

SHORT COMMUNICATION

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Genetic characterization of T-DNA insertions in the genome of the *Arabidopsis thaliana* *sumo1/2* knock-down line

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ABSTRACT

Sumoylation is an essential post-translational modification in *Arabidopsis thaliana*, which entails the conjugation of the SUMO protein onto lysine residues in target proteins. In *Arabidopsis*, 2 closely related genes, *SUMO1* and *SUMO2*, act redundantly and are in combination essential for plant development, i.e. the combined loss of *SUMO1* and *SUMO2* results in embryo-lethality. To circumvent this lethality, *SUMO2* was previously knocked down in a *sumo1* knockout background by expressing an artificial microRNA that targets *SUMO2* (*amiR-SUMO2*). This *sumo1/2^{KD}* line with low *SUMO2* levels represents a valuable genetics tool to investigate SUMO function in *planta*. Here, we re-sequenced the whole-genome of this *sumo1/2^{KD}* line and identified 2 *amiR-SUMO2* insertions in this line, which were confirmed by PCR-genotyping. Identification of these 2 insertions enables genetics with this tool.

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Arabidopsis; SUMO; T-DNA insertions; whole genome resequencing

Introduction

Sumoylation is a post-translational modification resulting in conjugation of SUMO (Small Ubiquitin-like Modifier) proteins onto targets through the side chain of lysine residues. SUMO is encoded by a single copy gene in many eukaryotes like budding yeast (*Saccharomyces cerevisiae*), *Caenorhabditis elegans* and fruit fly (*Drosophila melanogaster*).¹ In contrast to these species, the genome of *Arabidopsis* (*Arabidopsis thaliana*) contains 8 *SUMO* genes. *SUMO1* and *SUMO2* are the main isoforms used for sumoylation.^{2–5} They act redundantly and are essential in *Arabidopsis*, as both the *sumo1–1* and *sumo2–1* single null mutants do not display any aberrant development phenotype, while the corresponding double mutant is embryo-lethal.⁴ To understand the function of sumoylation in *planta*, we created a transgenic line where *SUMO1* is knocked out (KO) and *SUMO2* knocked down (^{KD}).⁶ These lines were obtained by crossing the *sumo1–1* null mutant with the *SUMO2^{KD}* line B, a line silenced for *SUMO2* using an artificial microRNA (*amiR*) targeting *SUMO2* transcripts (*amiR-SUMO2*); this *amiR-SUMO2* was engineered according to the instructions of WMD MicroRNA Designer: <http://wmd3.weigelworld.org/cgi-bin/webapp.cgi>.⁷ The *sumo1–1 SUMO2^{KD}* mutant (hereafter called *sumo1/2^{KD}*) displays a strong phenotype characterized by enhanced accumulation of salicylic acid (SA), accumulation of the Pathogenesis-Related proteins 1 and 2 (PR1/2), spontaneous cell death in leaves, early flowering,

partial sterility and a dwarf stature.⁶ Although *SUMO2* conjugation levels are strongly suppressed in this line, the low levels of *SUMO2* protein are apparently sufficient to maintain plant viability.

As the insertion site of the *amiR-SUMO2* construct is unknown, genotyping of ‘*SUMO2^{KD}* allele’ was till now based on the assessment of the presence of the *amiR-SUMO2* construct using PCR and on segregation for kanamycin-resistance in seedlings (the plant selection marker that was co-integrated with the *amiR-SUMO2* construct). Genetics with *sumo1/2^{KD}* is, therefore, tedious: homozygous lines can only be found by examining the segregation for kanamycin resistance in the next generation.

While out-crossing the *sumo1/2^{KD}* line to different mutant backgrounds, we noted that the dwarf phenotype segregated in the resulting F2 generation, albeit the F2 plants were genotyped as homozygous for the *sumo1–1* and *amiR-SUMO2* alleles. As stable transformation of *Arabidopsis* can result in multiple T-DNA insertions,⁸ we reasoned that the original *sumo1/2^{KD}* line might contain multiple *amiR-SUMO2* integration sites. Variation in the number of insertions potentially leads to different *SUMO2* silencing levels and could explain the heterogeneous phenotype of the F2 progenies. Identification of the insertion sites is, therefore, needed for reverse genetics with the original *sumo1/2^{KD}* line. Here we report on the *amiR-SUMO2* insertion locations in the genome of the

