A cytosine base editor toolkit with varying activity windows and target scopes for versatile gene manipulation in plants

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ABSTRACT

CRISPR/Cas-derived base editing tools empower efficient alteration of genomic cytosines or adenines associated with essential genetic traits in plants and animals. Diversified target sequences and customized editing products call for base editors with distinct features regarding the editing window and target scope. Here we developed a toolkit of plant base editors containing AID10, an engineered human AID cytosine deaminase. When fused to the N-terminus or C-terminus of the conventional Cas9 nickase (nSpCas9), AID10 exhibited a broad or narrow activity window at the protospacer adjacent motif (PAM)-distal and -proximal protospacer, respectively, while AID10 fused to both termini conferred an additive activity window. We further replaced nSpCas9 with orthogonal or PAM-relaxed Cas9 variants to widen target scopes. Moreover, we devised dual base editors with AID10 located adjacent or distally to the adenine deaminase ABE8e, leading to juxtaposed or spaced cytosine and adenine co-editing at the same target sequence in plant cells. Furthermore, we expanded the application of this toolkit in plants for tunable knockdown of protein-coding genes via creating upstream open reading frame and for loss-of-function analysis of non-coding genes, such as microRNA sponges. Collectively, this toolkit increases the functional diversity and versatility of base editors in basic and applied plant research.

INTRODUCTION

Enormous agronomic traits of crops and genetic diseases of humans are associated with genomic single nucleotide polymorphisms (SNPs) (1,2). To ease the investigation and manipulation of these SNPs, the clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR-associated (Cas)-derived cytosine or adenine base editors (CBEs or ABEs) have been invented (3,4), which can efficiently introduce C→T and A→G mutations, respectively, to a genomic target site in both plants and animals, independently of a DNA double-stranded break (DSB) or repair template. Although the so-called prime editing technology has recently been developed to support all types of nucleotide substitutions (5), this approach exhibits an overall low editing capability in monocots and works poorly in dicot species (6–11).

Both CBEs and ABEs are chimeric enzymes containing a Cas protein and a cytosine or adenine deaminase. The prototype of CBE is known as BE3, which is composed of the rat cytosine deaminase APOBEC1 at the N-terminus, the Streptococcus pyogenes Cas9 D10A nickase (nSpCas9) in the middle, and an uracil glycosylase inhibitor (UGI) at the C-terminus (3). The nSpCas9 moiety is responsible for single-guide RNA (sgRNA)-directed DNA targeting, while the APOBEC1 moiety is able to convert a cytosine to uracil within a favorable deamination window of the target sequence (referred to as the activity window). As the recovery of U→G to C→G through the endogenous uracil-excision repair pathway is inhibited by UGI, the U→G mismatch is subsequently corrected by DNA polymerases to T→A. Similarly, BE7.10, a canonical ABE, consists of a heterodimer between the Escherichia coli TadA adenine deaminase and TadA7.10, a laboratory-evolved TadA variant, at the N-terminus and the nSpCas9 at the C-terminus (4). Upon association with a genomic target sequence through the nSpCas9-sgRNA module, TadA7.10 deaminates an adenine within the activity window of the target sequence to inosine, which is subsequently read by DNA polymerases as guanine to induce an A→T to G→C substitution. In both CBE- and ABE-mediated base editing, nicking the non-
deaminted DNA strand by nSpCas9 can facilitate its repair using the deaminated strand as a template, thus promoting the conversion of the target base pair (3,4).

In general, CBEs have been employed more extensively than ABEs in basic and applied biological research (12), especially in plants (13). Notably, in addition to the originally designed use for installing targeted C·G to T·A mutations, CBEs have demonstrated a remarkable versatility in other gene manipulation applications. In the so-called CRISPR-STOP or iSTOP application, CBEs have been repurposed as gene loss-of-function tools by converting an exonic glutamine (CAA/CAG), arginine (CGA), or tryptophan (TGG) codon of a target gene to a stop codon (14,15). CBEs have also been explored to deactivate target genes through destroying intron splice sites to cause mRNA mis-splicing (16–18). These gene loss-of-function strategies have become attractive complements to the CRISPR/Cas technology because multiplex base editing by CBEs is unlikely to induce complicated chromosomal rearrangements, as observed in the CRISPR/Cas9-mediated genome editing (19–21). Moreover, CBEs have been utilized as gene gain-of-function tools to drive directed protein evolution and functional variant screening (22,23). Furthermore, CBEs have been leveraged as molecular recorders of cellular signaling events by translating extracellular stimuli into nucleotide changes at specific genomic loci (24,25).

In CBE-mediated base editing, the requirement for a protospacer adjacent motif (PAM) in the vicinity of a target cytosine and the activity window width are two major factors determining whether the target cytosine is editable or not (26). In addition, some CBEs (e.g. BE3) work inefficiently on target cytosines within unfavorable sequence contexts (3). To address these issues, researchers have combined different Cas proteins with natural or modified cytosine deaminases to build CBEs with distinct target scopes and activity windows. For instance, the lamprey CDA1, human AID (hAID), hAPOBEC3A (hA3A), and engineered hA3B have been used to replace the rat APOBEC1 in BE3 to obtain effective CBEs in plants (27–30). Theoretically, when the base editor arsenal is filled with more CBEs with different characteristics, any single cytosine within a given genome will likely fall into the activity window of at least one CBE (26).

To this end, in this study we developed a toolkit of base editors with varying activity windows and target scopes using an engineered variant of hAID cytosine deaminase (referred to as AID10) (31). Using this toolkit, we explored an unprecedented strategy to effectively down-regulate target gene activity in plants by creating upstream open reading frame (uORF). We also demonstrated that our base editor with a broad activity window was useful for probing the interactions between endogenous microRNA (miRNA) sponges and their cognate miRNAs in plants.

MATERIALS AND METHODS

Plant materials and growth conditions

The Arabidopsis thaliana ecotype Col-0 and rice (Oryza sativa) cultivar Zhonghua 11 (ZH11) plants were used as wild-type plants in this study. The seeds of bak1-4 null mutant (SALK_116202) were obtained from the Arabidopsis Biological Resource Center (ABRC). After stratification at 4°C for 2 days, Arabidopsis plants were germinated on the Jiffy soil (Jiffy Group) in a plant growth room under photoperiods of 12 h light (75 μmol/m²/s) at 23°C and 12 h dark at 21°C. For hypocotyl length comparison, Arabidopsis seeds were sterilized and sown on the 0.5 × Murashige and Skoog (MS) medium containing 1% sucrose and 0.6% agar. After stratification at 4°C for 2 days, plants were germinated in the dark at 23°C for 5 days before the hypocotyl length was measured. Rice plants were grown on the Jiffy soil in a growth chamber under photoperiods of 12 h light (200 μmol/m²/s) at 32°C and 12 h dark at 28°C. For the herbicide resistance assay, rice seeds were sterilized and grown on the 0.5 × MS medium supplemented with 0.1 mg/l imazethapyr herbicide for 9 days.

