# Targeted gene correction and functional recovery in achondroplasia patient-derived iPSCs

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# Abstract

6 Achondroplasia (ACH) is the most common genetic form of dwarfism and belongs to dominant 7 monogenic disorder caused by a gain-of-function point mutation in the transmembrane region of 8 FGFR3. Stem cells and gene-editing technology provide us with effective methods and ideas for 9 research and treatment of ACH. Here we generated non-integrated iPSCs from one ACH girl's skin and one ACH boy's urine via Sendai virus. We found that the chondrogenic differentiation 10 ability of ACH iPSCs was confined compared with healthy iPSCs. When the mutation of ACH 11 12 iPSCs from skin and urine was precisely corrected by CRISPR-Cas9, the chondrogenic differentiation ability of them could be restored. Furthermore, these corrected iPSCs displayed 13 pluripotency, maintained normal karyotype, and did not demonstrate off-target indels. Our study 14 may provide important theoretical and experimental basis for the stem cell research and treatment 15 of ACH. 16

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# 18 Introduction

19 Achondroplasia (ACH), the most common genetic form of short-limb dwarfism, is autosomal dominant monogenic disorder (MGD) caused by a gain-of-function point mutation in the 20 transmembrane region of fibroblast growth factor receptor 3 (FGFR3). There are currently two 21 mutation sites reported - Gly380Arg and Gly375Cys, the former occupies a vast majority of ACH 22 patients<sup>1,2</sup>. Because homozygous ACH patients have a much more severe phenotype and rarely 23 24 survive<sup>3,4</sup>, most of the ACH patients seen are heterozygous mutations. The estimated frequency of ACH is 1 in 25,000 (Male 1/15000), with at least 80% of the cases being sporadic<sup>5</sup>. Like many 25 26 other MGDs, there are no effective therapeutic methods for ACH even though the mutation has

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been found<sup>1</sup>. Fortunately, stem cell research provides new hope for potential ACH treatments.

Patient-specific stem cells can aid scientists in the investigation of specific molecular mechanisms and can be used to screen for the discovery of new drugs<sup>6,7</sup>. In addition, with powerful genome editing tools used, such as clustered regulatory interspaced short palindromic repeat (CRISPR)-Cas9 system<sup>8</sup>, mutation can be corrected in ACH stem cells *in vitro*. Through screening and safety assessment in animal models *in vivo*, stem cell transplantation therapy may provide an effective and novel treatment method<sup>6</sup>.

In this study, we isolated and cultured skin, urine, and white adipose-derived somatic cells 42 from three ACH patients of Gly380Arg mutation. Further we generated non-integrated iPSC lines 43 from one ACH girl's skin and one ACH boy's urine by Sendai virus. However adipose-derived 44 mesenchymal stem cells (AD-MSCs) could not be reprogrammed to iPSCs. We found the 45 46 chondrogenic differentiation ability of ACH iPSCs was confined compared with that of healthy iPSCs. When the mutation of ACH iPSCs from skin and urine was precisely corrected by CRISPR-47 Cas9, the chondrogenic differentiation ability of them was restored. Via cell immunofluorescence, 48 49 Alkaline Phosphatase (AP) Staining, karyotyping and sequencing analysis, we found these corrected iPSCs continued to keep pluripotency, maintained normal chromosomal number and 50 51 structure, and didn't display off-target. Our study may provide important theoretical and 52 experimental basis for the stem cell research and treatment of ACH.

#### 53 **Results**

#### 54 Identification of ACH Patients and Isolation and Culture of Somatic Cells

55 We recruited three ACH patient donors, including an 8-year-old girl, a 7-year-old boy and one 56 37-year-old adult man. Genetic identification was confirmed by DNA genome sequencing, and 57 the donors are all heterozygous mutations of a G-A transition at nucleotide 1138 of the cDNA of 58 FGFR3 (Fig. 1a). In addition, their clinical phenotypes are consistent with the NIH-defined characteristics, such as short arms and legs, enlarged head, and prominent forehead. We 59 punched skin tissue and collected urine from the two children to culture skin fibroblasts (SFs) and 60 urine-derived cells (UCs). Via liposuction, we obtained adipose tissue from the adult man to 61 62 culture AD-MSCs. After the small pieces of skins were placed and cultured in the dish for 2 weeks, SFs gradually climbed out to begin to proliferate (Fig. 1b, c). After urine-derived pellets were 63 cultured for one week, UCs formed colonies and showed epithelial cell morphology (Fig.1d, e). 64 After SVF from adipose tissue was seeded and cultured for 2-3 days, AD-MSCs grew out and 65 66 exhibited a typical fibroblast-like morphology (Fig. 1f, g).

