Multigene Editing via CRISPR/Cas9 Guided by a Single-sgRNA Seed in
*Arabidopsis*

**A Single sgRNA Seed for Multigene-editing**

**KEYWORDS**
Single sgRNA Seed, CRISPR/Cas9, Multigene Editing, *AtRPL10*

**Summary** We report that a single-sgRNA seed is capable of guiding CRISPR/Cas9 to simultaneously edit multiple genes *AtRPL10A*, *AtRPL10B* and *AtRPL10C* in *Arabidopsis*. Our results also demonstrate that it is possible to use CRISPR/Cas9 technology to create *AtRPL10* triple mutants which otherwise cannot be generated by conventional genetic crossing. Compared to other conventional multiplex CRISPR/Cas systems, a single sgRNA seed has the advantage of reducing off-target gene-editing. Such a single sgRNA seed-induced gene editing system might be also applicable to modify other homologous genes or even less-homologous sequences for multiple gene-editing in plants and other organisms.

Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR)/CRISPR-associated 9 (CRISPR/Cas9) is an adaptive immune mechanism that protects bacteria and archaea from extrachromosomal DNA and viral invasions (Jinek et al. 2012). CRISPR/Cas9 generates double-stranded breaks (DSBs) under specific guidance of a single guide RNA (sgRNA). These DSBs can then be repaired either by homologous recombination or predominantly by non-homologous end-joining, which leads to introduction of mutations such as nucleotide substitution, insertion or deletion into the targeted DNA molecules (Jinek et al. 2012; Cong et al. 2013). Such an ancient defense has been exploited for efficient genome/gene editing in organisms across kingdoms (Jinek et al. 2012; Cong et al. 2013; Fu et al. 2013; Mao et al. 2013; Gao and Zhao, 2014; Ma et al. 2015; Yan et al., 2015; Kim et al., 2016; Shen et al., 2016).
Moreover, multiplex CRISPR/Cas9-based gene editing can also be simultaneously achieved through the use of different sgRNAs in animals and plants (Cong et al. 2013; Feng et al. 2014; Wang et al. 2015; Yan et al. 2015). It remains to be elucidated, however, whether multigene-editing via CRISPR/Cas9 can be directed by a single sgRNA seed.

To address this, we searched the Arabidopsis genome database and identified the \textit{AtRPL10} family that includes three homologous members \textit{AtRPL10A} (AT1G14320), \textit{AtRPL10B} (AT1G26910) and \textit{AtRPL10C} (AT1G66580) coding for the Ribosomal Protein Large 10 subunits. The three \textit{AtRPL10} genes reside at different loci on Arabidopsis chromosome 1 (Figure S1), sharing 81-88% nucleotide (nt) identities, and their protein products are 95-98% identical (Table S1) \textit{AtRPL10A} and \textit{AtRPL10B} are expressed in female and male reproductive organs whilst \textit{AtRPL10C} is restricted to pollen grains. The three multifunctional genes are involved in protein translation and plant response to viral infection and abiotic stress (Falcone Ferreyra et al. 2013; Zorzatto et al. 2015). Homozygous \textit{AtRPL10A} T-DNA insertion mutation is lethal and RNAi of \textit{AtRPL10B} affects plant growth, although \textit{AtRPL10C} knockout results in no phenotypic change (Falcone Ferreyra et al. 2010). Interestingly, genetic crosses can generate \textit{AtRPL10A}, \textit{AtRPL10B} or \textit{AtRPL10C} heterologous double, but not triple, mutants in Arabidopsis (Falcone Ferreyra et al. 2013).

