17–20], or embryonic stem cell (ESC) method to get the chimera [21, 22] (Figure 1).

Animal models constructed by embryo engineering have played an important role in understanding gene functions and molecular mechanisms of diseases. However, many small animals are different from humans in terms of genetics and evolution. Small animal models of the disease have shortcomings in simulating disease processes and
pathological phenotypes, which cannot be ignored. Due to the high similarities between nonhuman primates (NHPs) and human beings, NHP models have been recognized as an indispensable bridge from basic research to clinical transformation. In this review, we systematically reviewed the research progress of generating NHP disease models using embryo engineering and discussed the advantages and disadvantages of the NHP models. Finally, solutions to NHP model construction problems and potential applications of NHP models in translational medicine are prospected.

2. Development of the NHP Models

Although there are numerous technical and ethical limitations, many NHP disease models were constructed after unremitting exploration, even produced offspring models with a uniform genetic background. The first transgenic NHP model was generated 20 years ago [17]. Seven years later, the first gene-edited NHP disease model was constructed [23]. In general, more than 20 types of different gene-edited NHP models have been successfully generated in the past 20 years (Table 1). Here, we reviewed the existing gene-edited NHP models and major achievements from five aspects: viral vector-mediated transgenic NHP models, precise knockout genetically modified NHP models, precise knockin genetically modified NHP models, precise point mutation NHP models, and somatic cell nuclear transfer NHP models, to gain experience and develop translational medicine.

2.1. Viral Vector-Mediated Transgenic NHP Models. This method uses viral vectors to integrate related genes into the target cells. Chan and his colleagues successfully produced the first genetically modified monkey in 2001 [17]. They injected the pseudotyped replication-defective retroviral vector into the perivitelline space of 224 mature rhesus oocytes. Oocytes were fertilized by intracytoplasmic sperm injection (ICSI) and 40 embryos were transferred to 20 surrogates. One out of three newborns were detected to contain the GFP gene integration. Although the transgenic mice accelerated the advancement of biomedical sciences to a great extent, many differences between humans and rodents cannot be ignored. The advent of genetically modified monkeys that are more similar to humans was undoubtedly a significant breakthrough.

Based on the same technology, the first transgenic monkey model of Huntington’s disease (HD) was created in 2008 [23]. Yang et al. injected a high-concentration lentivirus carrying 84 CAG trinucleotide repeats of the human huntingtin (HTT) gene (HTT-84Q) into the perivitelline space of mature rhesus oocytes and successfully constructed five HD rhesus monkeys. Generally, healthy individuals contain up to 36 CAG repeats, whereas the disease occurs when the number of CAG repeats increase to more than 40. CAG repeat expansion translates into polyglutamine which further forms the polyQ aggregates, a master pathological feature of HD patients [24]. By studying those five HD monkey models, Yang et al. found that the levels of mutant HTT in the tissues seem to be associated with the severity of illness. This supported the theory that N-terminal mutant HTT fragments are pathogenic. The successful construction of gene-edited disease monkey models has opened up a new avenue for studying disease mechanisms and treatment strategies. It supplements the deficiencies of other animal models, especially mouse models.

Compared with Old World monkeys, such as rhesus monkeys and cynomolgus monkeys, the New World common marmosets (C. jacchus) have shorter reproductive cycles and smaller sizes. Therefore, using marmosets as animal models to study complex human diseases is faster and more feasible. In 2009, Sasaki et al. obtained different embryos through natural intercourse and IVF. By injecting lentiviral...
Table 1: NHPs models constructed by different methods.