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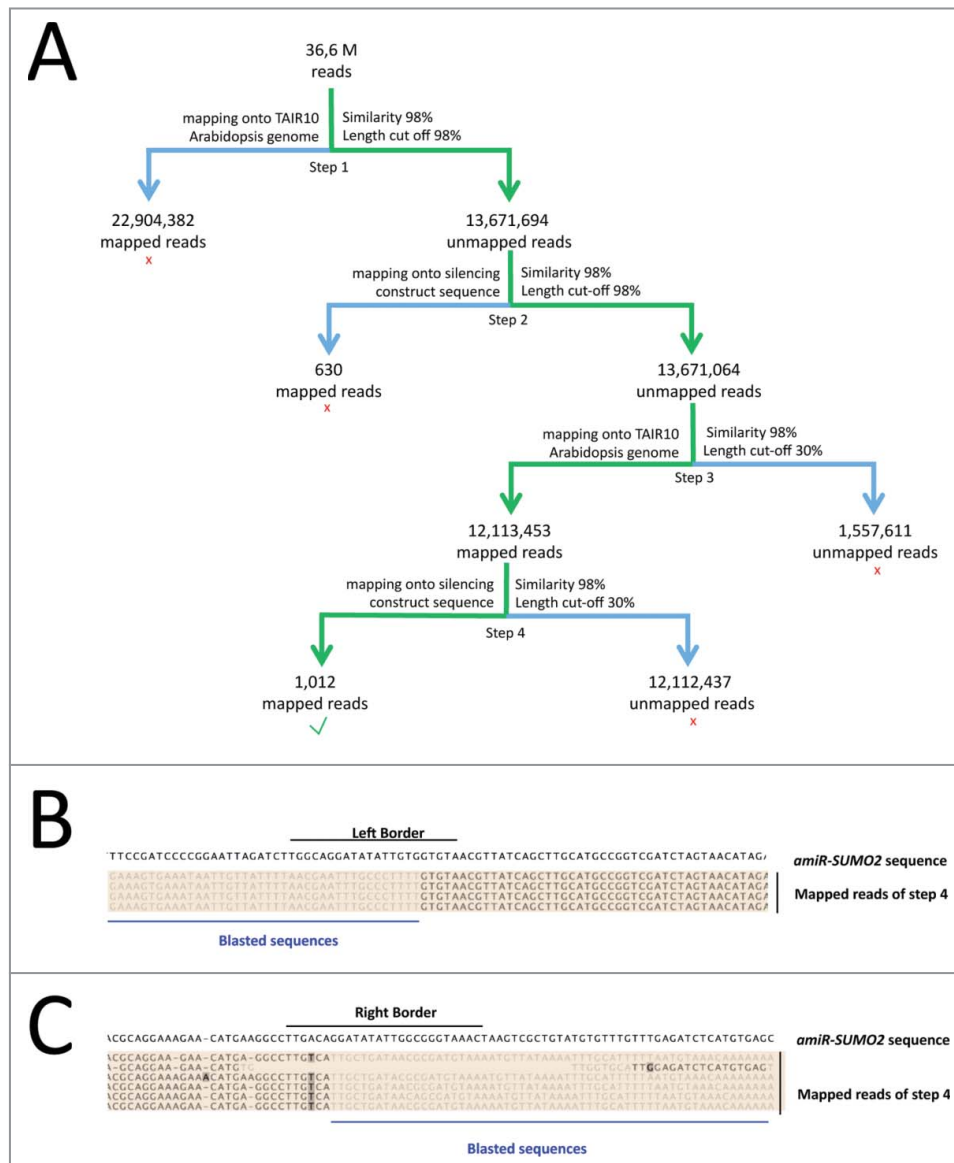


Figure 1. Identification of the 2 *amiR-SUMO2* insertion sites using NGS sequencing. (A) Pipeline used to identify the T-DNA insertion sites. (Step 1) Reads were mapped onto the Arabidopsis genome assembly (TAIR10), using a sequence similarity cut-off of > 98% and read length cut-off of > 98% sequence overlap. (Step 2) To remove the reads that fully matched to the *amiR-SUMO2* construct, the unmapped reads were then mapped to the *amiR-SUMO2* construct using similar parameters. (Step 3) We then selected in the remaining set of unmapped reads, the reads that partially mapped to the Arabidopsis genome, using >98% similarity and a length cut-off of > 30%. (Step 4) The retained reads (from Step 3) were then mapped to the *amiR-SUMO2* construct, with >98% similarity and a length cut-off of > 30% to obtain the reads that map across single integration site boundaries with at least 30 bp. We found 1,012 reads, which mapped to 2 different genomic sites. The insertions were identified by blast searches with these latter reads using the part of the reads that did not map onto the *amiR-SUMO2* construct. (B) and (C) Visualization of the mapped reads of Step 4 from panel A (shown on color background) on the *SUMO2* silencing construct sequence at the Left Border (B) and Right Border (C). Within the mapped reads of step 4, black sequences indicates the regions of the reads that map onto the *amiR-SUMO2* construct, while gray sequences indicate the regions of the reads from Step 4 that do not map on the *amiR-SUMO2* construct.

