A large-scale whole-genome sequencing analysis reveals highly specific genome editing by both Cas9 and Cpf1 (Cas12a) nucleases in rice

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Abstract

Background: Targeting specificity has been a barrier to applying genome editing systems in functional genomics, precise medicine and plant breeding. In plants, only limited studies have used whole-genome sequencing (WGS) to test off-target effects of Cas9. The cause of numerous discovered mutations is still controversial. Furthermore, WGS-based off-target analysis of Cpf1 (Cas12a) has not been reported in any higher organism to date.

Results: We conduct a WGS analysis of 34 plants edited by Cas9 and 15 plants edited by Cpf1 in T0 and T1 generations along with 20 diverse control plants in rice. The sequencing depths range from 45× to 105× with read mapping rates above 96%. Our results clearly show that most mutations in edited plants are created by the tissue culture process, which causes approximately 102 to 148 single nucleotide variations (SNVs) and approximately 32 to 83 insertions/deletions (indels) per plant. Among 12 Cas9 single guide RNAs (sgRNAs) and three Cpf1 CRISPR RNAs (crRNAs) assessed by WGS, only one Cas9 sgRNA resulted in off-target mutations in T0 lines at sites predicted by computer programs. Moreover, we cannot find evidence for bona fide off-target mutations due to continued expression of Cas9 or Cpf1 with guide RNAs in T1 generation.

Conclusions: Our comprehensive and rigorous analysis of WGS data across multiple sample types suggests both Cas9 and Cpf1 nucleases are very specific in generating targeted DNA modifications and off-targeting can be avoided by designing guide RNAs with high specificity.
Background

Bacterial type II CRISPR-Cas9 systems can effectively induce RNA-guided DNA double strand breaks (DSBs) [1], making them popular tools for genome editing in bacteria [2], animal cells [3], mammalian systems [4–7], and plants [8–11]. The most widely used Streptococcus pyogenes Cas9 (SpCas9) uses ~20 nucleotides (nt) of a single guide RNA (sgRNA) to recognize a complementary target DNA site along with an NGG protospacer adjacent motif (PAM) [1, 12]. More recently, type V CRISPR-Cpf1 (CRISPR-Cas12a) was shown to mediate target DNA with a TTTV PAM [13]. RNA-guided nucleases (RGNs) such as Cas9 and Cpf1 represent versatile genome editing tools that promise to advance basic science, enable personalized medicine, and accelerate crop breeding. However, Cas9 may cause undesired off-target mutations due to sgRNAs recognizing DNA sequences with one to a few nucleotide mismatches, albeit with reduced nuclease binding and cleavage activity [1, 6, 17, 18]. Although similar rules apply to Cpf1, recent studies in human cells [19, 20] have shown Cpf1 is generally more specific than Cas9.

Understanding the scope of off-target mutations in Cas9- or Cpf1-edited crops is critical for research and regulation. Previously, whole-genome sequencing (WGS) was applied for detecting off-target mutations by Cas9 in Arabidopsis [21], rice [22], and tomato [23]. Unfortunately, these studies either only looked at potential off-target sites predicted by computer programs or fell short of full analysis of all the mutations identified by WGS in edited plants. Without inclusion of enough necessary controls, such WGS studies had limited power for isolating off-target mutations in edited plants because they were unable to fully assess the levels of preexisting mutations, spontaneous mutations, and mutations caused by tissue culture- and Agrobacterium-mediated transformation. Genome-wide identification of off-target mutations by Cas9 or Cpf1 will be empowered only if all background mutations can be isolated. Furthermore, WGS-based off-target analysis of Cpf1 has not been reported in any higher organism. In recent years, WGS studies on Cas9-edited mice have generated contrasting results; one study found few off-target mutations [24] while another found many [25]. This controversy raised the urgency for comprehensive and rigorous analyses of off-target mutations using WGS in edited animals and plants. We reasoned a large-scale and well-designed study is required for comprehensive assessment of off-target effects in crops by Cas9 and Cpf1, two leading CRISPR genome editing systems. Here, we describe a large-scale WGS study to assess off-target effects of Cas9 and Cpf1 in rice, an important food crop. Our results suggest off-target mutations of Cas9 and Cpf1 are largely negligible when compared to spontaneous mutations or mutations caused by tissue culture and Agrobacterium infection in edited plants. The resulting knowledge is likely to serve as an important reference for plant researchers and regulatory agencies.