| Type of models | Year | Species | Method | Gene | Stages | Samples | Implanted blastocysts | Pregnancy rate | Total neonatus | Models | Mutation rate | Contribution | Author/reference |
|----------------|------|---------|--------|------|--------|---------|------------------------|----------------|--------------|--------|--------------|-------------|-----------------|
| Retroviral vector | 2001 | Rhesus monkeys | GFP | Mature oocytes | 224 oocytes | 40 | 5/20 = 25% | 3 | 1 | 33% | The first transgenic monkey | Chan et al. [17] |
| Lentiviral vector | 2008 | Rhesus monkeys | Htt-84Q and GFP | Mature oocytes | 130 oocytes | 30 | 6/8 = 75% | 5 | 5 | 100% | The first monkey model of HD | Yang et al. [23] |
| Lentiviral vector | 2009 | Common marmosets | EGFP | Preimplantation embryos | 201 embryos | 91 | 7/50 = 14% | 5 | 5 | 100% | The first germline transmission | Sasaki et al. [18] |
| Lentiviral vector | 2010 | Rhesus monkeys | EGFP | Early cleavage-stage embryos | 70 embryos | 30 | 5/8 = 63% | 5 | 2 | 40% | The first transgenic monkey in China | Niu et al. [26] |
| Lentiviral vector | 2015 | Rhesus monkeys | α-Syn (A53T) | Mature oocytes | 133 oocytes | 75 | 11/25 = 44% | 7 | 6 | 86% | The first germline transmission | Niu et al. [27] |
| Lentiviral vector | 2016 | Cynomolgus monkeys | MECP2 | Mature oocytes | 53/105 | 9/18 = 50%; 7/36 = 19% | 8 | 2 | Total 8 | 80% | The first monkey model of autism-like disease | Liu et al. [28] |
| Lentiviral vector | 2016 | Cynomolgus monkeys | GFP | 24 hours after ICSI; 4 hours before ICSI | 18; 69 | 5; 17 | 2/3 = 67%; 5/17 = 29% | 0; 2 | 0; 2; 0% | 100% | Construction of cynomolgus monkeys expressing GFP | Seita et al. [25] |
| Lentiviral vector | 2019 | Rhesus monkeys | MECP2 | One-cell embryos | 21; 54 | 2/7 = 29%; 6/19 = 32% | 0; 1 | Total 4 | — | — | The first monkey model of RTT; the first monkey model of TALEN | Liu et al. [33] |
| TALEN | 2014 | Rhesus monkeys and cynomolgus monkeys | TALEN | MECP2 | One-cell embryos | 123 embryos | 123 | 14/41 = 34% | 7 | 6 | 86% | Research on RTT | Chen et al. [34] |
| TALEN | 2017 | Cynomolgus monkeys | TALEN | MECP2 | One-cell embryos | 52 | 3/9 = 33% | 3 | 1 | 33% | The first monkey model of human microcephaly | Ke et al. [19] |
| TALEN and ZFN | 2016 | Common marmosets | ZFN and TALEN | IL2RG | Pronuclear stage embryos | 250 embryos | 179 | 19/113 = 17% | 21 | 9 | 43% | The first monkey model of ZFN; the first monkey model of X-SCID | Sato et al. [35] |
| CRISPR/Cas9 | 2014 | Cynomolgus monkeys | CRISPR/Cas9 | Pparg and Rag1 | One-cell embryos | 186 zygotes | 83 | 10/29 = 34% | 5 | 2 | 40% | The first monkey model of CRISPR/Cas9 | Niu et al. [36, 37] |
| CRISPR/Cas9 | 2015 | Cynomolgus monkeys | CRISPR/Cas9 | P53 | Zygotes | 108 zygotes | 62 | 4/13 = 31% | 3 | 2 | 67% | The first live p53 biallelic mutant monkey | Wan et al. [20] |
| CRISPR/Cas9 | 2015 | Rhesus monkeys | CRISPR/Cas9 | Dystrophin | Zygotes | 488 embryos | 179 | 17/59 = 29% | 14 | 9 | 64% | The first monkey model of DMD | Chen et al. [38] |
| CRISPR/Cas9 | 2015 | Cynomolgus monkeys | CRISPR/Cas9 | Nrb1 (Dax1) | One-cell embryos | 186 zygotes | 83 | 10/29 = 34% | 5 | — | — | The first monkey model of AHC-HH | Kang et al. [39] |
| CRISPR/Cas9 | 2017 | Cynomolgus monkeys | CRISPR/Cas9 | SHANK3 | One-cell embryos | 116 embryos | 116 | 3/37 = 8% | 1 | 1 | 100% | The first monkey model of ASD | Zhao et al. [45] |
| CRISPR/Cas9 | 2018 | CRISPR/Cas9 | SIRT6 | Zygotes | 98 zygotes | 48 | 4/12 = 33% | 3 | 3 | 100% | | Zhang et al. [46] |
Table 1: Continued.