sumo1/2^{KD} line B21, which facilitates classical genetics with this line. We have identified both the number of insertions and their localization based on whole genome re-sequencing of the *sumo1/2^{KD}* line using next-generation sequencing. By mapping the generated sequencing reads, we found 2 genomic insertions. Using PCR-based genotyping, we could confirm the location of both insertions in the *sumo1/2^{KD}* line. Using these PCR primers, the presence of both *amiR-SUMO2* insertions can now be quickly assessed in the offspring of out-crosses with this *sumo1/2^{KD}* line.

Materials and methods

Genomic DNA extraction, re-sequencing and short read mapping

We isolated gDNA from pools of seedlings of *sumo1/2^{KD}* from van den Burg et al.⁶ using the Nucleospin II plant kit (Macherey-Nagel). The gDNA isolation yielded 38.8 ng/uL, with A260/280 ratios of 1.87 and A260/230 ratios of 2.47. The gDNA was sequenced according to the manufacturer instructions on the Ion Torrent platform (ThermoFischer). The obtained short sequencing reads (average length 150 bp)

were then mapped onto both Arabidopsis genomic sequence (TAIR10) and the *amiR-SUMO2* plasmid using the CLC workbench v6.5 software by applying the strategy outlined in Fig. 1A. The parameters used for mapping were: miss match cost = 2; insertion cost = 3; deletion cost = 3.

Primer design and PCR genotyping

PCR genotyping was performed on *sumo1-1*, *SUMO2^{KD}* line B (*i.e.*, parental lines), and 2 *sumo1/2^{KD}* lines: *sumo1/2^{KD}* line B21 and *sumo1/2^{KD}* line B22#1. Both are lines obtained from the same cross between *sumo1-1* and *SUMO2^{KD}*. Primer sequences and primer combinations used for genotyping are summarized in Table 1. *SUMO1* genotyping was done with the primers 3039 and 6541 for the wild type *SUMO1* (*SUMO1^{WT}*) allele, and primers 6541 and 3249 for *sumo1-1*. *PFK7* genotyping was done with primers 4904 and 4980 for *PFK7* wild type (*PFK7^{WT}*), 4904 and 4719 for *amiR-SUMO2* in *PFK7* (*PFK7^{amiR-SUMO2}*). *proCYP98A3* genotyping was done with the primers 5733 and 5578 for *proCYP98A3* wild type (*proCYP98A3^{WT}*), 5733 and 4714 for *amiR-SUMO2* in *proCYP98A3* (*proCYP98A3^{amiR-SUMO2}*). The fragments were amplified using a touch-down PCR (35 cycles): (i) a melting temperature of 95°C for 30s, (ii) an annealing temp of 60°C with -1°C each cycle for 10 cycles and then 25 additional cycles at 50°C, (iii) an elongation time of 1m 15s at 72°C, and back to (i).

Results

After extraction of the gDNA from the *sumo1/2^{KD}* line B21 used in van den Burg *et al.*⁶ samples were sequenced using

Table 1. (A) Primer combinations used for genotyping of the different alleles and (B) sequences of the primers used for PCR genotyping.