We generated an ‘\textit{AtRPL10} sgRNA+CRISPR/Cas9’ construct in pCAMBIA1300 (Figure 1A). The \textit{AtRPL10} sgRNA consists of an identical 19 nucleotides (ATGTTGGTATGAAGAGGAA) targeting the three genes. However, the protospacer adjacent motif (PAM) is AGG in \textit{AtRPL10B} and \textit{AtRPL10C}, but GGG in \textit{AtRPL10A} (Figure 1B). \textit{A. thaliana} ecotype Col-0 was transformed with the binary vector via the floral dip method (Supplemental Materials and Methods). Four independent Line7, Line9, Line10 and Line11 were created. Transgenic T1 plants from Lines7, 9 and 10 showed severe growth retardation and delayed flowering whilst Line11 had slightly weaker growth compared to the wild-type Col-0 plants (Figure 1C, D). These lines
showed similar phenotypes to *AtRPL10B* RNAi plants, but differed from *AtRPL10A*
T-DNA insertion mutants or *AtRPL10C* knockout plants. To detect potential
multigene-editing events in these transgenic lines, we first analyzed the
sgRNA-targeted sequences using a high-fidelity PCR-RFLP (restriction fragment
length polymorphism) assay. An *EarI* site is located 4-9 nucleotides upstream of the
*AtRPL10* sgRNA PAM sequence (Figure 1B), the region in which
CRISPR/Cas9-mediated DSBs frequently occur (Jinek et al. 2012). We extracted
genomic DNA from transgenic and non-transformed Col-0 plant leaf tissues and
amplified the *AtRPL10* target sequences using gene-specific primers (Table S2).
Incomplete *EarI*-digestion of the resultant PCR products suggests that *AtRPL10A* and
*AtRPL10C* were successfully edited in Line9 (Figure 1E).

To further characterize multigene-editing in these transgenic lines, we cloned the PCR
products into pMD19-T (Supplemental Materials and Methods). Sequencing analyses
showed that nucleotide deletions and/or replacements were introduced into *AtRPL10A*,
*AtRPL10B* and *AtRPL10C* in all transgenic lines (Figures S2-13; Table S3; Dataset
S1). However, the efficiency of multigene-editing of all target sequences was
relatively higher (Figure 1F; Figures S5-7) although varied among *AtRPL10A* (8.8%),
*AtRPL10B* (3.8%) and *AtRPL10C* (23.6%) in Line9 (Table 1). Using an alternative
assay, we identified 7 more (4 deletion and 3 substitution) mutations that were
introduced into *AtRPL10B* in Line9 (Figure 1G-I). In Line7 (Figures S2-4) and
Line10 (Figures S8-10), we detected nucleotide deletions in *AtRPL10A* or *AtRPL10C*
but not in *AtRPL10B*, whilst only point mutations were found in the three *AtRPL10*
genes in Line11 (Figure S11-13). In total we sequenced 1,222 clones and identified 75
different mutations, 37 of which were a deletion of 2 nucleotides. There were single
cases of 1nt or 4nt-deletions, and 36 cases of 1nt-substitution (Table S3). Nevertheless,
multiple deletion and/or point mutations introduced by a single-sgRNA seed-directed
CRISPR/Cas9 were correlated with the abnormal phenotypes in the transgenic lines
(Figure 1C).
Multiplex gene editing through CRISPR/Cas9 that is directed by a number of different sgRNAs has been previously reported in animals and plants (Cong et al. 2013; Feng et al. 2014; Wang et al. 2015; Yan et al. 2015). In this letter, we show that a single-sgRNA seed is capable of guiding CRISPR/Cas9 to edit multiple genes in Arabidopsis. Secondly, we demonstrate that it is possible to use CRISPR/Cas9 technology to create AtRPL10A/B/C triple mutants which otherwise cannot be generated by conventional genetic crossing. Thirdly, we observe that most of mutations resulted from the single-sgRNA seed-guided CRISPR-Cas9 are 2nt-deletion or 1nt-substitution within the sgRNA-target sequences. This differs from a previous report that mutations induced by CRISPR/Cas9 were predominantly 1nt-insertion and short deletions of nucleotides (Feng et al. 2014), but consistent with others (Wang et al. 2015; Yan et al., 2015). Fourthly, the different AtRPL10A/B/C-editing efficiencies (Table 1), particularly in Line9, suggest that chromosomal locations of genes along with the contexts of their surrounding-sequences, heterochromatin architectures and/or DNA/histone methylation may affect the CRISPR/Cas9 system for editing multiple homologous genes (Kleinstiver et al. 2015). Nevertheless, Line9 may prove to be a valuable model to investigate positional effects on the ability of single sgRNA-directed CRISPR/Cas9 to target and edit multiple genes in plants. Lines7, 10 and 11 may be also useful to explore why the single-sgRNA directed CRISPR/Cas9 system preferably causes nucleotide substitution, rather than deletion mutations in target genes. It is interesting to note that all deletion mutations created in our transgenic lines result from removal of 1, 2 or 4 nucleotides, causing frameshifts of the target genes. Compared to conventional multiplex CRISPR/Cas systems (Fu et al. 2013), a single-sgRNA seed has the advantage of reducing off-target gene-editing. This approach is also applicable for the modification of other homologous genes. Moreover, considering how CRISPR/Cas9 recognizes canonical or non-canonical PAMs such as NGG, NGA, NGCG, TTN and YTN (Kleinstiver et al. 2015; Zetsche et al. 2015; Fonfara et al. 2016) as well as how sgRNAs interact with their target sequences (Jinek et al. 2012), it should also be possible to design a single ‘less-stringent’ sgRNA seed that may target less-homologous sequences for
multigene-editing in plants and other organisms.