67 Generation of Non-Integrated iPSCs from ACH Patients

68 To generate non-integrated iPSCs from ACH patients, we used Sendai virus to transfect cells. 69 Three weeks after transduction of SFs and UCs, lots of ES cell-like colonies appeared (Fig. 2a, 70 b). After picking up and expanding single colony, we established non-integrated iPSC lines from girl's skin (GF) and boy's urine (BU) (Fig. 2c, d). These iPSCs expressed pluripotent protein -71 NANOG, OCT4, SOX2, SSEA-4, and TRA1-60, while didn't express SSEA-1 (Fig. 2g). They also 72 73 showed AP positive (Fig. 2h). In addition, karyotyping analysis indicated that they maintained 74 normal chromosomal number and structure (Fig. 2i). However, it was surprised that iPSCs could 75 not be generated from AD-MSCs. Although small and unhealthy colonies could appear (Fig. 2e), they gradually became apoptotic and eventually died (Fig. 2f). 76

#### 77 Chondrogenic Differentiation Capability of ACH iPSCs

78 To Fig.out whether the mutations of ACH iPSCs from skin and urine would affect their function, we performed detection of their chondrogenic differentiation capability. We used healthy human 79 80 iPSCs and ACH iPSCs side by side to induce chondrogenic differentiation in a multi-step method 81 (Fig. 3a). We found that the chondrogenic clusters from ACH iPSCs were less and smaller than 82 those from healthy iPSCs (Fig. 3b). By Safranin O staining, we discovered the cartilage tissue 83 derived from ACH iPSCs showed less and weaker positive areas and lower cartilage density than that from healthy iPSCs (Fig. 3c). These results indicated that the cartilage tissue derived from 84 ACH iPSCs could produce less and thinner cartilaginous extracellular matrix. RT-gPCR results 85 also exhibited that cartilage tissue derived from ACH iPSCs expressed lower chondrocyte-specific 86 genes - SOX9, COL2A1, and ACAN (Fig. 3d). Our results suggested the chondrogenic 87 differentiation ability of ACH iPSCs was confined compared with that of healthy iPSCs. 88

#### 89 Gene Correction of ACH iPSCs by CRISPR-Cas9

To design sgRNAs, we input 64 bp DNA sequences of ACH FGFR3 around the point mutation 90 91 site into Guide Design Resouces. More than 10 sgRNAs were generated and we selected the 92 sg2RNA of highest score for use (Fig. 4a). To correct the point mutation in ACH iPSCs, we used 131 nucleotides ssODN donor as the homology arm which contained the point mutation site (Fig. 93 94 4a). 24-48 hours after CRISPR-sgRNAs and ssODNs were transected into ACH iPSCs from skin 95 and urine, we detected the transfection efficiency by FACS. The RFP positive ratio was 3.6% (Fig. 96 4b). By sequencing analysis of more than 100 RFP positive single cell colonies from skin and more than 100 RFP positive single cell colonies from urine, we found one completely corrected 97 cell line from skin and one completely corrected cell line from urine (Fig. 4c), and the total 98 99 efficiency was less than 1%. We analyzed the two corrected iPSC lines and found they expressed 100 pluripotent protein - NANOG, OCT4, SOX2, SSEA-4 and TRA1-60, while didn't express SSEA-1

(Fig. 4d). They also indicated AP positive (Fig. 4e). The karyotyping analysis verified they
 maintained normal chromosomal number and structure (Fig.4f). Sequencing analysis of potential
 offtarget sites given by Guide Design Resouces showed that no off-target indels were identified.

