

A high-fidelity RNA-targeting Cas13 restores paternal *Ube3a* expression and improves motor functions in Angelman syndrome mice

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Angelman syndrome (AS) is a rare neurodevelopmental disorder caused by loss of function mutations in maternally expressed *UBE3A*. No gene-specific treatment is available for patients so far. Although intact and transcriptionally active, paternally inherited *UBE3A* is silenced by elongation of antisense long noncoding RNA *UBE3A-ATS* in neurons. Here, we demonstrated that RNA targeting of paternal *Ube3a-ATS* with a high-fidelity CRISPR-Cas13 (hfCas13x.1) system could restore *Ube3a* expression to similar levels as that of maternal *Ube3a* in the cultured mouse neurons. Furthermore, injection into lateral ventricles with neuron-specific *hSyn1* promoter-driven hfCas13x.1 packaged in adeno-associated virus (AAV-PHP.eb) could restore paternal *Ube3a* expression in cortex and hippocampus of neonatal AS mice for up to 4 months after treatment. Behavioral tests showed that expression of paternal *Ube3a* significantly alleviated AS-related symptoms, including obesity and motor function. Our results suggested that hfCas13x.1-mediated suppression of the *Ube3a-ATS* lncRNA potentially serves as a promising targeted intervention for AS.

INTRODUCTION

Angelman syndrome (AS) is a severe neurodevelopmental disorder, with an incidence of approximately 1 in 12,000–1/20,000 worldwide, which is characterized by seizures and severely impaired motor, cognitive, and language skills.¹ AS is mainly attributable to loss of maternally inherited ubiquitin protein ligase *E3A* (*UBE3A*), which mediates the targeted degradation of several protein substrates.^{2–6} Only the maternally inherited *UBE3A* allele in neurons is active due to selective silencing of paternal *UBE3A* by its antisense transcript, *UBE3A-ATS*. *UBE3A-ATS* is a polyadenylated transcript processed from *SNHG14 small nucleolar RNA host gene 14* (*SNHG14*) transcription and it overlaps with the *UBE3A* gene locus, resulting in early termination and degradation of *UBE3A* transcripts.^{7–10} Approxi-

mately 75% of AS cases are reportedly caused by deletions of the maternal chromosomal region *15q11.2–q13*. The remaining cases are attributable to other mutations in maternal *UBE3A*, paternal uniparental disomy, or imprinting defects.¹

There are currently no specific therapies available for the treatment of AS. However, activation of the intact, functional, paternal copy of *UBE3A* by eliminating the *UBE3A-ATS* long non-coding RNA (lncRNA) represents a promising potential therapeutic strategy. Genetic restoration of *Ube3a* by interferences with *Ube3a-ATS* has been reported to dramatically correct the aberrant neurobehavioral phenotypes in AS model mice.¹⁰ Antisense oligonucleotides have been developed to reactivate *Ube3a* expression by inhibiting *Ube3a-ATS* in AS mice and are now in phase I and II clinical trials.^{11,12} However, this treatment strategy requires frequent administration through the lumbar. Directed targeting of *Ube3a-ATS* with SaCas9 can also effectively decrease *Ube3a-ATS* transcription in mice by introducing double-stranded breaks (DSBs), insertions and deletions (indels), or adeno-associated virus (AAV) vector integration,^{13,14} which may induce unexpected genomic change.¹⁵

CRISPR-Cas13, a class 2 type VI RNA endonuclease, can bind and cleave single-stranded RNA guided by an engineered CRISPR RNA (crRNA).^{16–20} Gene therapies based on RNA-targeting nucleases are currently under exploration as a safer alternative to DNA editing Cas endonucleases, since they are accompanied by a lower risk of

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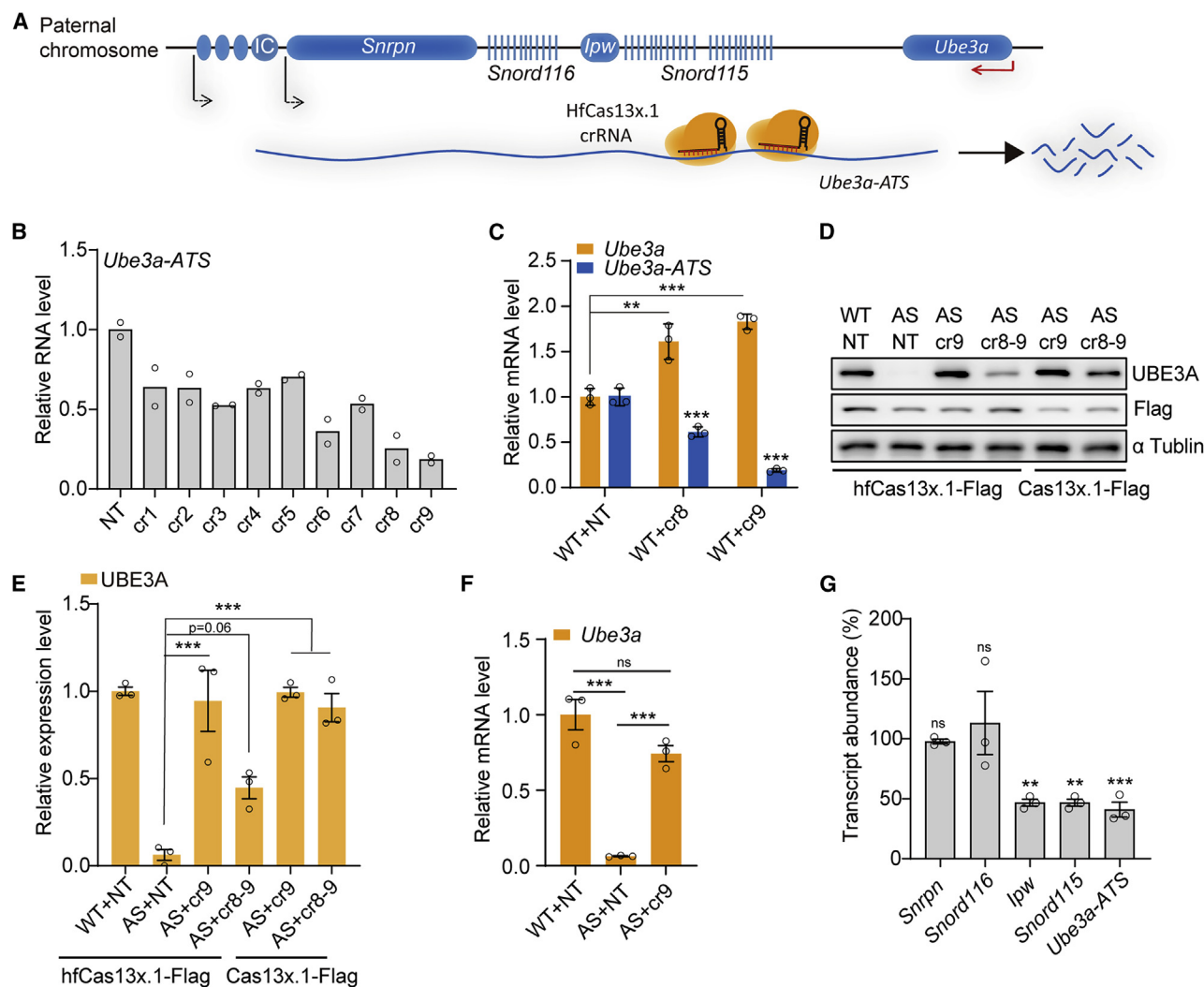