**ACKNOWLEDGMENTS**

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**AUTHOR CONTRIBUTIONS**

Z.Y. and Y.H. designed experiments; Z.Y. and Q.C. performed all experiments; W.C. and X.Z. analyzed bioinformatics data; J.N., F.M., P.Z., M.Z., X.W. and N.S. performed research. S.J. analysed the data and helped write the paper; Z.Y. and Y.H. wrote the paper.
REFERENCES


Additional Supporting Information is available.

Figure S1. Physical Positions of AtRPL10A, AtRPL10B and AtRPL10C in Arabidopsis Chromosome 1.
AtRPL10A (AT1G14320): 4,888,214 – 4,889,661;
AtRPL10B (AT1G26910): 9,321,650 – 9,322,965;
Chromosome Centromere: 14,899,838 – 14,906,596;

Figure S2. Triple Gene Editing in Line7.
(A) AtRPL10A sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.
(B) Representative chromatograms of AtRPL10A sequences with edited nucleotides.

Figure S3. Triple Gene Editing in Line7.
(A) AtRPL10B sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.
(B) Representative chromatograms of AtRPL10B sequences with edited nucleotides.

Figure S4. Triple Gene Editing in Line7.
(A) AtRPL10C sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.
(B) Representative chromatograms of AtRPL10C sequences with edited nucleotides.

Figure S5. Triple Gene Editing in Line9.
(A) AtRPL10A sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.
(B) Representative chromatograms of AtRPL10A sequences with edited nucleotides.

Figure S6. Triple Gene Editing in Line9.
(A) AtRPL10B sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.
(B) Representative chromatograms of AtRPL10B sequences with edited nucleotides.

Figure S7. Triple Gene Editing in Line9.
(A) AtRPL10C sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.
(B) Representative chromatograms of AtRPL10C sequences with edited nucleotides.

Figure S8. Triple Gene Editing in Line10.
(A) AtRPL10A sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.
(B) Representative chromatograms of AtRPL10A sequences with edited nucleotides.

Figure S9. Triple Gene Editing in Line10.
(A) AtRPL10B sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.
(B) Representative chromatograms of AtRPL10B sequences with edited nucleotides.
are highlighted red.

(B) Representative chromatograms of *AtRPL10B* sequences with edited nucleotides.

Figure S10. Triple Gene Editing in Line10.

(A) *AtRPL10C* sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.

(B) Representative chromatograms of *AtRPL10C* sequences with edited nucleotides.

Figure S11. Triple Gene Editing in Line11.

(A) *AtRPL10A* sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.

(B) Representative chromatograms of *AtRPL10A* sequences with edited nucleotides.

Figure S12. Triple Gene Editing in Line11.

(A) *AtRPL10B* sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.