Results

Detection of off-target, spontaneous, and background mutations

To comprehensively evaluate potential off-target effects of Cas9 in rice, we generated ten T-DNA constructs to target seven genes with 12 sgRNAs, including two dual-sgRNA constructs for editing two circular RNA loci (Additional file 1: Figure S1 and Additional file 2: Table S1). All ten CRISPR-Cas9 nuclease expression constructs were active at target sites and resulted in editing frequencies ranging from 15 to 100% in T0 lines (Fig. 1a, b and Additional file 2: Table S1). For each Cas9 construct, two independent T0 plants carrying non-mosaic mutations (Additional file 1: Figure S1) were chosen for WGS. To assess off-target effects of Cpf1, we followed three previously published Lachnospiraceae bacterium ND2006 Cpf1 (LbCpf1) targeting constructs that resulted in 100% editing efficiency in T0 lines [14] (Additional file 1: Figure S2 and Additional file 2: Table S1). Two Cpf1 T0 plants per construct carrying non-mosaic on-target mutations were chosen for WGS (Additional file 1: Figure S2). For four T0 lines edited by four different Cas9 sgRNAs and two T0 lines edited by two different Cpf1 crRNAs, we selected two to five plants from each T0 line in the T1 generation for WGS (Fig. 1a, b). In addition, four wild-type (WT) plants each from three consecutive generations were also included for WGS to survey spontaneous mutations (Fig. 1b). To ensure high confidence on base calling, all 69 individual plants were sequenced at 45× to 105× in depth (Additional file 2: Table S2). A stringent mutation mapping and calling pipeline was developed for WGS analysis (Fig. 1c). Single-nucleotide variants (SNVs) and small insertions and deletions (indels) were each identified with three variant-calling software programs, with high-confident variants shared by all software being further analyzed for mutation identification (Fig. 1c). Based on our criteria, mutations with frequencies below 10% may not be called out, as such low frequency mutations may have resulted from sequencing errors.

To survey pre-existing mutations in the WT population and estimate the level of spontaneous mutations across generations, we analyzed the WGS data from 12 WT plants across three consecutive generations (Fig. 1b and Additional file 1: Figure S3). After filtering shared pre-existing mutations, we estimated an average of 23 SNVs and 18 indels as spontaneous mutations from
Fig. 1 (See legend on next page.)
parents to progeny in rice (Fig. 2a, b). We calculated the spontaneous mutation rate at \( \sim 5.4 \times 10^{-8} \) per site per diploid genome per generation, which is in line with the rates previously reported in maize (2.2–3.9 \( \times 10^{-8} \)) [26] but higher than the rate in *Arabidopsis* (7–7.4 \( \times 10^{-9} \)) [27, 28].

To assess mutations generated by tissue culture and *Agrobacterium* infection, we produced and sequenced four types of control plants: tissue culture only, tissue culture with *Agrobacterium* transformation of Cas9 without sgRNA, and tissue culture with *Agrobacterium* transformation of Cpf1
without crRNA (Fig. 1b and Additional file 1: Figure S4). Tissue culture is known to be mutagenic, causing somaclonal variations [29]. Indeed, the two tissue culture-only samples contained an average of 114 SNVs and 36 indels (Fig. 2a, b), resulting in a background mutation rate of $1.86 \times 10^{-7}$, which is similar to the rates ($1.7-3.3 \times 10^{-7}$) previously published [30]. Importantly, similar numbers of SNVs were observed from Agrobacterium-infected or Cas9/Cpf1 backbone-transformed plants (Fig. 2a). These three controls generated ~15 to 41 more indels compared to tissue culture-only samples (Fig. 2b), suggesting Agrobacterium infection is mutagenic with a preference for introducing indels. This warrants further investigation as these three controls show large variations on indel counts. We mapped all identified mutations from these four control types to the rice genome across 12 chromosomes (Additional file 1: Figure S5). Further analysis of the genome-wide distribution of these background mutations revealed high enrichment of SNVs in transposable elements (TEs) and repeats (Fig. 2c), as well as high enrichment of indels in repeats (Fig. 2d).