Plasmid construction

The coding sequences of AID10-XTEN and 6 × Gly-Ser (GGS)-UGI were amplified from the pGAL1-AID10-dCas9-UGI plasmid (31), and those of the nuclear localization signal (NLS) and plant codon-optimized dCas9 (D10A, H840A) were amplified from the HBT-dCas9 plasmid (32). For transient expression in rice protoplasts, the sequences of the maize Ubiquitin-1 (ZmUbi) promoter, NLS, AID10-XTEN, and dCas9 were assembled in order into the Xhol/PstI-digested HBT-FLAG vector (33) to obtain HBT-AID10-dCas9 by recombination-based cloning using the CloneExpress MultiS One Step Cloning Kit (Vazyme, China). The 6 × GGS-UGI fragment was further inserted using the CloneExpress II One Step Cloning Kit (Vazyme, China) to generate HBT-AID10-dCas9-UGI. The H840A mutation in the dCas9 coding sequence was reversed by PCR-based site-directed mutagenesis to obtain HBT-AID10-ncas9 and HBT-AID10-ncas9-UGI. The coding sequence of APOBEC1-ncas9-UGI was amplified from the pmcas9-PBE plasmid (34) and was used to replace AID10-ncas9-UGI in the HBT-AID10-ncas9-UGI plasmid through restriction digestion and ligation. The coding sequence of AtCDA1 (At2g19570) was PCR amplified from Arabidopsis cDNA and was used to replace APOBEC1 to obtain HBT-AiCDA1-ncas9-UGI by recombination-based cloning. The coding sequences of plant codon-optimized SaCas9 (35) and Lb-Cas12a (36) harboring D10A and D832A mutations, respectively, were generated by overlapping PCR. Resultant PCR products, along with fragments of the ZmUbi promoter, NLS, AID10-XTEN, and 6 × GGS-UGI, were assembled into the Xhol/PstI-digested HBT-FLAG vector to generate HBT-AID10-nsaCas9-UGI and HBT-AID10-dLbCas12a-UGI, respectively. Seven or three point mutations were introduced into nSpCas9 and nSaCas9, respectively, to obtain the HBT-AID10-nSpCas9-NG-UGI and HBT-AID10-nSaCas9-KKH-UGI plasmids encoding Cas9 derivatives with relaxed PAM requirement (37,38). For transient expression in Arabidopsis protoplasts, the ZmUbi promoter in AID10-encoding plasmids was replaced by the Arabidopsis Ubiquitin 10 (AtUBQ10) promoter through SpeI and AvrII sites. To change the fusion orientation of AID10 to nCas9, coding sequences for AtUBQ10, NLS, nCas9, XTEN, AID10, and UGI were assembled in order into the Xhol/PstI-digested
HBT-FLAG vector to obtain HBT-nCas9-AID10-UGI by recombination-based cloning. The DNA fragment of XTEN-AID10-UGI was cut out from the HBT-nCas9-AID10-UGI plasmid by StuI and subsequently inserted into the HBT-AID10-nCas9 plasmid through the StuI site to generate the HBT-AID10-nCas9-AID10-UGI plasmid.

To construct dual base editors, we synthesized the plant codon-optimized coding sequence of adenosine deaminase ABE7.10 and used to replace APOBEC1-nCas9-UGI between the Sacl/Shfl and EcoRI/EcorV sites, respectively. The three sgRNA expression cassettes in tandem were PCR amplified and cloned into the binary plasmid at the HindIII site by recombination-based cloning.

Protoplast isolation and transfection

Protoplasts were isolated from leaves of 4-week-old Arabidopsis or leaf sheaths of 10-day-old rice plants as previously described (43,44). Briefly, plant tissues were cut into 0.5-mm strips and digested in 10 ml enzyme solution containing 1.5% Cellulase R10 (Yakult Honsha, Japan), 0.4% Macerozyme R10 (Yakult Honsha, Japan), 0.4 M mannitol, 20 mM MES, pH 5.7, 20 mM KCl, 10 mM CaCl2, 0.1% BSA for 3 h. After adding 10 ml W5 solution containing 154 mM NaCl, 125 mM CaCl2, 5 mM KCl, 2 mM MES, pH 5.7, the digestion mixture was filtered through a 75-μm cell strainer to remove tissue debris. Isolated protoplasts were collected by 2-min centrifugation at 100 g for Arabidopsis or 150 g for rice, and were resuspended in 10 ml W5 solution and rested on ice for 30 min. After centrifugation for 1 min, cells were resuspended in MMG solution containing 0.4 M mannitol, 15 mM MgCl2, 4 mM MES, pH 5.7, to a concentration of 2 × 105 cells/ml. To evaluate the editing efficiencies of base editors, 40 μg (2 μg/μl) plasmids expressing base editors and 40 μg (2 μg/μl) plasmids expressing sgRNAs were gently mixed with 400 μl protoplasts and 440 μl polyethylene glycol (PEG) solution containing 40% PEG 4000, 0.2 M mannitol, 0.1 M CaCl2 in a 2-ml round-bottom tube. The mixture was incubated at room temperature for 5 min for Arabidopsis or 15 min for rice before transfection was stopped by adding 1.6 ml W5 solution. Transfected protoplasts were collected by 3-min centrifugation at 100 g for Arabidopsis or 200 g for rice. Cells were resuspended in 200 μl W5 solution and transferred to 2 ml WI solution containing 0.5 M mannitol, 20 mM KCl, 4 mM MES, pH 5.7. After incubation at room temperature in the dark for 48 h, protoplasts were harvested by centrifugation at 200 g for 3 min for subsequent analysis.

Evaluation of base editing efficiencies in protoplasts

Genomic DNA was extracted from transfected protoplasts using the Plant Genomic DNA Kit (Tiangen, China) according to the supplier’s instruction. To evaluate base editing efficiencies by restriction fragment length polymorphism (RFLP) assay, target regions were amplified by PCR using the Q5 High-Fidelity DNA Polymerase (New England Biolabs, USA) and purified using the FastPure Gel DNA Extraction Mini Kit (Vazyme, China). Equal amounts of purified amplicons were subjected to restriction digestion with Xhol (New England Biolabs, USA) at 37°C overnight and then were resolved on a 2% agarose gel by electrophoresis. To confirm mutations at the target site, Xhol-resistant PCR amplicons were cloned into the pEASY-Blunt vector (TransGen, China) and 15 clones for each sample were randomly selected for Sanger sequencing.

To evaluate base editing efficiencies by amplicon deep sequencing, genomic fragments of 220–270 bp spanning the
target sites were PCR amplified using primers with a 6-nt unique barcode at the 5′ end, as listed in Supplementary Table S2. The PCR amplification was carried out with the Q5 High-Fidelity DNA Polymerase using 10 ng genomic DNA as template. The PCR cycle number was set as 28 to ensure the reaction was stopped within the linear range of amplification. Purified PCR amplicons using the FastPure Gel DNA Extraction Mini Kit were pooled at equal molar ratios and subjected to deep sequencing. Deep sequencing library construction (including dA-tailing and adaptor ligation to amplicons), sequencing, trimming of adapter sequences, and demultiplexing based on barcodes were fulfilled by the Genewiz company (Suzhou, China). Sequencing was performed on an Illumina HiSeq platform for 150-bp paired-end reads and at least 100,000 reads were obtained for each sample. FASTQ files were analyzed using CRISPRESSO2 (https://crispesso.pinellolab.partners.org) (45) or BE-Analyzer (http://www.rgenome.net/be-analyzer) (46) to quantify mutation efficiencies at individual target sequences.

Generation of transgenic plants

The binary plasmid co-expressing the base editor and sgRNA was introduced into Agrobacterium tumefaciens strain GV3101 for Arabidopsis transformation or strain EHA105 for rice transformation by electroporation. Correct A. tumefaciens cells were used to transform Arabidopsis floral tissues and rice embryogenic calli according to detailed protocols described previously (47,48). Transgenic Arabidopsis T1 plants and rice T0 plants were selected on the medium containing 50 mg/l hygromycin before resistant plants were transferred to the Jiffy soil.