A		
Locus	Allele	primer name
<i>SUMO1</i>	<i>SUMO1^{WT}</i>	3039
	<i>sumo1-1</i>	6541
<i>PFK7</i>	<i>PFK7^{WT}</i>	4904
	<i>PFK7^{amiR-SUMO2}</i>	4980
		4719
<i>proCYP98A3</i>	<i>proCYP98A3^{WT}</i>	5733
	<i>proCYP98A3^{amiR-SUMO2}</i>	5578
		4714
B		
primer name	primer sequence (5' to 3')	
3039	TCTGCAAACAGGAGGAAG	
4714	CATTAAATGAATCGGCCAACGCGCG	
4719	TCGCCCTTCTGACGAGTTCTTCTGA	
4904	AGTTTCTTGGGCCTAAGGATACA	
4980	AGTGTGAAAAACATATACAAGAAC	
5578	CACCGCTATTAGAAACACGAC	
5733	CAGCAGACGAAACCAACAACACT	
6249	TGGTTCACGTAGTGGCCATCG	
6541	TAGGATCCGATACCAACGACAA	

next-generation sequencing. We obtained 36.6 M reads with a median length of 177 bp. To localize the genome insertion sites of the *amiR-SUMO2* T-DNA, we identified the reads that partially (*i.e.*, >30%) mapped to both the Arabidopsis genome and

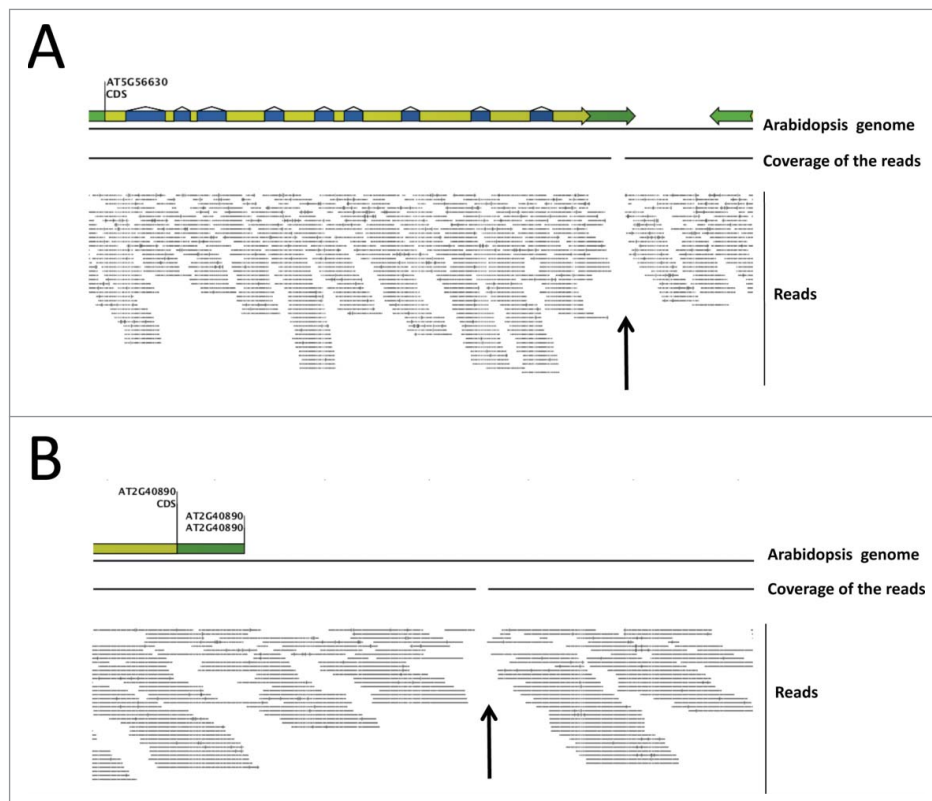


Figure 2. The *SUMO2* silencing construct is homozygous at both the *PFK7* (AT5G56630) and *proCYP98A3* (AT2G40890) integration site in the *sumo1/2^{KD}* line (B21). The 36.6 M short sequencing reads were mapped onto the Arabidopsis genome assembly (TAIR10), with a similarity match of > 98% and length cut-off of > 98%. Visualization of the reads on (A) *PFK7* genomic and (B) *CYP98A3* promoter (*proCYP98A3*) sequences shows the gap in read coverage in the 2 identified insertion sites (black arrows).