#### 104 Functional Recovery of Corrected ACH iPSCs

To detect whether the function of corrected ACH iPSCs from skin and urine was restored, we 105 performed the detection of chondrogenic differentiation ability of them. Through morphological 106 107 observation, we found that chondrogenic clusters from corrected ACH iPSCs obviously increased 108 compared with those from uncorrected cells (Fig. 5a). Via Safranin O staining, we found there 109 were more and stronger positive areas in cartilage tissue from corrected ACH iPSCs than that from uncorrected cells (Fig. 5b). EdU cell proliferation assay displayed corrected cells possessed 110 111 higher positive radio than uncorrected cells (Fig. 5c). Compared with cartilage tissue from ACH iPSCs, RT-qPCR results also revealed cartilage tissue from corrected ACH iPSCs expressed 112 113 higher chondrocyte-specific genes - SOX9, COL2A1, and ACAN (Fig. 5d). These results 114 suggested the chondrogenic differentiation ability of corrected ACH iPSCs was restored.

#### 115 **Discussion**

Whenever about 25,000 children are born, one of them may be an ACH patient<sup>5</sup>. For ACH 116 patients, not only will they lose a lot of opportunities to get education and job because of their 117 118 short stature, but they also have many complications, such as hydrocephalus, obesity and sleep 119 apnea etc. The older they are, the more serious they will be, which will seriously affect their 120 longevity. The average life expectancy for this cohort was decreased by 15 years compared with 121 the US population<sup>5</sup>. Moreover, the social concern of this disease is very low. At present there are 122 no effective therapeutic methods for ACH. With the rapid development of stem cell biology and gene-editing technology, the research and treatment of ACH not only show promising, but also 123 124 provide ideas and experimental basis for the investigation of other MGDs<sup>6</sup>.

125 In this study we first collected and isolated three different tissue samples from ACH patients to culture somatic cells, including the most commonly used skin, the most easily available urine, 126 127 and a large amount of acquired adipose tissue that can produce multipotent MSCs with 128 chondrogenic differentiation ability. We found that, like skin cells, ACH patient urine-derived cells 129 could be efficiently reprogrammed into iPSCs, which may provide new donor cells for research of MGDs. However, to our surprise, ACH patient-derived AD-MSCs could not be reprogrammed into 130 iPSCs. In fact, previously reported study<sup>9,10</sup> and our unpublished results found that healthy human 131 AD-MSCs could be reprogrammed more efficiently to iPSCs than skin cells. Given ACH is a 132

regeneration dysfunction disorder of MSC-derived chondrocyte caused by FGFR3 mutation, our 133 134 results suggested that perhaps the point mutation affected the reprogramming ability of AD-MSCs. 135 Our initial hypothesis was that point mutations might affect the chondrogenic differentiation ability of ACH iPSCs. Indeed, our experimental results confirmed chondrogenic differentiation 136 ability of ACH iPSCs was confined compared with that of healthy iPSCs. When we used CRISPR-137 Cas9 to correct point mutation of ACH iPSCs, we obtained two completely corrected cell lines 138 (one from skin, another from urine) from more than two hundred single cell colonies. The efficiency 139 140 of precise homology directed repair (HDR) was less than 1%. Fortunately, we found these two completely corrected iPSC lines still displayed pluripotency and maintained normal karyotype. 141 142 Sequencing analysis of potential offtarget sites suggested no off-target indels were identified. Finally, we detected whether the function of corrected ACH iPSC could be improved. Via 143 144 chondrogenic differentiation, EdU cell proliferation assay, and RT-qPCR experiments, we found their chondrogenic differentiation ability was indeed restored compared with uncorrected cells. 145

In summary, our study results of ACH iPSCs *in vitro* provided a solid and important foundation for the further exploration of ACH research and treatment *in vivo*. At present, we are constructing point mutation mouse model of ACH. Next, we will transplant the corrected ACH patient-derived MSCs or chondrocyte precursor cells into the mice to verify their function *in vivo* and explore the effect of this cell replacement therapy to ACH. Although challenges remain, the clinical application of patient-specific stem cells will be pursued through further advances in basic research (Fig. 6).