Genotyping of transgenic plants

For each transgenic plant, the crude extract of genomic DNA was obtained by homogenizing a total of 15 mg tissues collected from 3 different leaves in 30 μl lysis buffer containing 100 mM Tris–HCl, pH 9.5, 1 M KCl, 10 mM EDTA and then incubating at 70°C for 30 min. The 10-fold diluted lysate was used as PCR template for genotyping using the Hi-TOM method (49), which utilizes two rounds of PCR and deep sequencing to identify mutations. Briefly, genomic fragments spanning the target sites were PCR amplified using target-specific primers and barcoded primers sequentially. Resultant amplicons were pooled at equal molar ratios and sequenced with an Illumina HiSeq platform by the Novogene company (Beijing, China). Mutations in the transformants were identified by analyzing the deep sequencing data with the Hi-TOM tool (http://www.hi-tom.net/hi-tom) and further confirmed by Sanger sequencing of PCR amplicons of target regions. PCR primers used are listed in Supplementary Table S2.

RNA extraction and qRT-PCR

Leaf tissue of 20 mg from 4-week-old Arabidopsis plants were ground in liquid nitrogen and total RNA was extracted using 1 ml RNAiso Plus reagent (TAKARA, Japan) according to the manufacturer’s instruction. To quantify the transcript levels of Arabidopsis miR166 target genes, the second internodes in rachis branches of 8-week-old Arabidopsis plants were sampled. Reverse transcription (RT) was performed with 1 μg RNA and oligo dT primer using the PrimeScript RT reagent kit with gDNA Eraser (TAKARA, Japan) in a reaction volume of 20 μl. Resultant cDNA products were used as PCR template after 8-fold dilution. Quantitative RT-PCR (qRT-PCR) was performed in a LightCycler 96 Instrument (Roche, USA) using the TB Green Premix Ex Taq (TAKARA, Japan) according to the manufacturer’s instruction. The relative expression levels were quantified using the 2−ΔΔCt method and the Arabidopsis Actin2 gene was used as a reference gene. Primers used for qRT-PCR are listed in Supplementary Table S3.

Immunoblot analysis

Crude protein extracts were obtained from leaf tissues of 4-week-old Arabidopsis plants by boiling homogenized tissues in the SDS-PAGE loading buffer at 95°C for 10 min. Proteins were resolved on a 10% SDS-PAGE gel and transferred to a polyvinylidene fluoride (PVDF) membrane. ABK1 proteins were detected by immunoblotting using the anti-BAK1 antibody, which was raised against the C-terminal peptide DSTSQIENEYPSGPR of AtBAK1 (50). The primary antibody and the horseradish peroxidase (HRP)-conjugated anti-rabbit IgG antibody (Cell Signaling Technology, USA) served as the secondary antibody.

Luciferase reporter assay

To measure the activity of the wild-type or uORF-containing AtBAK1 promoter (1,864 bp upstream of the start codon), 32 μg (2 μg/μl) promoter::LUC reporter plasmid and 8 μg (2 μg/μl) transfection control plasmid (AtUBQ10:GUS (51)) were co-transfected into 200 μl Arabidopsis protoplasts. After incubation at room temperature for 12 h, protoplasts were harvested and homogenized in 100 μl lysis buffer containing 25 mM Tris–HCl, pH 7.8, 2 mM DTT, 2 mM trans-1,2-diaminocyclohexane-N,N′,N′′-tetraacetic acid, 10% (v/v) glycerol, 1% (v/v) Triton X-100. After incubation on ice for 5 min and centrifugation at 12,000 g for 30 sec, 10 μl supernatant was mixed with 100 μl luciferase assay reagent containing 20 μM Tricine, 1.07 mM (MgCO3)4Mg(OH)2, 2.67 mM MgSO4, 0.1 mM EDTA, 33.3 mM DTT, 0.27 mM Coenzyme A, 0.47 mM D-luciferin potassium salt, 0.53 mM ATP, pH 7.8, in a 96-well white microplate. Luminescence was measured in a Varioskan LUX Multimode Microplate Reader (ThermoFisher Scientific, USA). In parallel, another 10 μl protoplast lysate was transferred to a 96-well black microplate with 50 μl MUG solution containing 10 mM Tris–HCl, pH 8.0, 1 mM 4-methylumbelliferyl-β-D-glucuronide (MUG), 2 mM MgCl2. The mixture was incubated at 37°C for 30 min and cooled in the ice-water bath to terminate the reaction. The GUS expression was measured by the Varioskan LUX Multimode Microplate Reader with excitation at 365 nm and emission at 455 nm. The promoter activity was calculated as the ratio between the luciferase readout and the GUS readout for each sample.
RESULTS

AID10-nCas9-UGI confers a broad activity window at the PAM-distal protospacer

To explore CBEs with new properties in plants, we replaced the rat cytosine deaminase APOBEC1 in BE3 (i.e. APOBEC1-nCas9-UGI) with AID10 (Figure 1A and Supplementary Sequence), a newly engineered hAID variant (31). AID10 exhibits low genome-wide guide-independent off-target effects in Drosophila (31) and, as a hAID derivative, may inherently reduce guide-independent RNA off-target editing relative to APOBEC1 and hA3A (51). Unlike hAIDΔΔ, another hAID variant that has been successfully applied to cytosine editing in rice (28), AID10 harbors more mutations, including 9 amino acid substitutions (i.e. K10E, K34E, S38C, T82I, T110A, H130R, V152A, E156G, R174G) and a deletion of the C-terminal 18 residues relative to hAID (Supplementary Figure S1). These modifications have been shown to dramatically enhance the deamination activity of hAID (31). For comparison purpose, we also constructed AID10-nCas9, which lacked the UGI moiety, and AID10-dCas9-UGI and AID10-dCas9, both of which were based on the nuclease-dead Cas9 (dCas9) rather than the nCas9 nickase (Figure 1A). Meanwhile, an Arabidopsis cytosine deaminase, namely AtCDCA1 (52), was also used to replace APOBEC1 in BE3 to obtain AtCDCA1-nCas9-UGI (Figure 1A).

The performance of these CBEs was quickly evaluated using the RFLP assay (53) in rice protoplasts co-expressing individual CBEs and an sgRNA targeting the endogenous OsER1-T1 locus that contains an XhoI restriction site (Figure 1B). Any cytosine change at the protospacer position 5 or 7 (counting the PAM as 21–23) would eliminate the XhoI site and produce an XhoI-resistant allele. As expected, we detected XhoI-resistant target amplicons in protoplasts expressing APOBEC1-nCas9-UGI (Figure 1B), which has previously been shown to enable efficient C-to-T conversions at the positions 3–9 in plant cells (34). Expression of AID10-nCas9 or AID10-nCas9-UGI, but not AID10-dCas9 and AID10-dCas9-UGI, also yielded XhoI-resistant target amplicons (Figure 1B). This observation was reminiscent of a previous finding that APOBEC1-dCas9-UGI is unable to induce C-to-T conversions in rice plants (34), implying that nicking the non-deaminated DNA strand is required for efficient cytosine editing in plant cells. Sanger sequencing of the XhoI-resistant target amplicons further validated C-to-T conversions at the OsER1-T1 locus by AID10-nCas9 and AID10-nCas9-UGI (Supplementary Figure S2). No XhoI-resistant target amplicon was detected in protoplasts expressing AtCDCA1-nCas9-UGI (Figure 1B), suggesting that AtCDCA1 is not suitable for building CBEs. Consistent with this finding, a recent study has shown that the ribose moiety of cytidine is required for substrate binding to AtCDCA1 (54).