**SNVs and indels identified in edited T0 plants are largely background mutations**

WGS of 20 Cas9 and six Cpf1-edited T0 lines confirmed all target site mutations that were initially identified with Sanger sequencing (Additional file 1: Figures S1 and S2 and Additional file 2: Table S1). We identified SNVs and indels in these Cas9 T0 lines (Additional file 1: Figure S6) and Cpf1 T0 lines (Additional file 1: Figure S7) and mapped these mutations to the rice genome (Additional file 1: Figure S5). We found their numbers are close to those in Cas9 or Cpf1 backbone controls, with about twice as many SNVs as indels (Fig. 3a, b). This mutation pattern is not consistent with Cas9- or Cpf1-generated mutations in rice which are largely indels [9, 14]. For example, all target site mutations in these selected 26 T0 lines are indels (Additional file 1: Figures S1 and S2 and Additional file 2: Table S1). The SNV and indel mutations in Cas9- and Cpf1-edited T0 samples share similar genome-wide distribution with the tissue culture-related controls (Additional file 1: Figure S5). We identified a total of 31 T-DNA insertion events in 26 T0 lines and found T-DNA copy numbers ranging from 1 to 3; most T0 lines had only one T-DNA insertion (Additional file 1: Figure S8). No significant difference was found for the numbers of SNVs and indels among T0 lines with different T-DNA copy numbers (Fig. 3c, d). Cas9-J and Cas9-K T0 lines each expressed a dual-sgRNA construct for simultaneous expression of two sgRNAs, targeting two putative circle RNA genes (Fig. 1a). No significant difference was found for the numbers of SNVs and indels in these four dual-sgRNA lines and the other 22 single sgRNA lines (Fig. 3e, f). Moreover, there is no correlation between the numbers of SNVs or indels and the on-target editing efficiency by Cas9 or Cpf1 in these T0 plants (Fig. 3g, h). All these analyses strongly suggest mutations in these genome-edited T0 lines are mostly background mutations caused during tissue culture and Agrobacterium-mediated transformation.

**Identification of true off-target mutations in T0 plants**

To identify true off-target mutations in the T0 plants, we first evaluated the specificity of 12 sgRNAs of Cas9 and three crRNAs of Cpf1 with CRISPOR [31] and Cas-OFFinder [32]. With a stringent criterion allowing only a 1-nt mismatch in the protospacer, three Cas9 sgRNAs (Cas9-D, Cas9-E, and Cas9-J-sgRNA01; Fig. 1a) had predicted off-target sites (Fig. 4a and Additional file 2: Table S3). When we mapped all identified mutations to these potential off-target sites by allowing up to 10-nt mismatches to the protospacers of Cas9 (Additional file 1: Figure S9) and Cpf1 (Additional file 3: Figure S10), only Cas9-J-sgRNA01 showed evidence of true off-targeting. It is worth noting that these off-target sites showed high sequence homology to the Cas9-J-sgRNA01 target site and could be accurately predicted by software such as CRISPOR and Cas-OFFinder (Additional file 2: Table S3). We reasoned true off-target mutations are likely to occur separately in independent T0 lines. Indeed, among 12 off-target sites identified for Cas9-J-sgRNA01, seven sites were overlapped between two T0 lines while the remaining five sites were only validated from one T0 line (Fig. 4b, c). All 12 off-target sites show very high sequence homology with the target site (Fig. 4c). Among them, one site at Chr1:22043904 is technically an on-target site because it has the same 20-nt protospacer with 1-nt silent mismatch in the PAM (CGG vs TGG). For the remaining 11 true off-target sites, eight sites carry one mismatch mutation in the 20-nt protospacer. For the additional three sites with two or three mismatch mutations, only one mutation is present in the 1–18-nt sequence from the PAM (Fig. 4c). Further analysis of these 12 off-target sites found four have silent mutations in NGG PAM and one has a non-canonical CAG PAM, which was reported as an alternative PAM (NAG) for SpCas9 nuclease [33] and recently shown to mediate Cas9 activity in rice [34]. All mutations at these 12 sites were indels, and, importantly, the two Cas9-J T0 lines carried distinct alleles at these sites (Fig. 4d and Additional file 3: Figure S11); validating these mutations were truly caused by Cas9.