| Type of models | Year | Species | Method | Gene | Stages | Samples | Implantated blastocysts | Pregnancy rate<sup>a</sup> | Total neonatus | Models | Mutation rate<sup>b</sup> | Contribution | Authors/reference |
|----------------|------|---------|--------|------|--------|---------|------------------------|--------------------------|----------------|--------|------------------------|--------------|-------------------|
| Cynomolgus monkeys | 2019 | Cynomolgus monkeys | CRISPR/Cas9 | SHANK3 | Embryos | 178 embryos | 178 | 12/26 = 46% | 9 | 5 | 56% | The first monkey model of perinatal lethality syndrome | Zhou et al. [47] |
| Cynomolgus monkeys | 2019 | Cynomolgus monkeys | CRISPR/Cas9 | BMAL1 | Zygotes | 88 embryos | 88 | 10/31 = 32% | 8 | 5 | 63% | Monkey model of ASD and Phelan–McDermid syndrome | Qiu et al. [48] |
| Rhesus monkeys | 2019 | Cynomolgus monkeys | CRISPR/Cas9 | PKD1 | Embryos | 423 embryos | 86 | 29/86 = 34% | 5 | 4 | — | The first monkey model of circadian and psychiatric disorders | Tsukiyama et al. [49] |
| Cynomolgus monkeys | 2019 | Cynomolgus monkeys | CRISPR/Cas9 | PINK1 | One-cell embryos | 158 embryos | 11 | 11/28 = 39% | 8 | 5 | 73% | Monkey model of PD | Yang et al. [50] |
| Cynomolgus monkeys | 2019 | Cynomolgus monkeys | CRISPR/Cas9 | HBB | Zygotes | 97 zygotes | 22 | — | 1 | 1 | 100% | The first monkey model of human β-thalassemia | Huang et al. [51] |
| Cynomolgus monkeys | 2017 | Cynomolgus monkeys | CRISPR/Cas9 knockin | mCherry | Zygotes | First: 26; second: 10 | First (high-concentration HMEJ donor): 4/5 mCherry<sup>+</sup> blastocysts; second (low-concentration HMEJ donor): 1/4 mCherry<sup>+</sup> blastocysts | Verified HMEJ-based knockin at the embryo level | Yao et al. [55] |
| Cynomolgus monkeys | 2018 | Cynomolgus monkeys | CRISPR/Cas9 knockin | Oct4-GFP | Zygotes | 198 zygotes | 120 | 12/40 = 30% | 5 | 4 | 80% | The first monkey model of KI | Gui et al. [56] |
| Rhesus monkeys | 2019 | Cynomolgus monkeys | CRISPR/Cas9 knockin | c-kit | Pronuclear-stage embryos | 40 embryos | Optimal injection combination: 36 nt sense ssODN and CRISPR/nuclease mixture; editing efficiency: 30.8% | KI/KO efficiency verified at embryo level | Kumita et al. [54] |
| Chimera embryo | 2015 | Cynomolgus monkeys | Retroviral vector | GFP | ESC | — | 14 | 1/5 = 20% | 0 | 2 | — | The first monkey chimeras | Chen et al. [22] |
| SCNT | 2018 | Cynomolgus monkeys | — | — | — | — | 6/21 = 29% | 2 | 2 | 100% | The first monkey model of SCNT | Liu et al. [16] |
| Cynomolgus monkeys | 2019 | Cynomolgus monkeys | — | — | — | — | 325 | 16/65 = 25% | 5 | 5 | 100% | The first disease monkey model of SCNT | Liu et al. [64] |
| Cynomolgus monkeys | 2020 | Cynomolgus monkeys | CBE | LMNA | Zygotes | 86 zygotes | 41 | 6/11 = 55% | 5 | 4 | 80% | The first monkey model of BE | Wang et al. [60] |
| Rhesus monkeys | 2020 | Rhesus monkeys | CBE | MECP2 | Zygotes | — | Embryo level | Established a stable embryo editing system of RTT | Zhou et al. [62] |
| Cynomolgus monkeys | 2020 | Cynomolgus monkeys | CBE and ABE | Multiple loci | Zygotes | — | Embryo level | Simultaneously edit multiple loci at the embryo level | Zhang et al. [63] |

Pregnancy rate<sup>a</sup>: pregnancy/surrogate*100%; mutation rate<sup>b</sup>: mutation neonatus/total number of neonates*100%.
vectors carrying the enhanced green fluorescent protein (EGFP) gene into the perivitelline space or blastocyst cavity of early embryos, they successfully obtained five transgenic marmosets expressing the EGFP gene [18]. This is the first successfully constructed transgenic marmoset model, providing an efficient and practical animal model for the biomedical field. Moreover, they collected semen samples from the first-generation (F1) monkey model to obtain an offspring that carried the EGFP gene and proved the lineage inheritance of transgenic monkeys. However, the expression organization and strength of the five first-generation transgenic marmosets were not the same and the copy of EGFP and the integration site were also different, suggesting a random integration and uncontrollable expression strength. This is the common disadvantage of constructing transgenic monkey models with viral vectors. Following GFP transgenic rhesus monkeys and common marmosets, GFP transgenic cynomolgus monkey has been generated by Seita et al. [25]. They compared and analyzed the effects of lentivirus injection before and after fertilization and found that cynomolgus monkeys with lentivirus injection before fertilization can express GFP all over the body, indicating that the earlier the virus injection, the easier to obtain a homozygous animal model. However, injecting virus before fertilization can increase the rate of miscarriage, raising the question that the efficiency of virus integration and the development rate of embryos need to be balanced.

In 2010, the first genetically modified monkey in China was created by infecting embryos with simian immunodeficiency virus (SIV) in the early cleavage stage [26]. Two living infant monkeys expressed EGFP stably. Both in vivo and in vitro experiments showed that SIV infection had no significant effect on the development rate of embryos compared to the noninfected negative control group. The construction efficiency of transgenic monkeys has been significantly improved by improving and developing ARTs and using an SIV-based vector to infect early cleavage-stage embryos. However, flow cytometry showed that less than 30% of the cells from the transgenic baby monkeys expressed EGFP, suggesting that only part of the early embryonic cells was infected by the virus, resulting in the transgenic rhesus monkey chimera mutants.