Next, we compared cytosine editing patterns and efficiencies between AID10-nCas9-UGI, AID10-nCas9, and APOBEC1-nCas9-UGI at four C-rich genomic target sites (Supplementary Table S1) in rice or Arabidopsis protoplasts by amplicon deep sequencing. We found that AID10-nCas9-UGI induced higher levels of C-to-T conversions than AID10-nCas9, although they exhibited similar activity windows (Figure 1C). Notably, the activity window of AID10-nCas9-UGI spanned from the protospacer positions −1 to 9, which was considerably wider than that of APOBEC1-nCas9-UGI (i.e. positions 4–8) (Figure 1C). Moreover, AID10-nCas9-UGI was able to simultaneously edit more cytosines than APOBEC1-nCas9-UGI at the same target sites (Figure 1D and Supplementary Figure S3), suggesting that AID10-nCas9-UGI has a broader activity window and higher processivity than APOBEC1-nCas9-UGI.

To verify the activity window of AID10-nCas9-UGI in planta, we generated transgenic rice plants expressing AID10-nCas9-UGI or APOBEC1-nCas9-UGI by the ZmUbi promoter and an sgRNA targeting the OsLAZY1 locus. Genotyping of T0 rice transformants identified nucleotide substitutions in 26 out of 40 (65.0%) plants expressing APOBEC1-nCas9-UGI and 24 out of 33 (72.7%) plants expressing AID10-nCas9-UGI (Supplementary Table S4). Both CBEs induced unwanted indels but at different frequencies, namely 42.5% (17 out of 40) by APOBEC1-nCas9-UGI and 21.2% (7 out of 33) by AID10-nCas9-UGI. For simplicity in editing pattern comparison, we only focused on non-chimeric (i.e. homozygous, heterozygous or biallelic) mutant plants that carried genetically identical homogeneous mutations. In mutant plants generated by APOBEC1-nCas9-UGI, the C-to-T conversion only occurred at the protospacer position 7 with a frequency of 31.9% (Figure 1E and Supplementary Figure S4A), while the C-to-G mutation was also detected at the position 7 with a frequency of 13.9%. By contrast, AID10-nCas9-UGI enabled cytosine editing in a broader window, including C-to-T conversions at the positions 2, 7, 13, and 15 with frequencies of 5.6–51.9% and C-to-G conversions at the positions 2 and 7 with frequencies of 1.9% (Figure 1E and Supplementary Figure S4A). Of note, only in mutant plants generated by AID10-nCas9-UGI, mutant alleles (e.g. #20 and #25) carried homogeneous C→G to T→A mutation at the protospacer position 2 (Supplementary Figure S5A). This mutation would disrupt the splicing of the first intron of OsLAZY1 and cause gene inactivation (18). Consistently, these lines displayed increased tiller angles compared to wild-type plants (Supplementary Figure S5B), resembling the previously reported rice lazy1 null alleles (55). We also compared APOBEC1-nCas9-UGI and AID10-nCas9-UGI for editing the OsGL1-I locus in rice plants. Similarly, APOBEC1-nCas9-UGI only edited cytosines at the protospacer positions 7 and 8, whereas AID10-nCas9-UGI allowed cytosine substitutions at the positions −5, −2, 1, 7, and 8 (Figure 1F and Supplementary Figure S4B).

We further compared the activity windows between AID10-nCas9-UGI and APOBEC1-nCas9-UGI by targeting the AtCER10 locus in transgenic Arabidopsis plants, where both CBEs were expressed by the ZmUbi promoter. Targeted cytosine editing was identified in 27 out of 42 (64.3%) T1 plants expressing APOBEC1-nCas9-UGI and 32 out of 34 (94.1%) T1 plants expressing AID10-nCas9-UGI (Supplementary Table S4). Cytosine changes by APOBEC1-nCas9-UGI only occurred at the protospacer position 8, whereas those induced by AID10-nCas9-UGI were detected at the positions 2, 3 and 8 (Figure 1G and Supplementary Figure S6). All mutant plants cre-
Figure 1. AID10-nCas9-UGI exhibits a broader activity window than APOBEC1-nCas9-UGI. (A) Schematic diagram of CBEs with an N-terminally fused cytosine deaminase. UGI, uracil glycosylase inhibitor. NLS, nuclear localization signal. (B) Restriction fragment length polymorphism assay identifies active CBEs in rice protoplasts. The target sequence containing an XhoI site (green) is shown with the PAM in bold. bp, base pair. (C) Comparison of cytosine editing efficiencies between APOBEC1-nCas9-UGI, AID10-nCas9-UGI, and AID10-nCas9 at endogenous genomic loci in rice (left) or Arabidopsis (right) protoplasts by amplicon deep sequencing. Data are shown as mean ± s.d. of two biological replicates performed at different times. (D) Mutant allele composition of the OsNAL1 target site in protoplasts expressing APOBEC1-nCas9-UGI, AID10-nCas9-UGI, or AID10-nCas9. C-to-T conversions are highlighted in bold and the frequencies of individual alleles are shown on the right. Dashed-line box marks the protospacer and the PAM is underlined. The numbers on the top indicate the protospacer positions. Data are collected from one replicate in (C). (E, F) Comparison of cytosine editing efficiencies between APOBEC1-nCas9-UGI (top) and AID10-nCas9-UGI (bottom) at OsLAZY1 (E) and OsGL1-1 (F) target sites in transgenic T0 rice plants. Reverse complements of the protospacers and the PAMs are shadowed in blue and green, respectively. All guanines in the protospacers are in red with positions indicated by subscripted numbers. The percentages of indicated alleles among total alleles are shown and those corresponding to base substitutions are shadowed in yellow. (G) Comparison of cytosine editing efficiencies between APOBEC1-nCas9-UGI and AID10-nCas9-UGI at the AtCER10 target site in transgenic T1 Arabidopsis plants. The reverse complement of the protospacer is underlined and the PAM is in bold. The intron splice site is in lowercase letters.

ated by both CBEs appeared to be chimeric (Supplementary Table S4), reminiscent of our earlier observation that Arabidopsis transgenic T1 plants expressing APOBEC1-nCas9-UGI under the constitutive promoter only produce chimeric mutations (18). Since two earlier studies reported that heterozygous base edited plants can be obtained in Arabidopsis T1 generation when CBEs are expressed by the egg cell-specific promoter (17,56), we subsequently switched to the Arabidopsis ECI.2en-EC1.1p fusion promoter (42) for AID10-nCas9-UGI expression in transgenic Arabidopsis plants. Indeed, we identified 4 out of 157 (2.5%) transgenic T1 plants carrying heterozygous C•G to T•A mutations at the AtCER10 target site, which produced homozygous mutant alleles in the T2 generation (Supplementary Figure S7A and B). In T2 lines #41-16 and #61-3, the cytosine change at the protospacer position 2 by AID10-nCas9-UGI was expected to induce mis-splicing of the first intron of AtCER10, leading to gene deactivation. Consistently, these lines exhibited severe growth defects (Supplementary Figure S7C), resembling the previously reported Arabidop-
Combining AID10 with orthogonal or engineered Cas9 variants expands target scopes

To expand the target scope of AID10-nCas9-UGI, we replaced nSpCas9 with other orthogonal Cas proteins, including the *Staphylococcus aureus* Cas9 nickase (nSaCas9) that recognizes a 3'-NNGRRT PAM (58) and the nuclease-dead *Lachnospiraceaebacterium* Cas12a (dLbCas12a) that recognizes a 5'-TTTV PAM (59) (Figure 2A). To compare the performance of these CBEs in protoplasts by ampiclon deep sequencing, we purposely selected four genomic loci in rice or Arabidopsis including the *AtBIK1* and *AtHDC1* with overlapping target sequences for SpCas9, SaCas9 and LbCas12a (Supplementary Table S1). Both AID10-nSpCas9-UGI and AID10-nSaCas9-UGI induced C-to-T conversions at all target sites, whereas AID10-dLbCas12a-UGI failed to do so (Figure 2B). We reasoned that the failure of AID10-dLbCas12a-UGI was due to its inability to nick the target loci, similar to the case of AID10-dCas9-UGI (Figure 1B). Of note, AID10-nSaCas9-UGI appeared to exhibit a comparable or even higher editing efficiency and wider activity window than AID10-nSpCas9-UGI at these target loci in protoplasts (Figure 2B).