#### 152 Materials and Methods

#### 153 Research subjects and ethical statement

154 Research subjects in this study included two children and one adult man. Skin and urine 155 samples were obtained from the children. Liposuction was performed on the adult man. All human 156 subject protocols were reviewed and approved by the Ethical Review Board of the Renji Hospital, 157 Shanghai. All subjects gave signed informed consent.

#### 158 Isolation and culture of SFs

- 159 Skin biopsies were performed by using a sterile 3 mm skin punch from the locally anesthetized 160 lower legs. The skin tissue was cut in smaller pieces, placed in a 6-well plate and allowed to grow 161 for 2 weeks in fibroblast medium (DMEM, 10% fetal bovine serum (FBS), penicillin/streptomycin 162 (P/S) and glutamine). Above regents were all from ThermoFisher.
- 163 Culture of UCs

The culture of UCs followed the method established by Pei laboratory.<sup>11</sup> Briefly, urine samples 164 165 were collected into 50 ml tube and were centrifuged at 400 g for 10 minutes. The supernatants 166 were aspirated carefully, and about 1 ml volume at the bottom was kept in the tube. The remaining was resuspended by washing buffer (PBS containing amphotericin B and P/S) and centrifuged 167 again. After discarding supernatants, the pellets were suspended by 1 ml primary medium. Then 168 the cell suspensions were seeded into 12-well plates coated with gelatin and cultured in cell 169 170 incubator. The primary medium contained DMEM/Ham's F-12 1:1 (ThermoFisher), 10% FBS, 171 REGM Renal Epithelial Cell Growth Medium SingleQuots Kit (Lonza), amphotericin B (Selleck) and P/S. After 4 days, the medium was carefully changed to proliferation medium (Lonza REBM 172 173 BulletKit).

### 174 Isolation and culture of AD-MSCs

Adipose biopsy was obtained by liposuction. Isolation and culture of AD-MSCs followed our previous reports.<sup>12</sup> Briefly, adipose tissue was digested with 0.1% type I collagenase (Sigma) in PBS solution on a shaker at 37°C for 1 hour. Cell suspensions were then centrifuged to obtain stromal vascular fraction (SVF). The SVF was suspended and cultured in fibroblast medium.

# 179 PCR and Sanger Sequencing

180 Total genomic DNA was extracted from patient peripheral blood, SFs, UCs, AD-MSCs, and

- iPSCs by using the Genomic DNA Extraction Kit (TaKaRa). PCR was performed by using Q5
   High-Fidelity DNA Polymerase (New England Biolabs, NEB). *FGFR3* primers were
- 183 Forward: 5'-AGGAGCTGGTGGAGGCTGA-3',
- 184 Reverse: 5'-GGAGATCTTGTGCACGGTGG-3'.
- 185 PCR reactions were then purified by using GeneJET Gel Extraction Kit (ThermoFisher) and
- 186 sequenced by the Genewiz.

# 187 Generation of non-integrated iPSCs

Induction of iPSCs from SFs, UCs, and AD-MSCs was performed by using CytoTune iPS 2.0 Sendai Reprogramming Kit (ThermoFisher). Briefly, approximately 200,000 donor cells were seeded in a 6-well plate to culture. Two days later, cells were transduced with the Reprogramming Kit. 24 hours after transduction, the medium was replaced with fresh medium every other day. On day 8 after transduction, the donor cells were trypsinized and 5000 cells were seeded into a 6-well plate. The media were replaced by E8 (StemCell) every day thereafter. 3-4 weeks after transduction, single cell colonies were picked for expansion and characterization.