Cas9-E sgRNA was predicted by CRISPOR and Cas-OFFinder to contain six off-target sites when up to a 3-nt mismatch was allowed (Fig. 4a and Additional file 2: Table S3). However, no off-target mutations were found at these predicted sites. Although the two Cas9-E T0 lines shared seven SNVs and three indels (Fig. 4b), these ten shared mutations had very poor sequence homology to
Fig. 3 Detailed analysis of mutations at Cas9- or Cpf1-edited T0 plants. 

- **a, b**: Average numbers of SNVs and indels detected in 26 T0 plants edited by Cas9 or Cpf1.
- **c, d**: Average numbers of SNVs and indels in edited T0 plants with different numbers of T-DNA insertions.
- **e, f**: Average numbers of SNVs and indels in Cas9-edited T0 plants expressing one or two sgRNAs (in Cas9-J and Cas9-K).
- **g, h**: Pearson correlation between on-target editing frequency and the numbers of SNV or indel mutations in Cas9- and Cpf1-edited T0 plants. Error bars in a–f indicate s.e.m.
Fig. 4 (See legend on next page.)

| Sample | Case A | Case B | Case C | Case D | Case E | Case F | Case G | Case H |
|--------|--------|--------|--------|--------|--------|--------|--------|--------|
|        |        |        |        |        |        |        |        |        |

**Nucleotide mismatch**

| Sample | Case A | Case B | Case C | Case D | Case E | Case F | Case G | Case H |
|--------|--------|--------|--------|--------|--------|--------|--------|--------|
|        |        |        |        |        |        |        |        |        |

Off-target sites: WGS identified via computer prediction based on different numbers of nucleotide mismatches.
the target site (Fig. 4e). Only five sites contained the NGG PAM. Among them, the site sharing highest sequence homology with the target site still contained a 10-nt mismatch, making it unlikely to be a true off-target site. Unlike indels found in Cas9-J samples, these putative off-target mutations are mostly SNVs (Fig. 4b, f). Furthermore, both independent T0 lines always carried the same mutant alleles (Fig. 4f and Additional file 3: Figure S11). These observations suggest that the ten shared mutations of two Cas9-E T0 lines were not caused by Cas9, but were pre-existing mutations from a parental line.

Cas9 was previously shown to induce off-target mutations at sites with missing or extra nucleotides when compared to the target site, which forms bulges when targeted by guide RNAs [35]. To detect such off-target mutations, we extracted all T0 mutation site flanking sequences (25 bp upstream and downstream) and aligned them to corresponding sgRNA/crRNA sequences using BLAST. Only Cas9-J1 and Cas9-J2 samples had alignments to the Cas9-J-sgRNA01 target (15 in Cas9-J1 and ten in Cas9-J2); other samples had no hit. None of the detected mutations were caused by bulge-forming DNA–sgRNA recognition. We also investigated whether DNA translocation events were induced by Cas9 or Cpf1 by searching for structural variants (SVs) and gene fusion events in the whole rice genome. We did not detect any translocation event in all T0 lines. Given the level of nuclease-induced DNA translocation can be used for assessing targeting specificity [36], absence of detectable translocation events in all T0 samples here indicates these Cas9 and Cpf1 reagents are indeed very specific, limiting cleavage activity almost exclusively to the target sites.