We next compared the performance between AID10-nSpCas9-UGI and AID10-nSaCas9-UGI in transgenic Arabidopsis plants by targeting the *AtHDC1*, *AtABI3* and *At-eTM166-I* loci with overlapping target sequences (Supplementary Table S1). For the *AtHDC1* target site, genotyping of transgenic T1 lines showed that AID10-nSaCas9-UGI induced C-to-T conversions with a similar efficiency to AID10-nSpCas9-UGI, with the former being 5.3% (3 out of 57) and the latter being 8.5% (10 out of 118) (Figure 2C and Supplementary Table S4). C-to-T conversions by AID10-nSpCas9-UGI were induced at the protosparse positions 1, 2, 4, 6, 7, 16, and 17 with frequencies of 0.9–5.1%, while those by AID10-nSaCas9-UGI occurred at the positions 2, 3, 5, 7, 8, 14 and 17 with frequencies of 1.8–3.5% (Figure 2C and Supplementary Figure S8A). Multiple cytosines could be concurrently mutated at the same target loci by both CBEs (Supplementary Figure S8A). Interestingly, AID10-nSaCas9-UGI was able to edit the cytosine at the protosparse position 14, whereas AID10-nSpCas9-UGI was unable to edit the same cytosine (i.e. at the position 13) (Figure 2C). An opposite observation was made for AID10-nSpCas9-UGI at the protosparse position 17 (corresponding to the position 18 for AID10-nSaCas9-UGI) (Figure 2C).

Further comparison between AID10-nSaCas9-UGI and AID10-nSpCas9-UGI for editing the *AtABI3* and *At-eTM166-I* loci revealed contrasting editing frequencies. For the *AtABI3* target site, AID10-nSaCas9-UGI induced C-to-T conversions with a frequency of 17.8% (18 out of 101 T1 lines), whereas only 4 out of 107 (3.7%) T1 lines expressing AID10-nSpCas9-UGI carried C-to-T conversions (Supplementary Figure S8B and Table S4). In addition, the editing frequencies of all five cytosines at the *AtABI3* target site by AID10-nSaCas9-UGI remarkably exceeded those by AID10-nSpCas9-UGI (i.e. 1.0–15.8% versus 0–3.7%) (Figure 2C). For the *At-eTM166-I* site, sequencing analysis of 139 T1 lines expressing AID10-nSaCas9-UGI identified no C-to-T substitution at the target site, whereas AID10-nSpCas9-UGI mediated efficient C-to-T editing with a frequency of 11.3% (8 out of 71 T1 lines) (Figure 2C, Supplementary Figure S8C and Table S4). These results underscore that CBEs containing the same cytosine deaminase but different Cas moieties can exhibit distinct editing efficiencies at the same genomic target site.

We also utilized SpCas9-NG (37) and SaCas9-KKH (38), two engineered Cas9 variants with 3’-NG and 3’-NNNRT PAMs, respectively, to construct AID10-based CBEs. Amplicon deep sequencing of edited targets in Arabidopsis protoplasts showed that AID10-nSaCas9-KKH-UGI enabled efficient C-to-T editing at the *AtFLS2*, *AtCERK1*, and *AtTPST* target sites containing non-canonical NNNRT PAMs (Figure 2D), while AID10-nSpCas9-NG-UGI was also able to induce C-to-T editing at these target loci containing non-canonical NG PAMs, despite with lower efficiencies (Figure 2D). Notably, when compared with the PAM-stringent CBEs, these PAM-relaxed CBEs showed impaired activities at the *At-eTM166-I* target site containing the canonical PAM (Figure 2D). Importantly, both AID10-nSpCas9-NG-UGI and AID10-nSaCas9-KKH-UGI maintained a wide activity window that spanned more than 10 nucleotides (Figure 2D). To evaluate the guide-dependent off-target effects that may arise from relaxed PAM requirements, we used ampiclon deep sequencing to examine C-to-T editing frequencies of AID10-nSpCas9-NG-UGI and AID10-nSaCas9-KKH-UGI at potential off-target sites with high sequence homology (no more than 3 base mismatches), including some with non-canonical PAMs. We detected no off-target C-to-T editing at all of these sites (Supplementary Figure S9). Taken together, these results suggest that AID10 is compatible with diverse Cas9 nickases to confer a broad activity window at genomic target loci containing various PAMs.

nCas9-AID10-UGI confers a narrow activity window at the PAM-proximal protosparse

Next, we wondered whether AID10 fused to the C-terminus of nSpCas9 (nCas9-AID10-UGI, Figure 3A) might give rise to a different activity window. Therefore, we employed ampiclon deep sequencing to assess the performance of nCas9-AID10-UGI at the *AtBAK1* and *AtHDC1* target sites in Arabidopsis protoplasts. Strikingly, unlike AID10-nCas9-UGI whose activity window was located at the PAM-distal protosparse, nCas9-AID10-UGI exhibited efficient cytosine editing at the PAM-proximal protosparse (Figure 3B). Moreover, we used nCas9-AID10-UGI to edit the *OsALS* gene in transgenic rice plants, which encodes an acetolactate synthase targeted by many herbicides. As substitution of the conserved Ala96 residue of OsALS can confer gain-of-function resistance to the herbicide imidazoli-
none (60,61), we designed an sgRNA to target the coding strand of \textit{OsALS}, placing the GCG codon of Ala96 at the protospacer positions 12–14 (Figure 3C). By genotyping 39 transgenic T0 rice plants, we identified five \textit{OsALS} mutant alleles with nucleotide substitution and one mutant allele with nucleotide deletion (Supplementary Table S4). Importantly, although there are nine G\textsuperscript{C}C base pairs at the \textit{OsALS} target site, base editing by nCas9-AID10-UGI only occurred at the position 14 with no bystander editing in all five mutant alleles (Figure 3C). We isolated two T0 alleles for further analysis, in which the line \#7 carried biallelic G\textsuperscript{C}C to A\textsuperscript{T}T and G\textsuperscript{C}C to C\textsuperscript{G}G conversions and the line \#25 had heterozygous G\textsuperscript{C}C to A\textsuperscript{T}T conversion (Figure 3C and D). The G\textsuperscript{C}C to A\textsuperscript{T}T or C\textsuperscript{G}G conversion would generate A96T and A96P mutations, respectively, both of which have been documented to render imidazolinone resistance (60,61). Indeed, we observed that all progeny of line \#7 and three quarters of the progeny of line \#25 exhibited imidazolinone resistance (Figure 3E). Altogether, our data from protoplasts and transgenic plants imply that AID10 fused to the C-terminus of nSpCas9 offers a narrow activity window at the PAM-proximal protospacer.