# 195 Chondrogenic differentiation of iPSCs

196 This method referred to report of Tsumaki laboratory (Yamashita et al., 2014). After iPSCs 197 formed high-density cell colonies in E8, the medium was changed to initial chondrogenic medium, including DMEM/F12 (ThermoFisher), 10 ng/ml of Wnt3A (R&D), 10 ng/ml of Activin A 198 (Peprotech), 1% ITS (ThermoFisher), and 1% FBS (ThermoFisher). On day 3, the medium was 199 changed to the basal medium, including DMEM (ThermoFisher), 1% ITS, 1% FBS, 2 mM L-200 glutamine, non-essential amino acids, 1mM Napyruvate (ThermoFisher), 50 µg/ml ascorbic acid 201 (Sigma), 10 ng/ml BMP2 (Peprotech), 10 ng/ml TGF<sub>β</sub>1 (Peprotech), and 10 ng/ml GDF5 202 203 (Peprotech). 10 ng/ml bFGF (Peprotech) was added to the chondrogenic medium from day 3 to 204 day 14. After Day 42, the medium was changed to fibroblast medium.

### 205 AP staining

AP staining was performed by using Vector Blue AP Substrate Kit (Vector). Procedure was conducted in accordance with the instruction.

# 208 Design of single guide RNAs (sgRNAs) and construction of vector

- sgRNAs were designed to target point mutation site by using the Guide Design Resouces of
- 210 Zhang lab (https://zlab.bio/guide-design-resources). Then oligo sgRNAs were annealed and
- ligated to the pSpCas9(BB)-2A-RFP plasmid which was digested with Bbs I (NEB) enzyme.

# 212 Transfection of CRISPR into iPSCs

One million iPSCs were resuspended in 800 µl cold PBS and mixed with targeting and singlestranded oligo DNA nucleotides (ssODNs) donor (5 µg CRISPR targeting plasmid and 40 µg donor). The cells were electroporated using the Human Stem Cell Nucleofector Kit 2 (Lonza) on Nucleofector 2b Device (Lonza). Cells were recovered and re-plated on plates with ROCK inhibitor. 24-48 hours after electroporation, about 5000 RFP positive cells were sorted and seeded into 100 mm plate by BD FACS Aria II. One week later, single colony was pick up and expanded for sequencing analysis.

# 220 Off-target effect analysis

18 potential "off-target" sites given by Guide Design Resouces were identified. DNA sequencing
 of PCR products amplified from these sites was performed.

#### 223 Cell immunofluorescence

iPSCs were fixed in 4% paraformaldehyde solution for 15 minutes at room temperature (RT). Next,
 they were permeabilized by using 0.1% Triton X-100 in PBS for 20 minutes at RT. After they were
 blocked for 1 hour in 5% goat serum, the cells were incubated with primary antibodies Nanog

(Abcam), SSEA-1 (Invitrogen), Oct4 (Cell signal), SSEA-4 (ThermoFisher, MA1-021), Sox2

(Epitomics, 2683-1), and TRA-1-60 (Abcam) overnight at 4°C. The cells were incubated with a
fluorescently coupled secondary antibody for 1 hour at RT, meanwhile the nuclei were stained
with 4',6-diamidino-2-phenylindole (DAPI; Sigma). Images were captured on a Leica confocal
microscope.

## 232 Karyotyping

The iPSCs were cultured in E8 with 0.1 µg/ml colchicine (ThermoFisher) for 2 hours. After medium being removed, they were incubated in 0.56% potassium chloride for 40 minutes, then fixed in methanol: acetic acid (3:1 in volume) overnight. The cell suspensions were dropped onto cool slides and stained with Giemsa (ThermoFisher) for 15 minutes. Finally, chromosome analysis was performed under the karyotyping image analysis system. More than 15 metaphase spreads of each sample were analyzed.

# 239 RNA extraction and RT-qPCR

Total RNAs were extracted from whole cells using TRIzol RNA Isolation Reagents (Thermofisher). cDNA was synthesized from total RNA with a Hifair II 1st Strand cDNA Synthesis Kit (Yeasen) for qPCR (TB Green Premix Ex Taq kit, TakaRa). PCR procedure was carried out according to the kit instructions. Primers used were as follows:

- 244 SOX9 F: AGACCTTTGGGCTGCCTTAT
- 245 R: TAGCCTCCCTCACTCCAAGA
- 246 COL2A1 F: TTTCCCAGGTCAAGATGGTC
- 247 R: CTTCAGCACCTGTCTCACCA
- 248 ACAN F: AGGCAGCGTGATCCTTACC
- 249 R: GGCCTCTCCAGTCTCATTCTC
- 250  $\beta$ -ACTIN F: TGGCACCACACCTTCTACAATGAGC
- 251 R: GCACAGCTTCTCCTTAATGTCACGC

# 252 EdU Staining

253 We used BeyoClick EdU-594 kit (Beyotime) for staining. Briefly, cells were incubated for 24

- 254 hours in 10 μM EdU medium at 37°C in incubator. After EdU solution was removed, the cells were
- washed twice in PBS. Then they were fixed and performed immunocytochemistry procedure.

# 256 Safranin O staining

257 Safranin O staining Kit (ScienCell Research Laboratories, Inc. #8384) was used to perform this

experiment. Procedure was in accordance with the instruction.

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#### 272 Conflicts of Interest

273 There is no potential conflict of interest.

#### 274 Author contributions

275 Z.H. and G.M.F performed experiments and analyzed data, L.Y., and L.F. performed experiments; W.W.Y. provided 276 experimental platform and financial support; Q.Y.R. conceived and designed the study, performed experiments,

analyzed data, provided financial support and wrote the manuscript.

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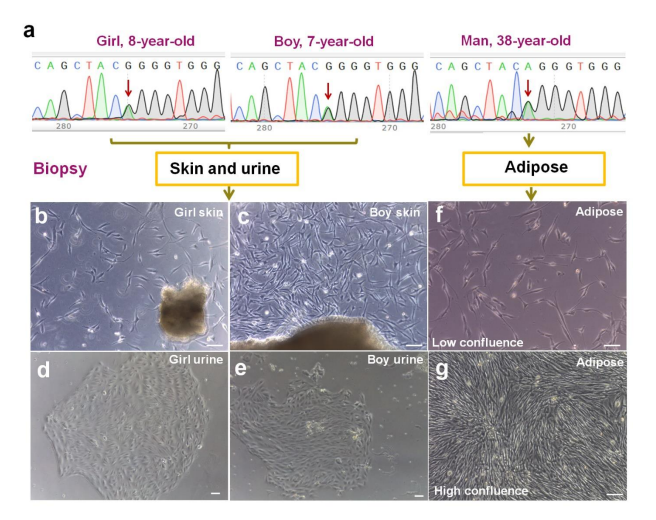
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# 301 Figures and figure legends

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- Fig. 1 Identification of ACH patients and isolation and culture of somatic cells. (a) Sequence of three ACH patient
   donors showed they are all heterozygous mutation of a G-A transition at nucleotide 1138 of the cDNA. (b, c) SFs from
   girl and boy. (d, e) UC colonies from girl and boy showed epithelial cell morphology. (f, g) AD-MSCs from adult man
   exhibited a typical fibroblast-like morphology. Bar in all panel: 10 μm.

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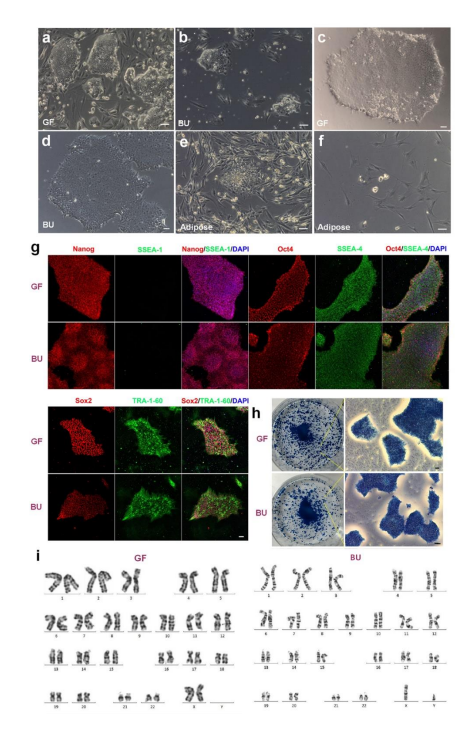