Besides HD, more transgenic NHP disease models have been reported. Parkinson’s disease (PD) models were established in rhesus monkeys by Niu et al. in 2015 [27]. Interestingly, in the lentiviral vector, mutant α-syn (A53T) and ECFP are linked via F2A (which could self-cleave in cells after transcription). In this way, the length of the integrated fragment is reduced due to one vector expressing two proteins. It is also convenient to judge whether the mutant gene is integrated successfully by detecting the expression of ECFP protein. It is worth mentioning that the transgenic PD monkey models showed similar nonmotor symptoms to humans, which provided a suitable animal model to monitor early nonmotor symptoms of PD. Nonmotor symptoms have a significant impact on patients. However, the clinical research is hindered by lacking data in early stages of PD and difficulties in follow-up of patients. Therefore, it is extremely important and necessary to identify biomarkers and treatment strategies based on established PD monkey models. The autism-like syndrome in cynomolgus monkeys was reported by Liu et al. in 2016 [28]. These monkeys carry methyl-CpG binding protein 2 (MECP2) gene duplication. They injected lentivirus carrying the hSynapsin-HA-hMECP2-2a-GFP cassette into the perivitelline space of mature oocytes. The F1 monkey models expressed the human MECP2 gene in their brain and exhibited autism-like behaviors. The F1 monkeys carried the human MECP2 gene and displayed reduced social activities, proving the success of germline transmission of the transgene. Recently, Cai et al. discovered that MECP2 coexpressed genes significantly enriched in GABA-related signaling pathways, which could reduce beta synchronization within fronto-parieto-occipital and clarify abnormal locomotive behaviors in MECP2 duplication syndrome individuals [29]. The findings based on this model imply the feasibility and reliability of using genetically engineered NHPs to study psychiatric diseases.

The evolution and development of the human brain have always been a research hotspot. Nonhuman primates are ideal experimental animals for studying the evolution of the human brain. After birth, the expression level of human MCPH1 is high during brain development but the expression level of MCPH1 in nonhuman primates such as monkeys is relatively low. This is perhaps due to the unique structure and function of human MCPH1 in the human brain. In order to explore the function of this gene, Shi et al. used lentivirus transfection to overexpress human MCPH1 in rhesus monkeys [30]. The results showed that the transgenic monkeys exhibited human-like brain developmental neoteny and exhibited enhanced short-term memory, but not enlarged brain size as expected. These suggested that a single genetic change cannot simulate the complex evolution of the human brain.

In general, using viral vectors to integrate exogenous genes into the embryonic genome is the earliest technology to generate transgenic monkeys. Initially, the exogenous gene is GFP or EGFP as a reporter; later, some pathogenic genes were integrated to generate disease models. Transgenic monkeys have made great contributions to disease research; however, it is only limited to the disease that could be induced by exogenous gene expression. Moreover, viral vector-mediated gene transfer has other fatal limitations, such as the gene size of insertion, uncontrollable expression strength, and random insertion [31]. In order to overcome these shortcomings, the second-generation gene editing technology has been developed.

2.2. Precise Knockout Genetically Modified NHP Models. The three most popular methods of second-generation gene editing technologies are zinc finger nucleases (ZFNs), transcription activator-like effector nucleases (TALENs), and clustered regularly interspaced short palindromic repeat/ Cas9 (CRISPR/Cas9) [31, 32]. Here, we reviewed the published targeted gene knockout monkey models generated by these three methods.

Liu et al. successfully constructed the first TALEN-mediated methyl-CpG binding protein2 (MECP2) knockout rhesus and cynomolgus monkeys. According to PCR and
gene sequencing analysis, no off-target or plasmid integration was found on the genome [33]. MECP2 is an X-linked gene, which is extremely essential to the growth and development of humans and monkeys. Duplication of MECP2 gene will cause MECP2 duplication syndrome, showing similar symptoms to autism, whereas MECP2 gene loss-of-function mutations will cause Rett syndrome (RTT), a severe neurodevelopmental disorder [28, 33]. The monkey model of RTT syndrome showed similar physiological, behavioral, and brain structural abnormalities to human RTT symptoms [34]. Therefore, it is a great opportunity to analyze the pathogenic mechanism and treatment methods of RTT through an in-depth exploration of monkey models. Ke et al. used TALEN to knock out biallelic MCPH1 and generated a cynomolgus monkey model of autosomal recessive primary microcephaly (MCPH), a genetically neurodevelopmental disorder [19]. The MCPH1 protein can inhibit the expression of telomere reverse transcriptase. If the MCPH1 gene mutation leads to a decrease in the MCPH1 protein level, the patients will experience a significant reduction in brain size and height. Some patients will have decreased intelligence and other neurological diseases. The main feature in patients with MCPH at the cellular level is premature chromosome condensation (PCC), which also appeared in the monkey MCPH model. Sato et al. used ZFN and TALEN technologies to target the IL2RG gene and constructed five and four X-linked severe combined immunodeficiency (X-SCID) marmoset models, respectively [35]. This is the first batch of monkey models constructed by ZFNs. By injecting mRNA into embryos at the prokaryotic stage and analyzing it after culturing for seven days, TALEN was found to have a higher mutation rate than ZFNs. This may be one of the reasons why TALENs are used more often than ZFNs, although both of them are not commonly used in gene editing.