Figure 1. Screen of crRNAs targeting the lncRNA *Ube3a-ATS* to unsilence paternal *Ube3a*

(A) Schematic diagram of the *Ube3a* locus in the mouse genome. IC, imprinting center. (B) *Ube3a-ATS* mRNA expression levels under targeting by different crRNAs in N2a cells ($n = 2$ for all groups). (C) RT-qPCR analysis of mRNA expression in wild-type (WT) primary cultured neurons of C57 mice. The WT neurons were infected with lentivirus containing *EFS*-hfCas13x.1/*U6*-cr8 (WT + cr8), *EFS*-hfCas13x.1/*U6*-cr9 (WT + cr9), or the non-targeted control crRNA *EFS*-hfCas13x.1/*U6*-NT (WT + NT) ($n = 3$ for all groups). (D and E) Western blot analysis (D) and quantification of band density (E) of UBE3A protein expression in cultured primary neurons of WT or AS mice infected with lentivirus-containing *hSyn1*-hfCas13x.1/*U6*-NT or *hSyn1*-hfCas13x.1/*U6*-cr9 ($n = 3$ for all groups). (F) RT-qPCR analysis of *Ube3a* mRNA levels in cultured primary neurons of WT or AS mice infected with *hSyn1*-hfCas13x.1/*U6*-NT or *hSyn1*-hfCas13x.1/*U6*-cr9 ($n = 3$ for all groups). (G) Expression of the indicated genes in cultured primary neurons of AS mice infected with *hSyn1*-hfCas13x.1/*U6*-cr9 to silence the paternal lncRNA *Ube3a-ATS* ($n = 3$ for all groups). Levels were relative to that in primary neurons of AS mice infected with the *hSyn1*-hfCas13x.1/*U6*-NT non-targeted control. Statistical significance was assessed by one-way ANOVA followed by Tukey's multiple comparison test. * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$. ns, not significant.

introducing permanent genomic alterations. To date, Cas13x.1 is the smallest known crRNA-guided RNA endonuclease, with only 775 amino acids, and a high-fidelity variant of this Cas, hfCas13x.1, has been shown to exhibit high on-target activity with markedly lower collateral activity. Here, we showed that treatment with hfCas13x.1 guided by crRNA targeting the intergenic region between *Ube3a* and *Snord115*, expressed under the neuron-specific *hSyn1* promoter and packaged in an AAV-PHP.eb vector effectively induced the

expression of paternal *Ube3a*, resulting in significant improvements to behavioral impairments in AS model mice.

RESULTS

A high-fidelity Cas13x.1 reinstates paternal *Ube3a* expression in primary cultured mouse neurons

To detect whether Cas13x.1 could inhibit the transcription of *Ube3a-ATS* lncRNA, we designed none crRNAs targeting premature

Ube3a-ATS transcripts in mice, with target sites between the *Snord115* cluster and the *Ube3a* locus (Figure 1A and Table S1), and then constructed *elongation factor 1 alpha short* promoter (*EFS*)-Nls-Cas13x.1-Nls-Flag-T2A-EGFP and each crRNA individually into all-in-one plasmid (Figure S1A). We transduced N2a neuroblastoma cells with the constructs above, and then sorted the EGFP⁺ cells by flow cytometry for quantification of RNA level of *Ube3a*-ATS regions downstream of *Snord115* genes by reverse transcriptase quantitative PCR (RT-qPCR). The results indicated that cr8 and cr9 provided the highest knockdown efficiency (i.e., >70%), compared with a non-targeting (NT) control (Figure 1B and Table S1). To evaluate the knockdown efficiency of *Ube3a* expression in primary cultured neurons, we isolated primary cortical neurons from C57 mouse embryos and infected these cells with lentivirus expressing Cas13x.1 and crRNA (Figure S1A). Quantification of *Ube3a*-ATS RNA and *Ube3a* mRNA in infected primary neurons showed that Cas13x.1/cr8 and Cas13x.1/cr9 induced significant up-regulation of *Ube3a* transcription concurrent with *Ube3a*-ATS knockdown (Figure 1C). Previous study showed that Cas13x.1 could well tolerate a single-nucleotide mismatch.²⁰ We predicted that cr9 could target multiple target sites with one or two base pair mismatches in the intergenic *Snord115*-flanking sequence (Table S1).