No evidence of off-target mutations in T1 plants
Our analysis of T0 plants suggested 11 out of 12 Cas9 sgRNAs and all three Cpf1 crRNAs are very specific as no off-target mutations were detected. However, lack of off-target mutations might be attributed to low expression or activity of Cas9 or Cpf1. It is also important to determine whether continued expression of the RGNs into the next generation will result in de novo off-target mutations. Therefore, we decided to sequence 14 T1 plants from Cas9 T0 lines with diverse levels of on-target editing efficiency (15, 60, 75, and 100%) at four target sites and nine T1 plants from Cpf1 T0 lines at two target sites (Fig. 1a, b and Additional file 1: Figure S1). Germline-transmitted on-target mutations in 14 Cas9-edited or nine Cpf1-edited T1 lines were validated by Sanger sequencing (Additional file 3: Figures S12 and S13). With WGS analysis, we identified all SNVs and indels in Cas9 T1 lines (Additional file 3: Figure S14) and Cpf1 T1 lines (Additional file 3: Figure S15). The WGS results confirmed the germline-transmitted on-target mutations (Additional file 3: Figures S12 and S13 and Additional file 2: Table S1). Among all other SNVs and indels, most of them were identified in the corresponding T0 lines, suggesting they have been fixed (Additional file 3: Figure S16). For the other new mutations identified in T1 lines, the average number of SNVs ranged from 9 to 29 (Fig. 5a), while the average of indels ranged from 10 to 28 (Fig. 5b). Such spontaneous mutation rates are consistent with the spontaneous mutation rates we found earlier in WT samples (Fig. 2a, b), which are also in line with a previous study [28].

These new mutations were mapped to the rice genome together with new mutations that were discovered in WT plants across two generations (Additional file 3: Figure S17). The genome distribution of these new mutations in T1 lines also showed enrichment in repeats (Additional file 3: Figure S16), consistent with the spontaneous mutations discovered in the WT (Fig. 2c, d). Detailed analysis of SNVs among all sample types revealed T1 lines have higher rates of G:C > A:T transitions than T0 lines (Additional file 3: Figure S18), consistent with the observation on spontaneous mutations in Arabidopsis [27]. Further analysis of T1 lines either with or without the Cas9 transgene did not reveal any difference in the numbers of new SNVs and indels among these two subpopulations (Fig. 5c, d). By applying similar methods from the analysis of T0 plants, we were unable to identify any off-target mutations by Cas9 or Cpf1 in T1 lines. Given most T1 lines analyzed still carry the RGN constructs, our results suggest continued expression of Cas9 or Cpf1 constructs did not cause de novo off-target mutations in T1 lines.

To further assess the new mutations found in T1 lines, we calculated and compared the allele frequency of SNVs and indels among four groups: tissue culture controls, T0 plants, T1 plants, and WT (Fig. 5e). The tissue
culture controls and Cas9/Cpf1 T0 lines share strikingly similar (mostly heterozygous-like) allele frequency distribution. This reiterates our earlier conclusion that all mutations in T0 samples (except a few found in Cas9-J samples) are background mutations. By contrast, T1 plants show more homozygous-like SNVs (0.75 to 1.0 in allele frequency) and somatic-like indels (0 to 0.25 in allele frequency). This trend of rapidly fixing SNVs and the increase of somatic indels in T1 is interesting and relatively in line with the observation in WT plants.