Due to the difficulties and time-consuming operations of ZFNs and TALENs, more and more genetically modified monkey models are now constructed by CRISPR/Cas9. CRISPR/Cas9 is simpler to operate, more versatile, and more efficient. In 2014, the same year that TALEN was first used to construct monkey models, the first monkey model that used CRISPR/Cas9 was also constructed successfully. Ni and colleagues injected Cas9 mRNA and gRNAs targeting Nr0b1, Ppar-g, and Rag1 into one-cell-stage embryos [36]. Ppar-g and Rag1 knockout cynomolgus monkey models were born when the article was published. However, no Nr0b1 gene mutation was detected in the newborn monkeys. One year later, a total of 3 surviving cynomolgus monkeys were born by the remaining 8 pregnant female monkeys but only one of the newborn monkeys was detected with the Rag1 mutation. Moreover, mutations have been detected in the somatic and germ cells of aborted fetuses, suggesting that gene mutations caused by the CRISPR/Cas9 technology can also be transmitted through the germ line [37]. In 2015, three knockout monkey models based on CRISPR/Cas9 were published, including the P53 knockout cynomolgus monkey model [20], Duchenne muscular dystrophy (DMD) thesis monkey model [38], and X-linked adrenal hypoplasia congenita-hypogonadotropic hypogonadism (AHC-HH) cynomolgus monkey model [39]. Here, we focused on the research progress of DMD in the NHP models. DMD is an X-linked recessive genetic disease. Patients are mostly male and the incidence is about 1/3500 to 1/5000 [40–42]. The main symptom of DMD is the deterioration of muscle function. The patient died from respiratory failure or heart failure around the age of 30s. In recent years, due to the use of ventilators to treat DMD patients with respiratory decline, the death of DMD patients has declined, which means that the death ratio of heart failure has gradually increased [43]. In 2015, a rhesus monkey model of DMD was generated [38] with no off-target occurrence [44]. Through the exploration and study of this model, it is very likely that new treatments for DMD could be developed. In 2017, the first cynomolgus monkey model of autism spectrum disorders (ASD) caused by SHANK3 gene mutation was generated by Zhao et al. [45]. The strategy of mutating the SHANK3 gene is to target the 6th and 12th exons of the SHANK3 gene, resulting in large fragment deletion of the SHANK3 gene. They implanted 116 embryos into 37 surrogate mothers, and only three female monkeys were pregnant. The pregnancy rate was extremely low, which is only 8.1%. The exact reason for the low pregnancy rate is unknown, but it suggests that the expression of the SHANK3 gene is very important for the early development of primates. In 2018, Zhang et al. constructed a cynomolgus monkey model with biallelic SIRT6 mutations, which provides a suitable model for studying human perinatal lethality syndrome [46]. SIRT6 gene deletion delays neuronal differentiation through transcriptional activation of long non-coding RNA H19 (a developmental inhibitor). The cynomolgus monkey model with SIRT6 mutation exhibits delayed embryonic development and died soon after birth. In 2019, Zhou et al. targeted exon 21 of the SHANK3 gene to construct five cynomolgus monkey models with ASD and Phelan–McDermid syndrome [47]. The F1 generation monkey model was generated using F0 generation germ cells, which made up for the shortcomings of an insufficient number of SHANK3 mutant monkey models. It is worth noting that there is a homozygous model for the mutant allele in the F0 generation, suggesting that the development of the CRISPR/Cas9 technology will eventually increase the mutation rate and mutation homozygous rate of the gene. Qiu et al. constructed five BMAL1 gene knockout circadian disruption cynomolgus monkey models, providing a suitable animal model for studying biological rhythm disorder [48]. Tsukiyama and colleagues constructed a monkey model of autosomal dominant polycystic kidney disease (ADPKD), an autosomal dominant genetic disease with a high incidence [49]. Yang et al. constructed a PD monkey model with large fragment deletion of the PINK1 gene. The monkey model showed similar symptoms of neuronal loss in humans, which has not been observed in mice or other models [50]. It is known that gene therapy may cure β-thalassemia; however, there is no suitable animal models for evaluating the safety and efficacy of this therapeutic strategies in vivo. Huang et al. constructed a HBB-deficient M. fascicularis monkey model that could be used to study the mechanism of β-thalassemia and the long-term safety and efficacy of gene editing therapies targeting HBB [51]. Since the mouse models of ADPKD, PD, and β-thalassemia exhibit large
differences to humans in physiology, monkey models of these diseases made up for the shortcomings of the mouse models, providing an ideal way to study disease pathogenesis and discover therapy strategies.

In summary, with the development and improvement of the second-generation gene editing technology, more and more genetically precise knockout NHP models have been constructed. These models provide a suitable platform for studying the mechanism of complex diseases and the exploration of treatment methods. Compared with retroviruses, gene knockout based on the second-generation gene editing technology has a significant advantage of controllable edit sites. But there are still many shortcomings in the genetically precise knockout technology. For example, CRISPR/Cas9 has the disadvantages of low gene knockout efficiency, mosaicism, restriction of PAM sites, and off-target effects.