We also isolated and infected AS (*Ube3a*^{m-/p+}) primary cortical neurons with lentivirus expressing Cas13x.1-Flag to investigate whether neuron-specific promoter, human *Syn1* (*hSyn1*) (Figures S1D and S1E), induced high efficiency of *Ube3a*-ATS silencing compared with the *EFS* (Figures S1B and S1C) promoter under the same conditions. Western blot results showed that the *hSyn1*-driven Cas13x.1 with cr8 reinstated UBE3A protein to 52.5% compared with the expression level of maternal UBE3A in WT cells, while *EFS*-driven Cas13x.1 reinstated UBE3A protein to 34.8% (Figures S1B and S1C). This result indicated that *hSyn1*-Cas13x.1 with crRNA could restore UBE3A expression more effectively in AS primary neurons. Since high-fidelity Cas13x.1 (hfCas13x.1) has markedly lower collateral cleavage activity compared with wild-type (WT) Cas13x.1,²¹ we examined UBE3A protein (Figures 1D, quantified in 1E) under *Ube3a*-ATS silencing by WT Cas13x.1 and hfCas13x.1 in primary AS mouse neurons. The results showed that hfCas13x.1 with cr9 could restore UBE3A protein expression through the suppression of *Ube3a*-ATS as effectively as WT Cas13x.1 with cr9 (Figures 1D and 1E). The *Ube3a* mRNA is also significantly reinstated by hfCas13x.1 (Figure 1F). Additionally, we found that the cr8–9 tandem crRNA with hfCas13x.1 could not restore UBE3A as efficiently as cr9 alone, indicating that the addition of cr8 did not positively affect hfCas13x.1 function (Figures 1D and 1E). To evaluate the localization of unsilenced paternal UBE3A, we also isolated and infected AS (*Ube3a*^{m-/p+}) primary cortical neurons with lentivirus expressing Cas13x.1-Flag. We co-stained UBE3A and Flag (hfCas13x.1-Flag) and found that the UBE3A expression restored by hfCas13x.1/cr9 was enriched in the nucleus of AS mouse primary neurons (Figure S1F).

Snrpn, *Snord116*, *Ipw*, and *Snord115* are all processed from the same *Shhg14* transcript. To detect the effects on these genes by hfCas13x.1/cr9, we evaluated the expression of *Snrpn*, *Snord116*, *Ipw*, and

Snord115 in AS neurons infected with lentivirus expressing hfCas13x.1 and cr9 by RT-qPCR, which indicated that the expression of *Ipw* and *Snord115*, relatively close to the crRNA target sites, were affected by the cr9, while *Snrpn* and *Snord116* were not (Figure 1G). Whole transcriptome RNA sequencing (RNA-seq) of infected AS mouse primary neurons indicated that relatively few genes were differentially expressed between the hfCas13x.1/U6-cr9 treatment group and neurons infected with the NT crRNA (Figure S1G and Table S2). We also checked the expression of predicted potential off-target sites and found no significantly decreased expression of these transcripts in RNA-seq data (Table S3), which suggested high targeting specificity for hfCas13x.1/cr9. Collectively, these data indicated that *hSyn1*-driven hfCas13x.1 coordinated by U6-driven cr9 crRNA effectively and specifically unsilences *Ube3a* in neurons.

AAV-packaged hfCas13x.1 guided by cr9 unsilenced paternal *Ube3a* expression in mouse cortex and hippocampus *in vivo*

Previous studies have shown that restoring *Ube3a* expression during early developmental stages could rescue most AS-associated phenotypes in AS model mice,²² while the AAV-PHP.eb capsid has been shown to efficiently cross the blood-brain barrier to infect cells in the central nervous system.²³ Based on these findings, we individually performed intracerebroventricular (I.C.V.) and intravenous (I.V.) injections with AAV-PHP.eb containing a CAG promoter-driven tdTomato reporter in neonatal mice. One facial vein was administered 1.9×10^{11} virus particles, or 0.3×10^{11} virus particles were injected into the bilateral ventricle of the neonatal mouse. When examined histologically at 3 weeks, animals injected by I.V. infusion showed remarkably high tdTomato signal in liver tissue, while I.C.V. injection with fewer virus particles led to obvious tdTomato expression in the central nervous system (Figure 2A).

In light of these results, we then packaged AAV-PHP.eb with *hSyn1*-hfCas13x.1 and U6 promoter-NT or cr9 (Figure S1A). To further determine whether hfCas13x.1/cr9 could unsilence paternal *Ube3a*, we also made *Ube3a*^{m+/pYFP} knock-in mice, which harbors a paternal *Ube3a*-YFP knock-in allele, and primary neurons from *Ube3a*^{m+/pYFP} embryos were isolated and infected with the packaged AAV with hfCas13x.1/NT or hfCas13x.1/cr9. RT-qPCR-based detection of YFP showed that AAV-packaged *hSyn1*-hfCas13x.1/U6-cr9 significantly induced the expression of paternal *Ube3a*-YFP in neurons (Figure S2A). To evaluate the efficiency of *Ube3a*-ATS silencing in brain, we co-injected 2.5×10^9 AAV-PHP.eb-CAG-tdTomato with a total of 5×10^{10} AAV-PHP.eb-*hSyn1*-hfCas13x.1/U6-cr9 or AAV-PHP.eb-*hSyn1*-hfCas13x.1/U6-NT particles I.C.V. into postnatal day 1 (P1) AS or WT mice (Figure 2B). We dissected the tdTomato⁺ cortex, hippocampus, spinal cord, or cerebellum tissues for quantification of *Ube3a*-ATS and *Ube3a* expression at 4 weeks. RT-qPCR results showed that treatment with *hSyn1*-hfCas13x.1/U6-cr9 led to a 77.5% knockdown of *Ube3a*-ATS transcripts in the cortex tissues (Figure 2C) and 76.6% knockdown in hippocampus tissues (Figure 2D) of AS mice compared with that in AS control mice administered with *hSyn1*-hfCas13x.1/U6-NT. Furthermore, paternal *Ube3a* mRNA levels increased to 44.3% in cortex tissues and 17.6% in