(B) Representative chromatograms of *AtRPL10B* sequences with edited nucleotides.

Figure S13. Triple Gene Editing in Line11.

(A) *AtRPL10C* sequences of individual clones. PAM sequence is boxed and the edited nucleotides are highlighted red.

(B) Representative chromatograms of *AtRPL10C* sequences with edited nucleotides.

Table S1. Comparisons of AtRPL10 Genes and Their Protein Products

Table S2. Primers Used in This Study

Table S3. Multigene Editing and Their Impacts on Protein Sequences in CRISPR/Cas9 Transgenic Lines

Dataset S1. Sequences of the PCR Products for the Three *AtRPL10* Genes.

(A-B) Restriction fragment length polymorphism (RFLP) analysis of *AtRPL10A* (A), *AtRPL10B* (B) *AtRPL10C* (C). Sequences corresponding to the ‘seed’ sgRNA are indicated in lowercase. The *Earl* digestion site (\(|\)\) is indicated and its recognition sequence is underlined.

**FIGURE LEGEND**

Figure 1. A Single sgRNA Seed Directs CRISPR/Cas9 to Simultaneously Edit Three *AtRPL10* Homologous Genes.

(A) Schematic of the single sgRNA seed and CRISPR/Cas9 construct in the binary vector pCAMBIA1300. Nucleotides corresponding to the sgRNA seed sequence are underlined. The *AtU6-26* promoter (arrow), sgRNA and the scaffold, enhanced 35S promoter (arrow), NLS (nuclear localization signal)-tagged Cas9, hygromycin (HYG) as well as the right and left borders (RB and LB) in the binary vector are indicated.

(B) Comparison of the sgRNA seed-targeted *AtRPL10* gene sequences. The *Earl* site is
underlined. The PAM sequences are highlighted red. Nucleotide coordinates are indicated.

(C) Phenotypes of transgenic plants of four independent lines. Bar = 3cm in Line7, Line9, Line10 and Line11. Bar = 5cm in Col-0.

(D) Confirmation of plant transformation. The Cas9 gene was detected in four transgenic lines as indicated. A BM2000 DNA ladder (Marker) as well as the size and position of the Cas9 transgene PCR fragment are indicated.

(E) PCR-RFLP assay of multiple gene-editing in four transgenic lines. Gene-specific PCR products were digested with EarI. Incomplete digestion shows three clear bands, indicating that successful editing of \( AtRPL10A \) and \( AtRPL10C \) in Line9. A BM2000 DNA ladder (Marker) was included in gel electrophoresis.

(F) Sequencing analysis of multiple gene-editing in Line9. Representative sequencings show indels in \( AtRPL10A \), \( AtRPL10B \) and \( AtRPL10C \). The sgRNA target sequences are shown in lowercase.

(G-I) PCR-RFLP and sequencing assays of \( AtRPL10B \) editing in Line9. After EarI digestion, residual DNA in the position of the red-box was extracted from the agarose gel (G) and subcloned for sequencing analysis (H). A BM2000 DNA ladder (M) was included in gel electrophoresis. Sequences of 19 individual clones for \( AtRPL10B \) were aligned, and mutations with two nucleotide-deletion (red arrow) or single nucleotide-substitution (highlighted red) are indicated (I). RD stands for restriction endonuclease digestion.

### Table 1. Summary of Multigene Editing Efficiency*

<table>
<thead>
<tr>
<th>Transgenic Lines</th>
<th>( AtRPL10A )</th>
<th>( AtRPL10B )</th>
<th>( AtRPL10C )</th>
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<tbody>
<tr>
<td></td>
<td>Deletion</td>
<td>Point Mutation</td>
<td>Deletion</td>
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<td>0/99</td>
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<tr>
<td>Line 11</td>
<td>0/93</td>
<td>2/93</td>
<td>0/109</td>
</tr>
</tbody>
</table>

*The number of CRISPR/Cas9 edited sequences (clones) out of the total number of sequenced samples for the three \( AtRPL10 \) genes in each of the transgenic lines.