2.3. Precise Knockin of Genetically Modified NHP Models. The second-generation gene editing technology can activate DNA damage repair mechanisms in cells by targeting double-strand breaks (DSBs) \([52]\). DNA damage repair mechanisms include nonhomologous end-joining (NHEJ) and homology-directed repair (HDR) \([53]\). NHEJ is the primary and efficient DNA damage repair mechanism. It will cause indel (including insertion or/and deletion) mutations while repairing DNA damage. HDR repair occurs in the presence of a donor template. Through the specific design of the donor template, specific gene knockout or knockin could be achieved. The existence of the template makes the gene editing more accurate, which can achieve the true "precision gene editing." However, the repair efficiency of HDR is very low. Kumita et al. verified the efficiency of specific gene knockin and knockin at the embryo level \([54]\). They found that the knockout efficiency of c-kit and Shank3 can reach more than 70% and even 100% editing efficiency. But c-kit knockin efficiency is only about 30% under optimal conditions.

In spite of technical difficulties, in recent years, accurate gene knockin animal models have been successfully constructed. In 2017, based on the principle of HMEJ and CRISPR/Cas9 technology, Yao et al. successfully conducted the precise mCherry gene knockin in cells, mouse embryos, monkey embryos, and mice, providing a new way to generate animal models \([55]\). In 2018, Cui et al. constructed the first Oct4-hrGFP precision knockin monkey models, achieving a major breakthrough in the gene editing field \([56]\). But the NHEJ repair is detected, and the HDR repair efficiency needs to be further improved. HDR repair is a promising gene repair mechanism, because it can achieve multiple types of gene mutations and does not have the uncertainty like NHEJ repair. The research and development of HDR repair will bring tremendous progress in the field of life sciences.

2.4. NHP Models via Precise Point Mutation. Genetic diseases are caused by multiple types of gene mutations, including point mutations, deletions, duplications, increase or decrease in copy numbers, indels, and insertions. Among them, the top three pathogenic gene mutations are point mutation, deletion, and duplication. It is worth noting that point mutations account for more than 50% of all pathogenic mutation types and it is mainly based on the change of AT to GC base pair (about 47.5% of point mutations) \([57]\). Since the CRISPR/Cas9 system often causes nonideal mutations such as indels and large fragment deletions, it seems not efficient and reasonable to construct animal models with single point mutations by CRISPR/Cas9. To address these difficulties, base editors (BEs) came into being. The BE system requires three elements: a Cas9 nickase fused to a nucleobase deaminase enzyme, a gRNA targeting Cas9 to a specific locus, and DNA glycosylase inhibitor. It includes cytosine base editors (CBE) (C to T mutation) and adenine base editors (ABE) (A to G mutation), which could cause precise point mutations in the target window without completely breaking the DNA double strand \([58, 59]\). This determines that BEs could introduce single point mutations more efficiently and more beneficial for modeling-related diseases. In 2020, Wang et al. injected BE mRNA and sgRNA targeting the LMNA gene into monkey zygotes to construct the Hutchinson-Gilford progeria syndrome (HGPS) monkey model, which is caused by single point mutation (1824 C>T) \([60]\). Mutations in LMNA can cause the accumulation of a toxic-truncated protein called progerin, causing changes in the structure and function of the nuclear membrane and ultimately leading to premature aging of children. At present, the treatment of this disease has been explored using CRISPR/Cas9 at the cellular level and in the HGPS mouse model. The results showed that gene editing therapy alleviated the phenotypes in cells and symptoms in mice models, manifested by the moderated nuclear phenotype and the prolonged life span of the mouse model. However, because of the organ targeting limitation of the adeno-associated virus (AAV), the immunohistochemical results showed that the nuclear structure of liver, heart, and skeletal muscle cells in mice improved significantly but there was no significant improvement in the lung, kidney, and aorta \([61]\). Zhou et al. verified the efficiency of the BE system at the embryonic level in mice and macaques, providing an effective reference for the later establishment of T158M mutant RTT animal models \([62]\). Zhang et al. coinjected SpCas9-based ABE mRNAs, SaCas9-based SaKKH-BE3 mRNAs, and their corresponding sgRNAs into monkey embryos and found that CBE and ABE can function in the same cell, suggesting an effective treatment strategy for polygenic diseases \([63]\). The NHP models will further improve our understanding of diseases and help us to comprehensively and accurately evaluate the safety and effectiveness of the drug before the clinic.

2.5. NHP Models via Somatic Cell Nuclear Transfer (SCNT). Because germ cells are difficult to culture and differentiate in vitro, they are mostly obtained in vivo. However, the number of germ cells is very limited, which greatly affects the development of NHP models. Somatic cell nuclear transfer (SCNT) enables CRISPR/Cas9 to edit somatic cells; therefore, the edited cells can be cultured in vitro to expand the number of cells. Part of the cells is used for genetic sequencing. If the sequencing is qualified, the remaining cells are transferred to
the enucleated oocytes and finally transplanted to the surrogate mother to produce genetically uniform monkeys.