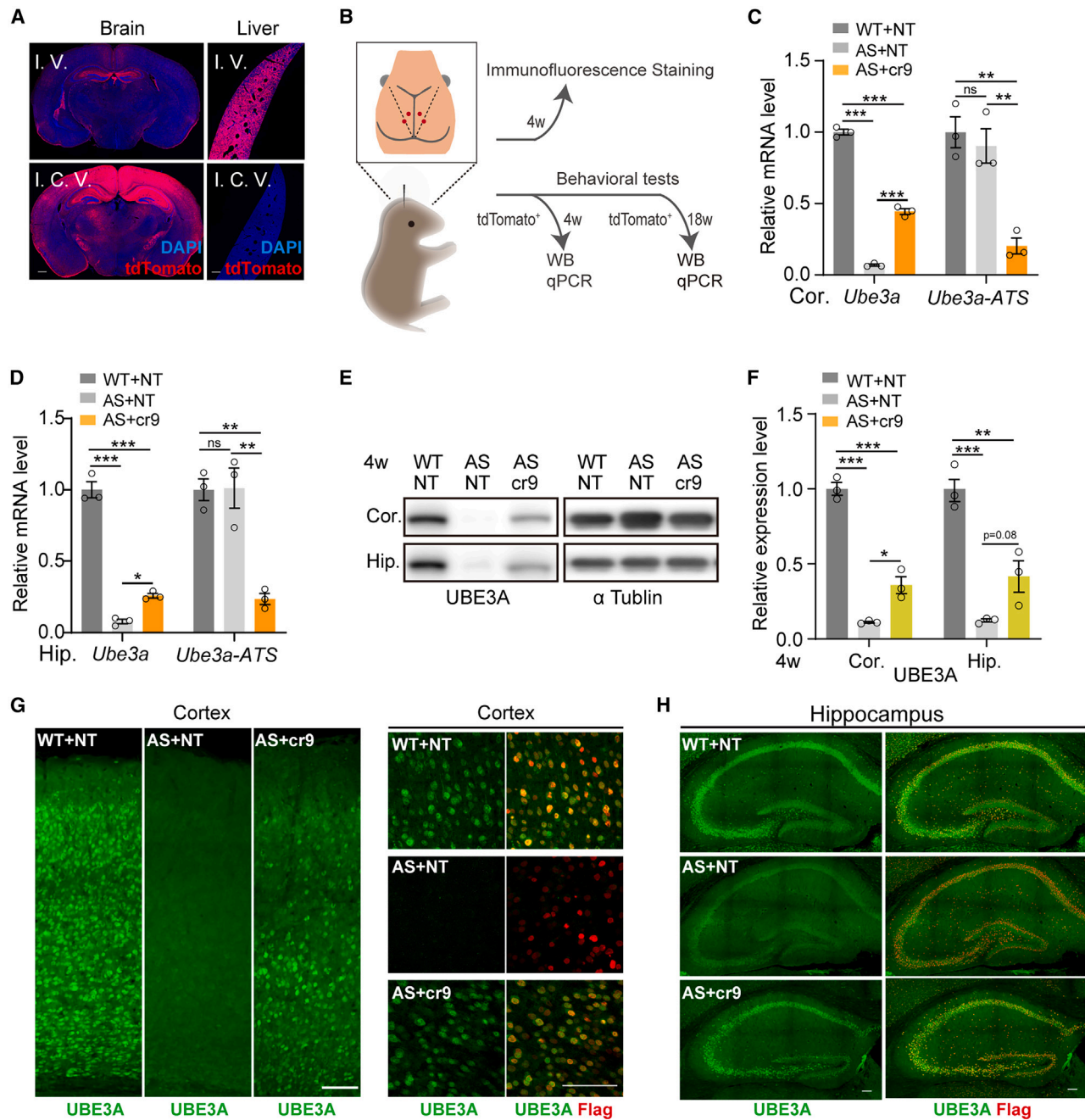


Figure 2. AAV delivery of the CRISPR-hfCas13x system restores paternal *Ube3a* expression in AS mouse brain

(A) TdTomato signal in brain and liver of WT mouse at 3 weeks after I.C.V. injection of 1.9×10^{11} or I.V. injection of 0.3×10^{11} AAV-PHP.eb particles containing CAG-tdTomato at P0. Scale bar, 500 μ m. (B) Experimental timeline for evaluation in WT and AS (*Ube3a*^{m-/p+}) mice after I.C.V. injection of AAV-PHP.eb containing *hSyn1*-hfCas13x.1/*U6*-NT or *hSyn1*-hfCas13x.1/*U6*-cr9 at P0. (Cand D) RT-qPCR analysis of mRNA expression of *Ube3a* and *Ube3a-ATS* in the cerebral cortex (Cor.) (C) and hippocampus (Hip.) (D) of WT and AS mice with indicated treatment at 4 weeks (n = 3 for all groups). (E and F) Western blot (E) and quantification (F) of protein expression in the cerebral cortex (cor.) and hippocampus (hip.) of WT and AS mice with indicated treatment at 4 weeks (n = 3 for all groups). (G and H) Immunofluorescence staining for UBE3A with FLAG in cortex (G) or hippocampus (H) of WT or AS mice with indicated treatment at 4 weeks. Scale bar, 100 μ m. Statistical significance was assessed by one-way ANOVA followed by Tukey's multiple comparison test. *p < 0.05; **p < 0.01; ***p < 0.001. ns, not significant.

hippocampus tissues of AS mice compared with expression levels of maternal *Ube3a* in C57 WT control mice (Figures 2C and 2D).

Western blots showed that paternal UBE3A protein expression reached 35.9% in cortex tissues and 41.6% in hippocampus tissues of AS mice at 4 weeks after infection with AAV-PHP.eb-*hSyn1*-hfCas13x.1/U6-cr9 (Figures 2E and 2F), compared with that in WT mice. By contrast, no paternal UBE3A protein expression could be detected in spinal cord or cerebellum samples of any mice at 4 weeks (Figures S2B and S2C). Paternal UBE3A protein expression maintained 21.1% and 39.1% expression in cortex and hippocampus, respectively, at 18 weeks; however, the difference in statistical analysis was not significant (Figures S2D and S2E). RT-qPCR-based detection of *Snord115* target gene transcripts, including *Crhr1*, *Ralgps*, *Taf1*, *Pbrm1*, and *Dpm2*, indicated no significant changes in the expression of any of these genes in cortical tissues of AS mice, despite partial inhibition of *Snord115* expression by the AAV-PHP.eb-*hSyn1*-hfCas13x.1/U6-cr9 system (Figure S2F). Moreover, the detection of hfCas13x.1 transcription driven by the *hSyn1* promoter in the cortex and liver of AS mice indicated that it was exclusively expressed in the brain (Figure S2G). When examined histologically at 4 weeks, mice showed uneven expression of hfCas13x.1-Flag in cortex and hippocampus cells (Figure S3A). AS mice treated with AAV-PHP.eb-*hSyn1*-hfCas13x.1/U6-cr9 mice had a significant expression of unsilenced UBE3A in the nucleus of cortex and hippocampus (Figures 2G, 2H, and S3B), and we also found the cells with strong Flag staining showed higher expression of UBE3A (Figure S3B). Co-staining of UBE3A and NeuN in treated mice indicated that UBE3A was specifically recovered in neurons (Figure S4). Moreover, we observed no obvious structural change in brain slices and no abnormal NeuN signal in immunostaining images (Figures 2G, S3, and S4). Collectively, these results indicated that hfCas13x.1/cr9 was delivered by the AAV-PHP.eb vector and could successfully suppress *Ube3a*-ATS transcripts to unsilence paternal *Ube3a* in AS mice *in vivo*.