**Discussion**

More attention has been given to the specificity of CRISPR-Cas RGN systems in humans than in animals or plants due to medicinal applications of RGNs. Earlier WGS studies in human cells found low incidence of off-target mutations by Cas9 [37, 38]. Recently, two WGS off-target studies in mice showed conflicting results [24, 25]. However, the study that claimed unexpected large-scale off-target effects by Cas9 may be flawed due to limitations in its experimental design and WGS data analysis [39]. Given the wide adoption of CRISPR-Cas systems in agriculture, with genome-edited crop products reaching market in record time [40], it becomes urgent to conduct large-scale and exhaustive WGS analysis of off-target effects by Cas9 and Cpf1 (Cas12a), two leading RGN systems, in agriculturally important crops. Such studies will help assess the safety of Cas9 and Cpf1 in precise crop breeding as well as provide valuable information to scientists, breeders, regulators, and consumers.

In this study, we conducted a large-scale WGS analysis for detecting potential off-target mutations caused by 12 Cas9 sgRNAs and three Cpf1 crRNAs in rice, an important food crop. We confirmed WGS-identified mutations by Sanger sequencing at randomly selected sites with a 100% success rate (Additional file 2: Table S4), which is consistent with the high quality of our WGS data. Our experimental design took into account background
mutations caused by tissue culture and Agrobacterium-mediated transformation, pre-existing mutations in parents, and spontaneous mutations that arise from seed propagation. Through sequencing 20 control plants of different types and 49 Cas9 or Cpf1-edited T0 and T1 plants, we only found true off-target mutations in two T0 lines expressing Cas9 protein with Cas9-J-sgRNA01. Importantly, these empirically validated off-target sites can be readily predicted computationally. Our examination of T1 plants that continue to carry Cas9-sgRNA or Cpf1-crRNA did not reveal off-target mutations, suggesting continued presence of the RGN reagents with varying activity in plants does not cause off-target mutations if the guide RNAs are well-designed for specificity. This observation is also highly significant because it encourages the use of Cas9 and Cpf1 in certain breeding applications that may require expression of RGNs across several generations. For example, a RGN cassette may be introduced from a transgenic line into a transformation-recalcitrant variety of the same plant species for genome editing with simple genetic crossing.

Our study also provided insights into avoiding off-target effects of Cas9 and Cpf1 in edited crops. To minimize off-target effects, many systems have been developed, including paired Cas9 nickases [41], high fidelity Cas9 proteins [42–44], FokI-dCas9 fusions [45, 46], truncated sgRNAs [47], and ribonucleotide protein (RNP) delivery [48]. To assess and identify off-target sites, in vivo [18, 36] and in vitro [49–51] tools have also been developed in human cells, which may be applied in plants. Our WGS analysis with WT SpCas9 and LbCpf1 proteins did not find off-target mutations for 14 out of 15 guide RNAs tested in T0 and T1 plants, suggesting utilization of a high-fidelity enzyme, which are typically of lower activity, may be unnecessary in crop applications. When a mismatch up to 3 nt of the protospacer is allowed, Cas9-OFFinder programs predicted a total of 37 off-target sites for 7 out of 11 Cas9 sgRNAs. Yet, we could not detect any mutations at these putative off-target sites. Alternatively, Cas9-OFFinder predicted all the off-target sites that we identified for Cas9-J-sgRNA01; many of the sites have just 1-nt mismatch to the protospacer of the target site. Therefore, we can deduce a simple rule to alleviate off-target effects: make sure even the highest scored potential off-target sites will have at least a 2-nt mismatch to the seed sequence of the protospacer. We note this may not always be possible if the target sequence shares many homologous sequences in the genome. For example, maize has a very repetitive genome and wheat has A, B, D sub-genomes that share high similarity. In these cases, targeted amplicon sequencing using next-generation sequencing technologies may be an appropriate and cost-effective method to look for off-target mutations [52].