After Dolly the cloned sheep was produced in 1996 [15], many somatic cell-cloned mammals have been reported. However, as the closest evolutionary relative to humans, the somatic cell-cloned NHPs have not been successfully constructed. It is the problem that the whole world is looking for a breakthrough. In 2018, Chinese scientist Liu et al. used histone demethylase and deacetylase inhibitors to increase the embryonic development rate of somatic cell clones and successfully constructed two SCNT cynomolgus monkey models [16]. In 2019, they isolated fibroblasts from a CRISPR/Cas9-mediated BMAL1 knockout monkey model [48] and successfully cloned five homozygous macaque monkeys with BMAL1 mutation-induced biological rhythm disorder without mosaicism [64]. The success of NHP SCNT provides a new and efficient way to construct NHP models with uniform genetic backgrounds.

Now, we can obtain gene-specific knockin, knockout, or point mutations NHP models through embryo engineering. However, gene editing technologies and the construction of NHP models still have many imperfections.

3. Challenges and Opportunities of Existing Animal Models

3.1. Shortcomings of Existing Animal Models. Rodents are the most widely used experimental animal models (more than 90%) [65–73]. Although the mouse model is low cost and easy to operate, it has limitations in revealing human biology. The biological system of mice, especially the immune system, is not completely consistent with the human immune system. This is characterized by many differences in the interaction of innate immune molecules. For example, mice can express immune molecules TLR11, TLR12, and TLR13 when lacking the functional immune molecule TLR10. However, humans cannot express those three immune molecules if lacking functional TLR10. Many experimental studies have been carried out in immunodeficient mice; therefore, it is difficult for the mouse model to truly simulate the immune response process of the human immune system, which reduces the reliability of prediction in clinical applications. Many human pathogenic factors and drugs are species specific. Some pathogens are only targeted primates, not mice. All of these problems directly affect the smooth transformation from biomedicine to clinical practice. In recent years, more and more researchers tend to use large animal models such as pig models. With more than 80% similarities in analysis parameters, domestic pigs are closely related to humans in terms of the immune system. Therefore, pigs can be used as a powerful animal model to study immune diseases [74, 75]. Pigs are also potential organ donors for humans. NHPs transplanted with pig organs can survive for months or even years [76]. This renewed the interest in its potential to ease organ shortages. However, widespread application of pig organ transplantation is limited by immunosuppressive complications, chronic rejection. In addition, porcine embryonic stem cell resources have not been widely developed and the culture system and differentiation system are not mature enough. These further limited the application of pigs as animal models.

3.2. Advantages and Challenges of NHP Models. NHPs are very similar to humans in the central nervous system, immune system, and cardiovascular system [77], NHPs played a vital role in scientific research in recent years. The similarity between NHPs and the human genome is as high as 98%, which can be used to extensively study human-related genetic diseases. NHPs have a sulcus structure and a well-developed prefrontal lobe, which are significantly important for study and memory. It is an excellent animal model for the research of central nervous system-related diseases. Currently, NHP models for Parkinson’s disease [27, 50], microcephaly [19], autism [28, 45], and other related diseases have been established and breakthroughs in those areas are expected. Meanwhile, NHPs can serve as important models for developing and evaluating effective treatment strategies. Because of those advantages, NHP models have become more and more vital in recent years [78–82].

However, there are also many challenges in the application of NHP models, such as the reproductive cycle is longer than other experimental animals and the genetic manipulation is difficult. A study in crab-eating monkey embryos showed that the targeting rate of CRISPR/Cas9-based gene editing is not precise enough and the single-nucleotide polymorphism (SNP) effect also needs to be considered [36]. Moreover, in a study of somatic cell-cloned monkeys, the survival rate after nuclear transplantation with cumulus cell or monkey fetal fibroblasts is very low [16], which brings great difficulties to the follow-up work. However, as long as the NHP models are constructed, the characteristics of NHPs are incomparable by any other animals. They are closer to the real situation of human beings and also have higher persuasiveness and credibility. In general, compared with other animal models, NHPs have unique advantages, but meanwhile, they also face many challenges (Figure 2).

3.3. Quantity and Cost of NHP Models. In the past few years, the use of animal models has gradually increased. Mice are the dominant animal models because of their low price and ease of operation. Although nonhuman primates are small in number and difficult to operate, they still have an irreplaceable position in the research of human genetic diseases and neurological diseases. At present, more than 170000 NHPs are used in biomedical experiments worldwide each year. Among them, the vast majority of NHPs (about 70000) were used by the United States [83]. Europe and Japan are also in the leading position. Compared with other animal models, even with other large animal models, the number of successfully constructed NHP models is very low. The main reasons include the low mutation rate of embryos and the low survival rate of birth. In terms of the cost of animal models, as the experimental value of monkeys becomes more and more prominent, the cost of buying monkeys is rising, from US $2000 to more than US $10000, plus daily feeding costs and labor costs; the price will continue to increase, which brings a high-cost problem at the same
time. But considering the highly physiological, anatomical, and histological similarities of NHPs to humans, therefore, NHPs are worth using.