CRISPR-hfCas13x.1 in the AAV-PHP.eb vector improves impaired motor function in AS mice

Previous reports have shown that AS mouse models exhibit deficiencies in motor coordination, obesity, hypoactivity, and memory deficit.^{24–27} To evaluate the phenotypic improvement in treated AS mice, we then conducted several behavioral tests for phenotypic analysis of WT or AS model mice I.C.V. with AAV-PHP.eb-*hSyn1*-hfCas13x.1/U6-cr9 or AAV-PHP.eb-*hSyn1*-hfCas13x.1/U6-NT control (Figures S5A, S5C–S5J). We found that WT mice injected with hfCas13x.1/U6-NT control exhibited a sex effect in body weight (Figure S5B), which was also observed in a previous investigation.¹³ In hindlimb clasping and beam walking tests, female AS mice with hfCas13x.1/NT showed more pronounced deficits when compared with sex-matched WT mice with hfCas13x.1/NT (Figures S5D, S5H, and S5I). In an open field test, the distance traveled showed no significant difference between WT and AS mice injected with hfCas13x.1/NT (Figure S5E). Moreover, a significant difference was observed between female and male AS mice with hfCas13x.1/NT in a rotarod test (Figure S5J). Hence, body weight and behavior results for each gender were analyzed separately.

We found that obesity in female AS mice administrated with hfCas13x.1/cr9 was significantly improved (Figure 3A), but not in male AS mice (Figure 3B). Furthermore, female AS mice treated with hfCas13x.1/cr9 also showed improvement in the dowel test (Figure 3C), beam walking test (Figures 3D and 3E), and rotarod test (Figure 3F), while male mice showed improvement in the hindlimb clasping test (Figure 3G) and dowel test (Figure 3H), indicating that motor coordination and balance were improved in AS mice treated with hfCas13x.1/cr9 compared with NT-treated AS control mice.

However, female AS mice injected with hfCas13x.1/cr9 showed no improvement in the hindlimb clasping test (Figure S6A), and male AS mice injected with hfCas13x.1/cr9 showed slight improvement in the rotarod test (Figure S6B). Female AS mice injected with hfCas13x.1/cr9 showed the trend of entering the center more times than AS + NT control mice (Figure S6C), but not the male AS mice (Figure S6D). Also, AS mice treated with the hfCas13x.1/cr9 vector showed no improvement in the marble burying test (Figures S6E and S6F), and the microcephaly phenotype was also not rescued (Figures S6G and S6H). In our hands, AS mice showed no deficits in contextual-based and cue-based learning during fear conditioning assays (Figures S6I–S6K), consistent with some reported results^{13,28}; therefore, this assay was not performed in hfCas13x.1/cr9-treated mice. We also compared the body weight between non-injected WT and WT with hfCas13x.1/NT group, and no difference in body weight was observed between the two groups (Figures S6L and S6M), indicating that AAV-packaged hfCas13x.1 was well tolerated in mice. In summary, these results demonstrate that the unsilencing of paternal *Ube3a* at the P1 stage by AAV-PHP.eb-*hSyn1*-hfCas13x.1/U6-cr9 significantly improved several AS-related behavioral impairments in mice.

DISCUSSION

In this study, we examined different combinations of tissue-specific promoters, modes of administration, Cas endonucleases, and crRNAs to explore the specificity, delivery efficiency, and therapeutic effects of silencing paternal *Ube3a*-ATS *in vivo*. We proved that an RNA-guided CRISPR-hfCas13x.1 system can effectively recover the expression of UBE3A via knockdown of the lncRNA *Ube3a*-ATS *in vitro* and *in vivo*. An increasing number of AAV-based gene therapies have been approved by the US Food and Drug Administration to treat inherited diseases.^{29,30} RNA-guided endonucleases have facilitated the development of precisely targeted gene therapies, and numerous clinical trials exploring the safety and efficacy of CRISPR/Cas-based strategies are currently in progress.³¹ CRISPR-hfCas13x.1 system will be a promising and safe treatment strategy for AS.

Other studies have reported a CRISPR-SaCas9 system that could unsilence paternal *Ube3a* by introducing indels, through AAV vector integration, or by inducing the formation of secondary or tertiary genomic structures at the *Ube3a*-ATS locus.^{13,14} However, the end fragments generated by DSBs can induce chromosomal translocations, rearrangements, or large fragment deletions.^{15,32,33} CRISPR-Cas13 enzymes do not contain the RuvC and HNH domains responsible for DNA cleavage, which avoid genomic alterations and