Finally, we hope our data can be a valuable reference for regulatory agencies and other entities. It is reasonable and necessary to scrutinize any new technology for its efficacy and safety. Cas9 and Cpf1, as new crop-breeding technologies, are no exception. Although Cas9-based off-target effects have been studied by WGS in plants [21–23], our study differs from previous studies significantly with regard to scale, depth, and comprehensiveness. Our research also represents the first report of using WGS to assess off-targeting by Cpf1 in any edited higher eukaryotic organism. We could not find any off-target mutations in 47 out of 49 rice plants edited by 11 Cas9-sgRNA and three Cpf1-crRNA constructs. This precise level of genome modification is in stunning contrast to many conventional breeding technologies. For example, we found that even the safest breeding approach, harvesting seeds from parental lines, introduces ~30 to 50 spontaneous mutations into the next generation in rice. We also observed ~200 tissue culture-introduced somaclonal variations per rice plant, even though few are affecting coding sequences. In conclusion, our data support a recent call to “Regulate genome-edited products, not genome editing itself” [53].

Methods

Plant material and growth conditions
This study used the rice variety Nipponbare (Oryza sativa L. ssp. Japonica cv. Nipponbare). All plants were grown in growth chambers under controlled environmental conditions with a 16/8 h light/dark regime at 28 °C and 60% relative humidity.

Vector construction
Plasmids encoding Cas9 and a single sgRNA were generated by ligating annealed oligos with a 4-bp overhang into a BsaI digested backbone (either pZHY988 or pTX172) [54–56]. Plasmids with two sgRNAs were created by ligating pZHY988 with a 485-bp fragment, after digestion with BsaI. This 485-bp fragment contains two sgRNAs generated by overlap extension PCR [55]. All CRISPR-Cpf1 (CRISPR-Cas12a) nuclease expression vectors were reported in our previous study [14]. The sequences of all primers used to construct vectors are shown in Additional file 2: Table S5.

Rice stable transformation

Agrobacterium-mediated rice transformation was performed as described in published protocols [57] with slight modification [16, 56]. The binary vectors were introduced into Agrobacterium tumefaciens strain EHA105 by the freeze-thaw method [58]. For rice transformation, dehusked seeds were sterilized with 70% ethanol for 1 min. Afterwards, seeds were washed five times with sterile water, then further sterilized for 15 min with a 2.5% sodium
hypochlorite solution containing a drop of Tween 20. The washing and sterilization step were repeated, this time without addition of Tween. Seeds were then rinsed an additional five times before being dried on sterilized filter paper and cultured on solid medium at 28 °C in a dark growth chamber for 2–3 weeks. Actively growing calli were collected for subculture at 28 °C in the dark for 1–2 weeks. *Agrobacterium* cultures were collected and resuspended in liquid medium (OD600 = 0.06–0.1) containing 100 μM acetosyringone. Rice calli were immersed in the *Agrobacterium* suspension for 30 min, then dried on sterilized filter paper and co-cultured for 3 days on solid medium at 25 °C in a dark growth chamber. The infected calli were moved to a sterile plastic bottle and washed five times with sterile water to remove excessive *Agrobacterium*. After being dried on sterilized filter paper, these calli were transferred onto screening medium at 28 °C in a dark growth chamber for 5 weeks. During the screening stage, infected calli were transferred to fresh screening medium every 2 weeks. After the screening stage, actively growing calli were moved onto regenerative medium for regeneration at 28 °C with a 16 h light/8 h dark cycle. After 3–4 weeks, transgenic seedlings were transferred to sterile plastic containers containing fresh solid medium and grown for 2–3 weeks before being transferred into soil. Transgenic rice plants were grown in a growth chamber at 28 °C with a 16 h light/8 h dark cycle.

**Mutagenesis analysis at target sites**

Genomic DNA was extracted from transgenic plants using the CTAB method [59]. The genomic region flanking the CRISPR target site for each gene was amplified and sequenced. Samples with heterozygous and biallelic mutations were decoded using CRISP-ID [60].