Fig. 2 Generation of non-integrated iPSCs from ACH patients and characterization of them. (a, b) 3 weeks ES
cell-like colonies from girl SFs (GF) and boy UCs (BU) after transfection. (c) Expanded iPSCs from GF. (d) Expanded
iPSCs from BU. (e) 3 weeks ES cell-like colonies from AD-MSCs after transfection, they were small and unhealthy. (f)
Colonies from AD-MSCs gradually became apoptotic and eventually died. (g) iPSCs from GF and BU expressed
pluripotent protein - NANOG, OCT4, SOX2, SSEA4 and TRA1-60, but didn't express SSEA-1. (h) iPSCs from GF and
BU showed AP positive. (i) iPSCs from GF and BU indicated normal chromosomal number and structure. Bar in all
panel: 10 μm.

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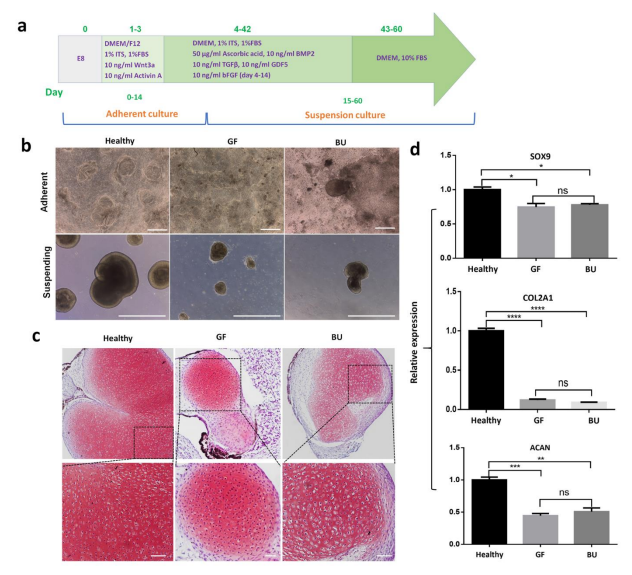


Fig. 3 Chondrogenic differentiation of ACH iPSCs. (a) A multi-step chondrogenic differentiation method. (b)
 Chondrogenic clusters from ACH iPSCs were less and smaller than those from healthy iPSCs. Bar: 100 μm. (c) Safranin
 O staining displayed there were less and weaker positive areas in cartilage tissue from ACH iPSCs than that from
 healthy iPSCs. Bar: 10 μm. (d) RT-qPCR results also exhibited that cartilage tissue derived from ACH iPSCs expressed
 lower chondrocyte-specific genes SOX9, COL2A1, and ACAN.

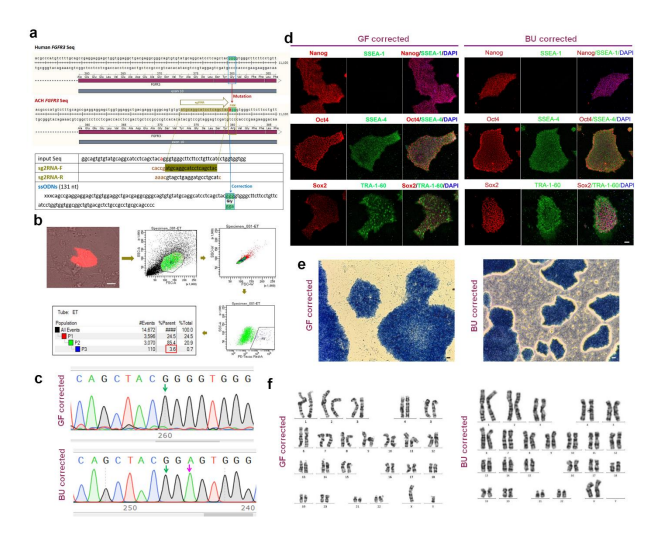
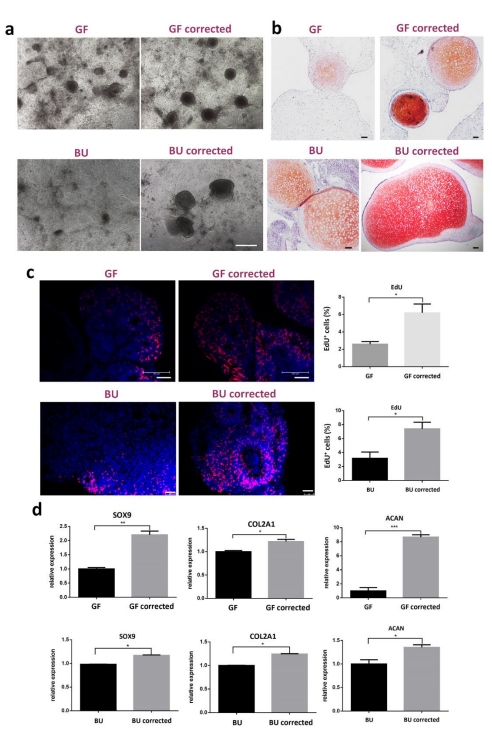
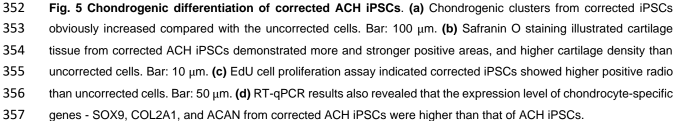