AID10-nCas9-AID10-UGI confers a superimposed activity window

The observations that the N-terminal or C-terminal AID10 enabled base editing with distinct activity windows prompted us to test whether concurrent fusion of AID10 to both termini of nSpCas9 could lead to an overlaid activity window. To this end, transgenic rice plants were generated to express AID10-nCas9-AID10-UGI (Figure 3F) and three sgRNAs targeting the \textit{OsSWEET11}, \textit{OsSWEET13} and \textit{OsSWEET14} loci, respectively. Cytosines at the target sequences of \textit{OsSWEET11} and \textit{OsSWEET14} are located at the PAM-distal protospacer, while the only cytosine at the \textit{OsSWEET13} target sequence lies at the position 13. In 77 transgenic T0 rice plants, AID10-nCas9-AID10-UGI successfully introduced cytosine changes to at least one \textit{OsSWEET} target site (Supplementary Table S4). In particular, cytosine editing by AID10-nCas9-AID10-UGI occurred at the positions –1, 5, 7, 8, 10 and 11 of the \textit{OsSWEET11} protospacer and positions 2, 7, 9 and 10 of the \textit{OsSWEET14} protospacer, reminiscent of the activity window of AID10-nCas9-UGI (Figure 3G). Meanwhile, AID10-nCas9-AID10-UGI also edited the cytosine at the

Figure 2. Combination of AID10 with orthogonal or engineered Cas proteins expands the target scope. (A) Schematic diagram of AID10-based CBEs derived from SpCas9, SaCas9, and LbCas12a. (B) Comparison of cytosine editing efficiencies between different CBEs at endogenous genomic loci in rice (top) or \textit{Arabidopsis} (bottom) protoplasts by amplicon deep sequencing. (C) Comparison of cytosine editing efficiencies between AID10-nSpCas9-UGI and AID10-nSaCas9-UGI at three endogenous genomic loci in transgenic T1 \textit{Arabidopsis} plants. (D) Comparison of cytosine editing efficiencies between CBEs derived from PAM-relaxed Cas9 variants at endogenous genomic loci with canonical or non-canonical PAMs in \textit{Arabidopsis} protoplasts by amplicon deep sequencing. The PAMs are shown in the parentheses. In (B) and (C), the numbers in red, green, and blue indicate protospacer positions for SpCas9, SaCas9, and LbCas12a, respectively. Data in (B) and (D) are shown as mean ± s.d. of two biological replicates performed at different times.
Figure 3. nCas9-AID10-UGI and AID10-nCas9-AID10-UGI exhibit distinct activity windows relative to AID10-nCas9-UGI. (A) Schematic diagram of nCas9-AID10-UGI. (B) Comparison of cytosine editing efficiencies between nCas9-AID10-UGI and AID10-nCas9-UGI at endogenous genomic loci in Arabidopsis protoplasts by amplicon deep sequencing. Data are shown as mean ± s.d. of two biological replicates performed at different times. (C) Cytosine editing at the OsALS locus by nCas9-AID10-UGI in transgenic T0 rice lines #7 and #25 and resultant amino acid changes. The reverse complement of the target sequence is underlined and the PAM is in bold. The subscripted number indicates the protospacer position. (D) Sanger sequencing chromatograms show base conversions (marked by red arrows) described in (C). (E) Amino acid substitutions in OsALS confer herbicide resistance. The seedlings were germinated on the medium containing 0.1 mg/l imazethapyr for nine days. Scale bar = 2 cm. (F) Schematic diagram of AID10-nCas9-AID10-UGI. (G) Frequencies of cytosine editing at three endogenous genomic loci in transgenic T0 rice plants expressing AID10-nCas9-AID10-UGI and three sgRNAs, each targeting one indicated OsSWEET locus. (H) Sanger sequencing chromatograms show multiplex base conversions (marked by red arrows) by AID10-nCas9-AID10-UGI at the OsSWEET loci in two T0 rice lines.

position 13 of the OsSWEET13 protospacer, showing a comparable activity window to nCas9-AID10-UGI (Figure 3G). Multiplex base editing at all three target loci was detected in two lines (i.e. #21 and #75) (Figure 3H). We also targeted AID10-nCas9-AID10-UGI to a C-rich sequence at the OsSLR1 locus in transgenic rice plants, in which the base editing efficiency appeared to be low (Supplementary Table S4). However, we still detected cytosine substitutions at either PAM-distal or -proximal protospacer in the OsSLR1-edited lines (Supplementary Figure S10). Taken together, these results indicate that AID10-nCas9-AID10-UGI possesses a highly flexible activity window at either PAM-distal or -proximal protospacer.

Dual base editors enable cytosine and adenine co-editing at the same target sites in plant cells

Inspired by AID10-nCas9-AID10-UGI, we reasoned that combination of AID10 and ABE7.10 (4) in a single base editor might allow flexible editing of either cytosine or adenine at a genomic target sequence. Since only the N-terminal ABE7.10 can work efficiently for A-to-G editing (4,62), we constructed ABE7.10-nCas9-AID10 (Figure 4A) and assessed its performance at the OsIPA1 or OsCKX2 locus in transgenic rice plants. However, we failed to detect any base change at the OsIPA1 site in 29 transgenic T0 lines (Supplementary Table S4), while only PAM-proximal C-to-T substitutions by AID10 were detected at the OsCKX2 site in 2 out of 23 transgenic T0 plants (Supplementary Figure S11 and Table S4).

To comprehensively evaluate the performance of ABE7.10-nCas9-AID10, we checked two more genomic loci containing AC or CA tandem repeats (referred to as AtChr3 and AtChr4, Supplementary Table S1) in Arabidopsis protoplasts by ampiclon deep sequencing. ABE7.10-nCas9, nCas9-AID10, and AID10-nCas9 were included for side-by-side comparison (Figure 4A). Amplicom deep sequencing showed that ABE7.10-nCas9-AID10 was able to catalyze C-to-T (Figure 4B) or A-to-G (Figure 4C) conversions at the same target site. Although there were multiple cytosines and adenes in the two protospacers, C-to-T and A-to-G conversions by ABE7.10-
Figure 4. Cytosine and adenine co-editing at the same target sequences by AID10-based dual base editors. (A) Schematic diagram of the dual base editor ABE7.10-nCas9-AID10. (B, C) ABE7.10-nCas9-AID10 mediates both C-to-T (B) and A-to-G (C) conversions at two endogenous genomic loci in Arabidopsis protoplasts as revealed by amplicon deep sequencing. Data are shown as mean ± s.d. of two biological replicates performed at different times. (D) Comparison of editing product compositions between ABE7.10-nCas9-AID10 and single-deaminase base editors. The percentage of indicated sequencing reads among total reads is presented as mean ± s.d. of two replicates. (E) The ten most abundant mutant alleles with concurrent cytosine and adenine substitutions by ABE7.10-nCas9-AID10 in one replicate of (D). Frequencies of indicated sequencing reads are shown on the right. The A-to-G, C-to-T, and C-to-A substitutions are in blue, red, and green, respectively. (F) Schematic diagram of eight dual base editors with different protein architectures. (G) Comparison of cytosine (top) and adenine (bottom) editing efficiencies between these base editors at the AtChr4 locus in Arabidopsis protoplasts by amplicon deep sequencing. Data are shown as mean ± s.d. of two biological replicates performed at different times. (H) Comparison of editing product compositions at the AtChr4 locus generated by these base editors. (I) The ten most abundant mutant alleles with concurrent cytosine and adenine substitutions by ABE8e-nCas9-AID10-UGI or AID10-ABE8e-nCas9-UGI in one replicate of (H). The A-to-G and C-to-T substitutions are in blue and red, respectively.

nCas9-AID10 mostly occurred at the positions 11–14 and 4–8, respectively, resembling the editing pattern by nCas9-AID10 or ABE7.10-nCas9 on its own (Figure 4B and C). The C-to-T editing efficiency of ABE7.10-nCas9-AID10 was comparable to or slightly lower than that of nCas9-AID10, whereas the A-to-G editing efficiency of ABE7.10-nCas9-AID10 was dramatically reduced relative to that of ABE7.10-nCas9 (Figure 4B and C). Nevertheless, deep sequencing reads revealed synchronous cytosine and adenine conversions at the same target sequences in protoplasts expressing ABE7.10-nCas9-AID10 with frequencies of 1.09–2.04%, whereas only cytosine or adenine changes were detected in protoplasts expressing nCas9-AID10 or ABE7.10-nCas9 alone (Figure 4D and E).