**WGS and data analysis**

For each sample, about 1 g of fresh leaves were collected from seedlings between 5 and 6 weeks old. DNA samples were extracted using the Plant Genome DNA Kit (Tiangen) as described by the manufacturer. All 69 samples were sequenced by Bionova (Beijing, China) using the Illumina X10 platform. Adapters were trimmed using SKEWER (v. 0.2.2) [61] and the Illumina TruSeq adapter. Cleaned reads were mapped to rice reference sequence TIGR7 (http://rice.plantbiology.msu.edu/) [62] with BWA (v. 0.7.15) software [63]. The Genome Analysis Toolkit (GATK) [64] was used to realign reads near indels and recalibrate base quality scores by following GATK best practices [65]. A known SNPs and indels database for GATK best practices was downloaded from Rice SNP-Seek Database (http://snp-seek.irri.org/) [66]. Whole genome SNVs were detected by LoFreq [67], MuTect2 [68], and VarScan2 [69]. Whole genome indels were identified using MuTect2 [68], VarScan2, and Pindel [70]. Bedtools [71] and BCFtools [72] were used to process overlapping SNVs/indels. Off-target sites were predicted with CRISPOR [31] online and Cas-OFFinder software [32] by allowing up to 10-nt mismatch. A genome-wide map of mutations was plotted with Circos software [73]. Structural variants and translocation events were analyzed using TopHat2 [74] with the ‘-fusion-search’ parameter, DELLY [75] with default parameters, and manually checking with IGV software [76]. NCBI BLAST+ with the ‘-task blastn-short’ parameter for off-target mutation site analysis, which includes mismatch, deletion, and insertion. Read-mapping screenshots were from Golden Helix GenomeBrowse visualization tool v2.1. Data processing and analyses were completed using R and Python. One T1 sample of Cas9-CC2 was excluded from analyses due to contamination of fungal DNA.

**Additional files**

**Additional file 1:** Figures S1–S9. Supplemental Figures- part I. (PPTX 12111 kb)

**Additional file 2:** Tables S1–S5. Supplemental Tables. (XLSX 43 kb)

**Additional file 3:** Figures S10–S18. Supplemental Figures-part II. (PPTX 13267 kb)

**Acknowledgements**

We thank Dr. Yusheng Qin at Soil and Fertilizer Research Institute in Sichuan Academy of Agricultural Sciences for plant care.

**Funding**

This work was supported by the National Natural Science Foundation of China (31330017 and 31771486) and the Sichuan Youth Science and Technology Foundation (2017JQ0005) to YZ. This work was also supported by grants, including the National Transgenic Major Project of China (2018ZX08022001–003) to YZ, and TZ, the Jiangsu Specially-Appointed Professor and the Priority Academic Program Development of Jiangsu Higher Education Institutions (PAPD) to TZ, and startup funds provided by University of Maryland to YQ.

**Availability of data and materials**

The WGS raw data reported in this article are listed in Additional file 2: Table S2 and have been deposited in the Sequence Read Archive in National Center for Biotechnology Information (NCBI) under the accession number PRJNA420933 (https://www.ncbi.nlm.nih.gov/bioproject?term=PRJNA420933) [77] and the Genome Sequence Archive in Beijing Institute of Genomics (BIG) under the accession number PRJCA000656 (http://bigd.big.ac.cn/bioproject/browse/PRJCA000656) [78].

**Authors’ contributions**

YZ proposed the project. YQ, TZ, and YZ designed the experiments. XT, JZ, DZ, and AM generated CRISPR-Cas nuclease vectors. XT, JZ, LT, XX, DZ, and XZ generated stable transgenic rice and identified the rice mutants. XT, JZ, QR, ZZ, BL, and XZ prepared samples for WGS and confirmed WGS results with Sanger sequencing. GL, QY, ZG, and TZ performed WGS data analysis. YQ, TZ, and YZ analyzed the data and wrote the manuscript. All authors read and approved the final manuscript.

**Competing interests**

The authors declare that they have no competing interests.

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