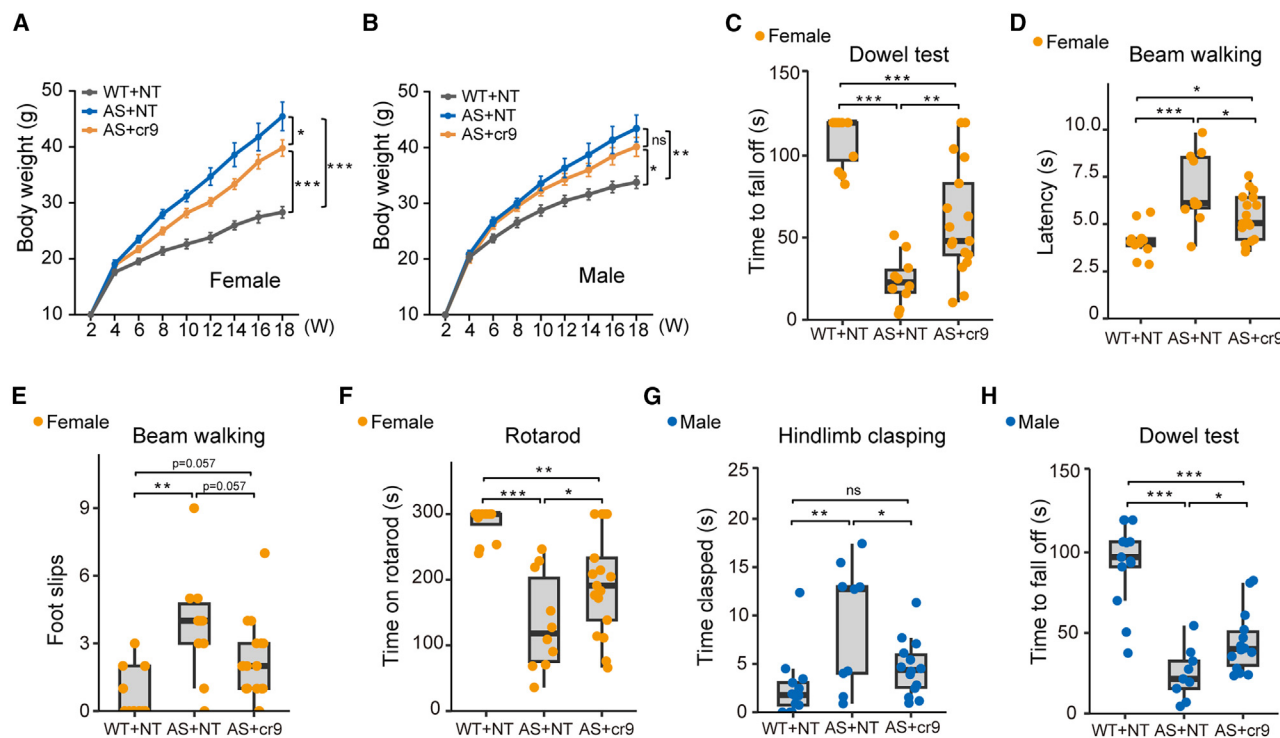


Figure 3. HfCas13x.1/cr9 improves motor functions in an AS mouse model

(A) Body weight of female mice was measured biweekly over 18 weeks. (B) Body weight of male mice was measured biweekly over 18 weeks. (C) Dowel test data in 13-week-old female mice. (D and E) Beam walking assays in 14-week-old female mice. Time to traverse the beam (D) and the number of foot slips (E) are shown. (F) Accelerating rotarod test data in 15-week-old female mice. (G) Hindlimb clasping assays in 7-week-old male mice. (H) Dowel test data in 13-week-old male mice. N = 12 for WT + NT; n = 10 for AS + NT; n = 17 for AS + cr9 in female mice. N = 13 for WT + NT; n = 9 for AS + NT; n = 14 for AS + cr9 in male mice. Statistical significance was assessed by one-way ANOVA followed by the Holm-Sidak comparison test. *p < 0.05; **p < 0.01; ***p < 0.001. ns, not significant.

off-targets.¹⁷ Therefore, using a high-specificity Cas13x system packaged in AAV may represent a safe and potential long-lasting therapeutic option for targeted treatment of AS.

However, we found that the expression of hfCas13x.1-Flag varies in each neuron cell and the UBE3A was significantly reinstated in cells with high expression of hfCas13x.1-Flag (Figure S3), indicating the uneven spreading of the AAV vector in the brain. This could explain that the recovered protein levels were not high enough in cortex and hippocampus, and AAV-PHP.eb-hSyn1-hfCas13x.1/U6-cr9 treatment partially improved motor functions of AS mouse model. By comparing male and female mice injected with NT control, we found that WT or AS mice showed sex effects in several behavioral tests (Figures S5B–S5K), which might explain gender-dependent improvements in the obesity phenotype and motor functions in AS mice treated with hfCas13x.1-Flag/cr9 (Figures 3A–3H and S6A–S6H). It is reasonable to believe that the further optimization of AAV vector and drug component will improve the therapeutic effect in disease model and promote translational research of this approach.

Snhg14, as a single long RNA, is co-transcriptionally spliced into several host genes that were further processed into *Snrpn* mRNA, *Snord116*,

Ipw, *Snord115*, and *Ube3a-ATS*. In this study, we found that hfCas13x.1/cr9 also decreased the expression of *Snord115* and *Ipw* in primary cells. As mentioned above, cr9 has multiple target sites with one or two mismatches (Table S1) between *Snord115* clusters, and these target sites are also close to the *Ipw* location in mice. Therefore, *Snord115* and *Ipw* might have no enough time to be spliced from the nascent RNA when hfCas13x.1/crRNA bound and degraded the nascent RNAs. This crRNA-specific effect could be solved by switching crRNA.

Adolescent reactivation in *Ube3a*^{Stop/p+;Cre^{ERT+}} mice only partially rescued motor coordination, while no improvement was seen in the adult reactivation group.²² This indicated earlier reinstatement of *Ube3a* is critical for AS treatment. However, humans have a very long childhood compared with mice, and until now the precise treatment window in AS patients remains uncertain.²² Therefore, whether patients of different ages can benefit from gene therapy needs to be thoroughly studied. Although the bilateral I.C.V. route is commonly used in preclinical and clinical studies for CNS diseases and the patient ages ranged from 1 day to 84 years,^{34,35} the same AAV serotype usually displayed variable delivery efficacy and safety among mice and primates; hence, preclinical studies in non-human primates with the AAV vector need to be conducted in the future.

Overall, we demonstrated that the hfCas13x.1-crRNA system effectively and specifically relieves *Ube3a* silencing by degrading *Ube3a*-ATS in primary neurons and *in vivo*. Moreover our data showed that one-time injection of AAV with hfCas13x.1-crRNA significantly improved several motor functions in AS mice, indicating its potential for treatment of AS.