The decreased A-to-G editing capability of ABE7.10-nCas9-AID10 relative to ABE7.10-nCas9 may suggest steric hindrance of the C-terminal AID10 on the activity of the N-terminal ABE7.10 (Figure 4C). Meanwhile, Hua and colleagues have recently reported that the TadA7.10 by itself can achieve higher A-to-G editing efficiency than ABE7.10.
(i.e., the TadA-TadA7.10 heterodimer) in rice (63). Therefore, we constructed additional dual base editors, including TadA7.10-nCas9-AID10, AID10-ABE7.10-nCas9 and AID10-TadA7.10-nCas9 (Supplementary Figure S12A), and compared their performance with that of ABE7.10-nCas9-AID10 in Arabidopsis protoplasts by amplicon deep sequencing. Intriguingly, dual base editors containing the N-terminal or C-terminal AID10 induced C-to-T conversions at the protospacer positions 1–11 and 11–14, respectively, suggesting that the fusion orientation of AID10 is the key determinant for the cytosine editing window (Supplementary Figure S12B). By contrast, adenine editing windows of the four dual base editors were located similarly at the protospacer positions 4–8 (Supplementary Figure S12B). All dual base editors were able to mediate simultaneous cytosine and adenine conversions at the same target sequences with frequencies of 0.41–2.06% (Supplementary Figure S12C and D). AID10-ABE7.10-nCas9, despite with a different cytosine editing window from ABE7.10-nCas9-AID10 (Supplementary Figure S12B), failed to improve the adenine editing or cytosine/adenine co-editing efficiency (Supplementary Figure S12C). Using TadA7.10 to replace nCas9 in protoplasts, Amplicon deep sequencing of the AtChr4 locus revealed that the addition of UGI could promote the cytosine and adenine co-editing efficiency. Indeed, ABE7.10-nCas9-AID10-UGI and AID10-ABE7.10-nCas9-UGI (Figure 4F) exhibited a dramatically increased C-to-T editing efficiency for the cytosine and adenine co-editing efficiency relative to AID10-nCas9-AID10, while there was a 3.4-fold increase with homozygous C-to-T mutation (Figure 5B). Among 325 transgenic T1 plants, five heterozygous mutant lines carrying the intended C-to-T conversion were identified (Supplementary Figure S13C and Table S4), from which we harvested two T2 alleles (#13-10 and #205-3) with homozygous C-to-T mutation (Figure 5D). Consistently, #13–10 and #205–3 seedlingsexhibitedshorterhypocotyls in the dark resembling the bak1-4 mutant (Figure 5E). Interestingly, dual base editors containing the ACG, ATA or ACA trinucleotide in the 5′ UTR of the target gene were able to generate efficient C-to-T editing of downstream primary ORFs (pORFs) through ribosome stalling (67), we reasoned that creation of a uORF by changing the ACG, ATA or ACA trinucleotide in the 5′ UTR to ATG using our base editor toolkit would provide an effective strategy to down-regulate target gene activity (Figure 5A).

To provide a proof of concept, we intended to generate uORFs in the 5′ UTR of the AtBAK1 gene in Arabidopsis, which encodes the co-receptor for the plant hormone brassinosteroid (68). As there are three ACG trinucleotides in the 5′ UTR of AtBAK1, we evaluated the effects of changing them to ATG, singly or combinatorially, on the AtBAK1 expression in protoplasts using a luciferase reporter as a proxy (Supplementary Figure S13A). Creation of different uORFs inhibited the luciferase expression to variable levels, among which the uORF-2 was able to reduce the luciferase expression by 84% (Supplementary Figure S13B). Therefore, we employed AID10-nCas9-UGI to generate an endogenous uORF-2 in the 5′ UTR of AtBAK1 by inducing a C-to-T change at the protospacer position 2 (Figure 5B). Among 325 transgenic T1 plants, five heterozygous mutant lines carrying the intended C-to-T conversion were identified (Supplementary Figure S13C and Table S4), from which we harvested two T2 alleles (#13-10 and #205-3) with homozygous C-to-T mutation (Figure 5B). Strikingly, immunoblot analysis using the anti-AtBAK1 antibody indicated that the accumulation of endogenous AtBAK1 in these two mutant alleles was almost abolished using the bak1-4 T-DNA insertion null allele (68) as a reference (Figure 5C). The basal immunoblot signals in these bak1 mutant plants were attributed to the cross-reactivity of the anti-AtBAK1 antibody to AtBKK1 (50), a paralog of AtBAK1. Consistently, #13–10 and #205–3 seedlings exhibited shortened hypocotyls in the dark resembling the bak1-4 null allele (Figure 5D and E), indicative of functional knockout of AtBAK1 in these mutant alleles. These results suggest that creation of a suitable uORF in the 5′ UTR of the target gene through base editing can quantitatively knockdown or even knockout target gene expression.

Base editing can be repurposed for tunable gene knockdown via creating uORFs

To expand the usefulness of our base editors in plant research, we explored whether tunable gene knockdown could be achieved by base editors via creating uORFs in the 5′ untranslated region (5′ UTR) of the target gene. As the uORFs have been shown to inhibit the translation initiation of downstream primary ORFs (pORFs) through ribosome stalling (67), we reasoned that creation of a uORF by changing the ACG, ATA or ACA trinucleotide in the 5′ UTR to ATG using our base editor toolkit would provide an effective strategy to down-regulate target gene activity (Figure 5A).

Base editing is useful for loss-of-function analysis of endogenous target mimics of miRNAs

Plant endogenous target mimics (eTMs) and animal competing endogenous RNAs (ceRNAs) are analogous eukaryotic machineries regulating miRNA activities through sequestering and preventing miRNAs from interacting with their authentic target mRNAs (69–71). While a large num-
Figure 5. Repurposing base editors as loss-of-function tools for studying protein-coding or non-coding genes. (A) Schematic diagram illustrating the gene knockdown strategy through creating a uORF in the 5′ UTR of the target gene by base editing. TSS, transcription start site. (B) Two homozygous mutant alleles of Arabidopsis BAK1 containing AID10-nCas9-UGI-created uORF in the 5′ UTR. The target sequence is underlined and the PAM is in bold. The ACG trinucleotides are in blue with subscripted numbers indicating the protospacer positions of target cytosines. (C) Immunoblot analysis indicates significantly decreased AtBAK1 abundances in bak1 mutant alleles relative to wild-type (WT) plants. The T-DNA insertion null mutant bak1-4 was used for reference. Note that the basal immunoblots were due to the cross-reactivity of the anti-BAK1 antibody to AtBKK1, a AtBAK1 paralog. RuBisCO, loading control. (D, E) Bak1 5′ UTR mutant seedlings resemble T-DNA insertion null seedlings. The photograph in (D) was taken at five days post germination. Scale bar = 0.5 cm. Quantification of hypocotyl lengths in (E) is presented as means ± s.d. of 20 seedlings. **P < 0.01 (two-tailed Student’s t test). (F) Predicted base pairing between miR166 and the miRNA binding site of At-eTM166-1 in Arabidopsis. (G) Two homozygous mutant alleles of the At-eTM166-1 locus generated by AID10-nCas9-UGI. The miR166 binding site is in green. The target sequence is underlined and the PAM is in bold. Cytosine conversions in mutant lines are in red. (H) qRT-PCR analysis reveals reduced transcript levels of miR166 target genes in At-eTM166-1 mutant lines. Data are presented as means ± s.d. of three biological replicates.