3.4. Ethics and Morality of Animal Models. Animal experimental ethics is a problem that must be faced by scientific research. The main problem in China is the contradiction between animal protectionism and experimental animal science, as well as the differences in culture and concepts in different regions. In recent years, the trend of keeping pets has developed rapidly, which makes people have strong feelings for animals. Without knowing the background of scientific research, they have some prejudices against animal experiments. At the same time, the laws and regulations in this field are not perfect, which leads to some bad phenomena. However, some principles have been introduced internationally to help solve such problems, such as the implementation of the 3Rs principle. 3Rs include the use of unconscious experimental materials to replace live animals (replacement), reduce the number of experimental animals in the experiment (reduction), and improve animal welfare (refinement) in the whole process of the experiment [84–89]. At present, this principle has also become a recognized global animal scientific standard in animal ethics. Therefore, the use of NHPs for research must be subject to strict moral review before it can be approved. For example, is it scientifically necessary to use NHPs? Is the design scientific? Is the degree of injury appropriate? These moral reviews will also promote the healthy development of animal ethics. In view of the challenges and difficulties faced by NHP experiments, some researchers have also put forward another point of view of animal experiments, that is, use high-throughput approaches to overcome these problems. Although this view is highly feasible, there is still a long way to go before it can be fully achieved. But on special scientific issues, if animal models must be used in experiments, they should also be used after strict scrutiny and ethical considerations.

4. Conclusion and Prospect

An important method for studying mechanisms of human disease and developing disease treatments is to establish effective animal models. However, many rodent disease models, such as PD, RTT, DMD, and ADS, are not able to recapitulate the same process in humans, leading to difficulties in using rodents to study these diseases. NHPs are very close to humans in terms of evolution, development, metabolism, and pathology. Fortunately, NHP models of these diseases show similar symptoms to human patients. Therefore, NHPs are the ideal animal models for studying human diseases, especially complex diseases.

Embryo engineering combined with different gene editing methods, in theory, can construct various NHP disease models. There are different methods to establish an animal model at different stages of embryo formation, such as sperm vector, SCNT, pronuclear microinjection, retrovirus infection, and embryonic stem cell method. SCNT has been successfully applied in NHPs, providing an efficient and practical way to construct NHP disease models. Because different gene editing methods and stages have different effects on the mutation rate, embryonic development conditions, and birth rate of NHPs, those methods should be selected according to different needs.

The most popular gene editing tools include virus-mediated transgene, ZFNs [90], TALENs [33, 91, 92], and the CRISPR/Cas system [93, 94]. Since the CRISPR/Cas9 system has the advantages of relatively simpler operation, shorter experimental period, and being more versatile and efficient than other gene editing methods, most NHP disease models were constructed by it. However, it is also necessary
to improve the repair efficiency of HDR or develop new methods to achieve the true “precision gene editing.” Nowadays, many new gene editing methods are emerging. For example, homology-independent targeted integration (HITI) strategy, a gene editing method based on NHEJ repair, can achieve targeted knockin in both mitotic and postmitotic cells [95]. The prime editing system includes three elements: Cas9 nickase, reverse transcriptase, and pegRNA. Interestingly, pegRNA has the dual role of guiding the other two elements to the target gene site and serving as a template for reverse transcription. In this way, prime editing could achieve targeted gene insertion, deletion, and all 12 types of point mutation without the donor template and DSBs [96]. The DdA-derived cytosine base editor (DdCBE) could achieve cytosine deamination modification on mRNA without unwinding the dsDNA [97]. With the emergence of these new technologies, now we can generate more animal models, bringing new hopes to the research and treatment of related diseases.

Due to the advancement of the gene editing technology, in vivo gene therapy becomes possible. How to deliver gene editing systems to target cells or organs has become an urgent problem. The primary delivery method in vitro is nonviral delivery systems (electroporation, microinjection, and lipid nanoparticles), and the in vivo delivery method is viral delivery systems (AAV and lentivirus) [98]. With the widespread use of viral vectors, potential problems are also gradually revealed. At present, the main problem faced by therapeutic genome editing is the safety and effectiveness of the delivery system [99]. In NHPs, a viral delivery system (AAV) was used to deliver meganuclease [100] and nonviral vectors (lipid nanoparticles) to deliver ABE [101]; they specifically reduce the expression of PCSK9. The results showed that LDL-C levels were significantly reduced for a long time, providing a safe and effective strategy for the treatment of cardiovascular diseases. However, intravenous injection of AAV in large doses can cause acute liver failure and shock in monkeys [102], suggesting that the use of AAV needs to consider more factors and safer delivery methods need to be developed. In general, the use of NHP models can more comprehensively reflect the safety and effectiveness of delivery systems or gene editing strategies, providing a strong reference for clinical applications and transformation.

In summary, the construction of NHP disease models provides a valuable research platform for human diseases, especially for those complicated diseases that cannot be successfully modeled in other animals. The use of NHP models to verify disease treatment strategies before the clinic can predict treatment effects more accurately and provide a powerful reference for clinical treatment. NHP disease models have promoted the development of translational medicine and brought new hopes to understand the underlying disease mechanisms and explore disease treatment methods. Therefore, NHPs have an indispensable status in the field of life sciences.

Conflicts of Interest
The authors declare that there is no conflict of interest regarding the publication of this article.

Authors’ Contributions
Mei Huang, Jiao Yang and Peng Li contributed equally to this work.

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