MATERIALS AND METHODS

Plasmid construction

The coding sequences of Cas13x.1 and crRNAs were constructed into different backbones with the NEBuilder HiFi DNA assembly Master Mix (New England BioLabs), as previously described.²⁰ All crRNA sequences are provided in Table S1. The plasmid diagram is shown in Figure S1A.

N2a culture, transient transfection, and cell sorting

The N2a cell line was purchased from the cell bank of the Shanghai Institute of Biochemistry and Cell Biology, Chinese Academy of Sciences, and cultured in an incubator at 37°C with 5% CO₂. The medium of N2a is DMEM (Gibco) supplemented with 10% fetal bovine serum (FBS) (Gibco), 1% penicillin/streptomycin (Gibco), and 1× non-essential amino acids (Gibco). Cells were transfected with plasmids and PEI (Polyscience) in a 1:2 ratio 12 h later after being plated. Two days after transfection, cells were digested with 0.05% trypsin (Thermo Fisher Scientific) and sorted by MoFlo XDP. GFP-positive cells are kept in DMEM and then in Trizol for RNA isolation.

Animals

All animal experiments were approved by the Animal Care and Use Committee of the Institute of Neuroscience, Chinese Academy of Sciences, Shanghai, China. *Ube3a*^{m-/p+} mice and *Ube3a*^{m+/pYFP} mice were maintained on the C57BL/6 background. All mice used in this study were housed in 12 h light/dark cycle. Mice were randomly assigned to the corresponding treatments. All experimental protocols were approved by the Animal Care and Use Committee of the Institute of Neuroscience, Chinese Academy of Sciences, Shanghai, China.

AS mouse model and *Ube3a*^{YFP} mice

Ube3a KO mice were generated by Jiang et al.³⁶ *Ube3a*^{YFP} mice were generated as previously described.³⁷ AS mouse models were generated by crossing the *Ube3a*^{m+/p-} heterogeneous females among *Ube3a* KO mice with C57BL/6 WT males from Shanghai SLAC Laboratory Animal Co., Ltd. *Ube3a*^{m+/pYFP} were generated by crossing the *Ube3a*^{YFP} heterogeneous male mice with C57BL/6 WT females. *Ube3a*^{matYFP/p+} were generated by crossing the *Ube3a*^{YFP} heterogeneous female mice with C57BL/6 WT males.

Recombinant lentivirus particles preparation

Lentiviral transgene plasmids were co-transfected with package plasmids and envelope plasmids into HEK293 cells to produce recombinant lentiviral particles delivering the hfCas13x.1 system. The HEK293 cell supernatant was collected for ultracentrifugation

(27,000 rpm, CP90NX, Hitachi). The DNA genome titer of lentiviral vector genomes was determined by detecting lentivirus vectors' copy numbers in cells with RT-qPCR.

Neuronal culture and lentivirus infection

The cortex and hippocampus were dissected from E14–E16 embryos, dissociated in digestion buffer with papain (Worthington Bio Corp), as described previously.³⁸ The digestion buffer contains (in mM): 161 NaCl, 5.0 KCl, 2.9 CaCl₂, 5.0 HEPES, 5.5 glucose, 0.53 MgSO₄, and 0.0056 phenol red, pH 7.4, with additional (in mM) 1.7 cysteine, 1 CaCl₂, and 0.5 EDTA. The tissues were digested for 30 min at 37°C, then plated on PDL (Sigma-Aldrich) in neuronal medium (Thermo Fisher Scientific) containing 2 mM Glutamax-I (Thermo Fisher Scientific), B27 (Thermo Fisher Scientific), and 5% FBS. Four hours later, the neurons were maintained in the same medium without FBS. On days *in vitro* 2, the virus was added to well for 24 h, and then half-changing the medium was carried out for three consecutive days. The subsequent experiments were performed 7–9 days later, after virus infection.

Quantitative PCR and RNA-seq analysis

RNA was extracted from cells or dissected tissues using TRIZOL (Invitrogen) and converted to cDNA using HiScript Q RT SuperMix for qPCR (Vazyme) according to the manufacturer's instructions. We performed qPCR on a Roche 480 II-A using AceQ Universal SYBR qPCR Master Mix (Vazyme). The primers used in qPCR are shown in Table S5. For transcriptome-wide RNA-seq, RNA was extracted and then reverse transcribed to cDNA, which was used for transcriptome-wide RNA-seq. An average of the three repeats is presented. Raw data from the sequencer were first undergone quality control using FastQC. Next, reads are cleaned using trim galore, where the Illumina adapter sequence and low quality (Phred score <20) at the 3' end were trimmed, and the paired reads were removed if any of the two reads did not meet the minimum length (20 bp). Salmon (v1.5) was then used to quantify the expression of transcripts. Differential analysis between the groups was done by a count-based method limma, which is implemented in R and voom is involved for normalization.³⁴ Significantly expressed genes were screened by BH-adjusted p value (<0.05) and the fold-change (>2).

Western blotting

Cells or dissected brain regions were homogenized in SDS lysis buffer (Beyotime) containing a 1× Protease inhibitor cocktail (Beyotime). We loaded 40–60 μg total proteins and separated by SDS-PAGE (Epizyme) and transferred to polyvinylidene fluoride membrane (Merk Millipore). The membrane was blocked in 5% skim milk powder in Tris-buffered saline with Tween buffer (Epizyme). The following antibodies were diluted and incubated overnight: anti-UBE3A (1:1,000; A300-352A, Bethyl), anti-Flag (1:3,000; F1804, Sigma), anti-α-Tubulin (1:5,000; AF0001, Beyotime). After primary antibody incubation, membranes were probed with goat anti-mouse HRP (1:5,000; A9044, Sigma-Aldrich) or goat anti-rabbit HRP (1:5,000; A0545, Sigma), and images were collected using AmershamImager600 (GE) and analyzed by ImageJ software.

AAV virus production

AAV-PHP.eb serotype was used in this study.^{23,35} The CAG-tdTomato, *hSyn1*-hfCas13x.1/*U6*-cr9, or hfCas13x.1/*U6*-NT plasmids were constructed and sequenced before packaging into AAV-PHP.eb vehicle; and the AAV vectors were packaged by PackGene Biotech. The vector titer was determined by qPCR specific for the inverted terminal repeat.