Fig. 4 Gene correction of ACH iPSCs by CRISPR-Cas9 and characterization of corrected iPSCs. (a) Designed
 sgRNA and ssODNs around the point mutation site. (b) FACS detection showed the RFP positive cells were 3.6%. (c)
 Two corrected iPSC lines showed normal DNA sequence (Pink arrow indicated synonymous mutation). (d) Corrected
 iPSCs expressed NANOG, OCT4, SOX2, SSEA4 and TRA1-60, but didn't express SSEA-1. (e) Corrected iPSCs
 indicated AP positive. (f) Corrected iPSCs showed normal chromosomal number and structure. Bar in all panel: 10 μm.

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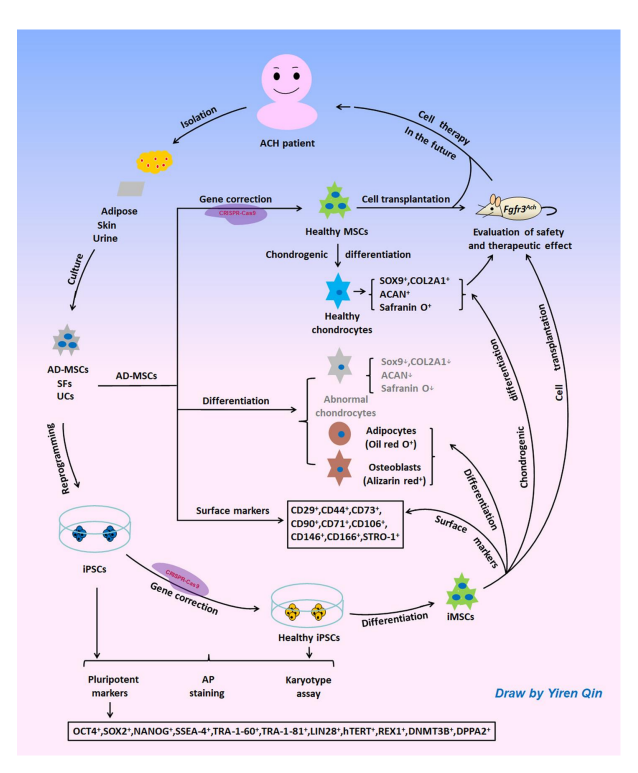


Fig. 6 Diagrammatic strategy of ACH patient-derived stem cell research. Somatic cells can be isolated and cultured
 from ACH adipose, skin, and urine, and further can be reprogrammed into iPSCs. After gene correction of iPSCs or
 AD-MSCs via CRISPR-Cas9, they can differentiate into healthy cells, such as MSCs or chondrocyte precursor cells.
 Then these healthy cells can be transplanted into ACH mouse model to assess their relative safety and therapeutic
 effects.