We conceived an approach for loss-of-function analysis of plant eTMs by introducing a cluster of nucleotide changes to disrupt the putative miRNA binding site in eTMs through base editing. To demonstrate a proof of concept, we employed AID10-nCas9-UGI to destroy the miR166 binding site in At-eTM166-1 (Figure 5F), an eTM that has been reported to negatively regulate the activity of miR166 in Arabidopsis (71). An sgRNA was designed to target the At-eTM166-1 locus, placing the four consecutive C•G base pairs at the protospacer positions 2–5 (Figure 5G). Among 71 transgenic T1 plants, eight lines harboring C•G to T•A mutations were identified (Supplementary Figure S8C and Table S4). We further isolated two T2 lines (i.e. #45–3 and #48–10) carrying homozygous C•G to T•A mutations at the positions 2–4 and 2–5, respectively (Figure 5G). If the interaction between At-eTM166-1 and miR166 was biologically relevant, we expected that these mutations would unleash miR166 to further down-regulate its target genes, including AtPHB, AtPHV and AtCNA (73). Indeed, we observed that the transcript levels of AtPHV and AtCNA were comparably decreased in these two lines (Figure 5H), implying that disruption of three internal cytosines has been sufficient to fully abolish the interaction between At-eTM166-1 and miR166. The transcript abundances of AtPHB were also diminished in both lines, whereas the line...
In summary, here we report a suite of plant base editors with different activity windows and target scopes (Figure 6 and Supplementary Table S5). This toolkit will be a valuable asset for plant research. On the one hand, we invented a strategy for gene inactivation that has a particular advantage in terms of tunability. By creating uORFs at different positions relative to the pORF, quantitative gene knockdown to varying degrees or even functional knockout can be selectively achieved for the target gene, making it a flexible and versatile tool for manipulating both lethal and non-lethal genes. When compared with other gene knockdown strategies, such as RNA interference (RNAi) and CRISPR interference (CRISPRi) (81), this strategy can mediate stable and permanent gene knockdown free of any transgene. On the other hand, we utilized base editing to effectively perturb the putative miRNA binding site in a plant eTM to study its biological function in miRNA regulation, which fills in the blank of loss-of-function tools for experimental validation of predicted plant eTMs. It is conceivable that such a strategy is immediately transferrable to studying regulatory mechanisms of predicted animal ceRNAs, which are functionally analogous to plant eTMs (70). The PAM requirement and the width of deamination window of a CBE combinatorially determine the total number of editable cytosines in a given genome. When compared to CBEs with a narrow activity window, those with a broad activity window, such as AID10-nCas9-UGI, not only make more genomic cytosines accessible but also increase mutations in the protospacers, which is advantageous for applications such as directed protein evolution (22,23,76,82) and loss-of-function analysis of eTMs/ceRNAs (Figure 5G and H). The downside of broad-window CBEs is the accompanied risk of bystander editing. Although the bystander editing can be circumvented by rational selection of an optimal base editor and sgRNA in some cases, it is inevitable in others. However, it should be emphasized that bystander mutations can be tolerated when they are synonymous in the exons or occur in the non-coding regions. Moreover, for the base editor-mediated gene inactivation, such as inducing intron mis-splicing (Supplementary Figures S3 and S7) (18,83,84) or creating premature stop codons (14,15) or uORFs (Figure 5A–E), the bystander editing is inconsequential. Also, it is possible in practice to screen a large quantity of edited plant alleles to identify those without or with acceptable bystander mutations.

In addition to developing useful plant base editors, we also explored unprecedented applications of base editing in plant research. On the one hand, we invented a strategy to down-regulate target gene activity by creating uORFs in the 5’ UTR of the target gene through base editing. When compared with the CRISPR/Cas9 technology or other base editor-mediated gene loss-of-function strategies, such as CRISPR-STOP (14) and mRNA mis-splicing (18), this new strategy for gene inactivation has a particular advantage in terms of tunability. By creating uORFs at different positions relative to the pORF, quantitative gene knockdown to varying degrees or even functional knockout can be selectively achieved for the target gene, making it a flexible and versatile tool for manipulating both lethal and non-lethal genes. When compared with other gene knockdown strategies, such as RNA interference (RNAi) and CRISPR interference (CRISPRi) (81), this strategy can mediate stable and permanent gene knockdown free of any transgene. On the other hand, we utilized base editing to effectively perturb the putative miRNA binding site in a plant eTM to study its biological function in miRNA regulation, which fills in the blank of loss-of-function tools for experimental validation of predicted plant eTMs. It is conceivable that such a strategy is immediately transferrable to studying regulatory mechanisms of predicted animal ceRNAs, which are functionally analogous to plant eTMs (70).

The PAM requirement and the width of deamination window of a CBE combinatorially determine the total number of editable cytosines in a given genome. When compared to CBEs with a narrow activity window, those with a broad activity window, such as AID10-nCas9-UGI, not only make more genomic cytosines accessible but also increase mutations in the protospacers, which is advantageous for applications such as directed protein evolution (22,23,76,82) and loss-of-function analysis of eTMs/ceRNAs (Figure 5G and H). The downside of broad-window CBEs is the accompanied risk of bystander editing. Although the bystander editing can be circumvented by rational selection of an optimal base editor and sgRNA in some cases, it is inevitable in others. However, it should be emphasized that bystander mutations can be tolerated when they are synonymous in the exons or occur in the non-coding regions. Moreover, for the base editor-mediated gene inactivation, such as inducing intron mis-splicing (Supplementary Figures S3 and S7) (18,83,84) or creating premature stop codons (14,15) or uORFs (Figure 5A–E), the bystander editing is inconsequential. Also, it is possible in practice to screen a large quantity of edited plant alleles to identify those without or with acceptable bystander mutations.

In summary, here we report a suite of plant base editors with different activity windows and target scopes (Figure 6 and Supplementary Table S5). This toolkit will be a valuable asset for plant research. On the one hand, we invented a strategy for gene inactivation that has a particular advantage in terms of tunability. By creating uORFs at different positions relative to the pORF, quantitative gene knockdown to varying degrees or even functional knockout can be selectively achieved for the target gene, making it a flexible and versatile tool for manipulating both lethal and non-lethal genes. When compared with other gene knockdown strategies, such as RNA interference (RNAi) and CRISPR interference (CRISPRi) (81), this strategy can mediate stable and permanent gene knockdown free of any transgene. On the other hand, we utilized base editing to effectively perturb the putative miRNA binding site in a plant eTM to study its biological function in miRNA regulation, which fills in the blank of loss-of-function tools for experimental validation of predicted plant eTMs. It is conceivable that such a strategy is immediately transferrable to studying regulatory mechanisms of predicted animal ceRNAs, which are functionally analogous to plant eTMs (70).
able addition to the current base editor arsenal in plants by expanding editable cytosines in a given plant genome or creating diverse base editing products at the same target sequences. We also open up novel routes for manipulating the function of protein-coding or non-coding genes through base editing.

DATA AVAILABILITY

Raw data of amplicon deep sequencing have been deposited in the NCBI Sequence Read Archive under BioProject accession code PRJNA762247 (http://www.ncbi.nlm.nih.gov/sra/PRJNA762247). Amino acid sequences of all base editors tested in this study are provided in the Supplementary Sequence.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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