I.C.V. injection and I.V. injection

For I.C.V. administration, neonatal mice were placed on ice for hypothermia anesthesia and injected with AAV virus into bilateral ventricles at four sites.^{39,40} A total of 5×10^{10} AAV (containing *hSyn1*-hfCas13x.1/*U6*-cr9 or *hSyn1*-hfCas13x.1/*U6*-NT) particles plus 2.5×10^9 AAV (CAG-tdTomato) in PBS was injected into each mouse within 24 h of birth using Nanoject III (Warner Instruments). For immunofluorescence experiments, no AAV-containing CAG-tdTomato was injected. TdTomato⁺ cortex or hippocampus tissues were dissected for RT-qPCR and western blotting. For I.V. injection, AAV solution was infused into the facial vein of neonatal mice as previously described.³⁹

Immunofluorescence

Primary neurons were fixed with 4% paraformaldehyde (PFA) for 1 h. Mice were perfused with PBS and then 4% PFA; brain tissues were removed for further fixation overnight and then dehydrated in 30% sucrose for 3 consecutive days. Tissues were sectioned to a thickness of 40 μ m with Thermo HM525-NX and were performed antigen retrieval at 70°C in sodium citrate solution for an hour. Antibodies were diluted in primary antibody dilution buffer (Epizyme) and incubated as following ratio: anti-Ube3a (1:500; A300-352A, Bethyl), anti-Flag (1:500; F1084, Sigma-Aldrich), anti-NeuN (1:2,000, MAB377, Chemicon) overnight at 4°C, and then incubated with Alexa Fluor 488 or Alexa Fluor 647 goat secondary antibody to rabbit IgG and Alexa Fluor 568 goat antibody to mouse IgG (1:500, Invitrogen) for 2 h at 4°C. Slides were sealed with fluoromount-G mounting medium. All images were visualized under a Nikon C2.

Behavioral tests

All behavioral experiments were performed blind to genotypes and injection treatment of animals. Mice were placed in the room for 30 min to acclimate to the environment before the test.

Hindlimb clasping

Each mouse was suspended 10 cm above the table for 20 s by being held by the tail. The video was recorded and time was scored offline. Hindlimb clasping time is the total time spent on clasping. Clasping was defined by the behavior of incomplete splay with one or both hind limbs.

Open field

Each mouse was placed on the edge of a 40 cm \times 40 cm open-field box and allowed to explore for 15 min. The center area is 20 cm \times 20 cm square in the center of the arena. The distance traveled is the total travel distance by each mouse in the arena, which was recorded and

analyzed by camera and EthoVision software (Noldus Wageningen). Center frequency means the number of center entries of each mouse for 15 min.

Dowel test

The 1-m-long dowel (diameter of 9 mm) was placed parallel to the ground at a height of 30 cm. Mice were individually placed on dowel and recorded the time on the dowel. The longest experimental record is 120 s.¹⁰

Beam walking test

The 1-m-long dowel (diameter of 16 mm) was placed parallel to the ground at a height of 30 cm. There was a safe platform at one end of the dowel. After 2 days of training, latency was quantified by measuring the time it took for the mouse to walk over the beam, as well as the number of footsteps was counted. The longest experimental record is 60 s.

Accelerating rotarod test

The test was performed on a rotarod system with the rod rotating from 5 rpm to 30 rpm over 5 min. Mice were trained for 2 days. Two trials were performed for each day with more than 1 h inter-trial interval. On the test day, mice were given two trials and recorded the time of remaining on the rod until falling off or making two consecutive turns. The average time of the two trials was calculated.

Marble burying test

Each mouse is placed in a mouse cage containing 20 small (1.5 cm) clean black marbles on top of 4 inches of corn cob bedding, forming 4 rows of 5 columns. After a period of 30 min of exploration, the mouse is removed from the cage and the number of marbles that are buried at least 50% is recorded.

Fear conditioning

On the training day, after 2 min in the conditioning chamber, mice received two pairings of a tone (2,800 Hz, 85 db, 30 s) with a co-terminating foot-shock (0.7 mA, 2 s), after which they remained in the chamber for 1 additional min and then were returned to their home cages. At 24 h after training, mice were returned to the conditioning chamber for 5 min and tested for freezing in response to the training context. The percent of time spent freezing was taken as an index of contextual learning and memory. At 48 h after training, mice were returned to the conditioning chamber with new walls, mice received a tone (2,800 Hz, 85 db, 30 s) without a foot shock; the percent of time spent freezing was taken as an index of cue learning and memory.

Statistical analyses

All results from analyses are presented as the mean \pm standard error of the mean, except for behavioral tests. Two-tailed, unpaired Student's t-test was used for comparisons between two normally distributed groups, and differences were considered significant when the p values was less than 0.05. For data consisting of more than two groups except for behavioral tests, varying in a single factor, one-way

ANOVA and Tukey's multiple comparison tests were used and differences were considered significant when the p value was less than 0.05. Behavioral tests are presented as boxplot whiskers with the middle line showing the median. For behavioral data consisting of more than two groups, varying in a single factor, one-way ANOVA and Holm-Sidak comparison test were used and differences were considered significant when the p values was less than 0.05.

DATA AND CODE AVAILABILITY

All RNA-seq data have been deposited in the NCBI SRA under project accession number PRJNA842160. Source data are provided in this paper.

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.ymthe.2023.02.015>.

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AUTHOR CONTRIBUTIONS

Conceiving the project: C.Z. and H.Y.; designing experiments: J.L., Z.S., and Y.L.; conducting experiments: J.L., Z.S., Y.L., Z.Y., X.L., J.T., R.L., G.G., and F.Y.; bioinformatic analysis: Y.L.; funding acquisition: H.Y. and C.Z.; writing – original draft: J.L. and Z.S.; writing – review and editing: J.L., Z.S., Y.L., H.Y., C.Z., and Z.X.

DECLARATION OF INTERESTS

The authors have no conflicts of interest to declare that are relevant to the content of this article.

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