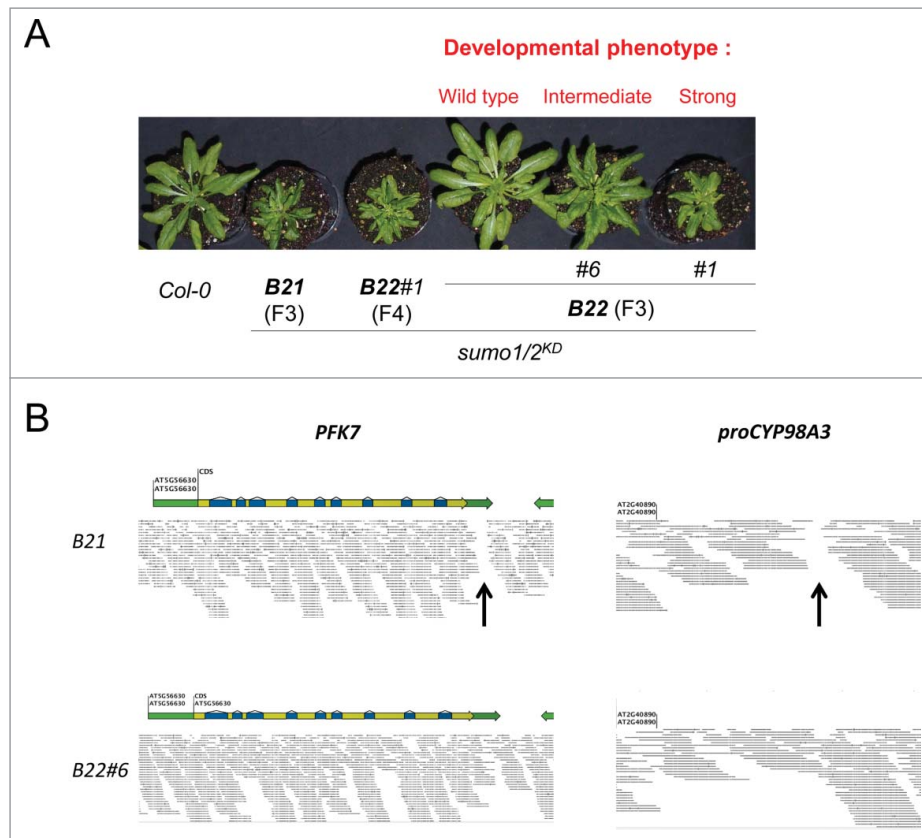


Figure 3. Both the *amiR-SUMO2* integration events (*i.e.*, at *PFK7* and *proCYP98A3*) need to be present in a homozygous state to obtain a strong *sumo1/2^{KD}* phenotype. (A) Seedlings from the self-pollination offspring of *sumo1/2^{KD}* lines B21 (F3 generation); B22 (F3 generation); and B22#1 (F4 generations) were grown for 4 weeks in short day conditions (11 hours in day light / 13 hours in dark). The lines B21 and B22 represent 2 sister plants obtained from the same cross between *sumo1-1* and *SUMO2^{KD}* line B; B22#1 and B22#6 exemplify selfings from the plant B22. Whereas the strong developmental phenotype did not segregate for the progeny of B21, the progeny of B22 displayed 3 different phenotypes: normal, intermediate, and strong (*i.e.*, ‘B21-like’). A set of B22#6 intermediate plants was pooled and their combined gDNA was sequenced using next-generation sequencing. (B) Visualization of the reads obtained from re-sequencing of the pool of the B22#6 at the *PFK7* and *proCYP98A3* promoter (*proCYP98A3*) genomic regions reveals no gap in the read coverage for both insertion sites (bottom row), while a gap in the read coverage is visible for both genomic regions in the case of the B21 plant (top row, black arrows).

the *amiR-SUMO2* construct (Fig. 1). Briefly, we first removed the reads that align for >98% with either the Arabidopsis genome or with the *amiR-SUMO2* construct. From the remaining reads, we then selected the reads that partially mapped onto the *amiR-SUMO2* construct (minimum overlap of 30%). These reads were then mapped onto the Arabidopsis genome allowing again a min. match of 30%. With this method 1,012 reads remained (Fig. 1A). Some of these reads partially mapped onto the *amiR-SUMO2* construct at the Left (LB) or Right Border (RB) sequence. The sequence fragment of these latter reads, which could not be mapped to the plasmid, was then blasted to NCBI to identify their location in the Arabidopsis genome. Considering the reads that mapped onto the LB (Fig. 1B) or RB (Fig. 1C) of the *amiR-SUMO2* construct, we identified 2 locations: (i) the 3’UTR of *PFK7* (AT5G56630), a gene coding for PHOSPHOFRUCTOKINASE 7 located on chromosome 5, and (ii) the promoter region of *CYP98A3* (*proCYP98A3*; AT2G40890) located on chromosome 2. Our *sumo1/2^{KD}* line contains, therefore, 2 different insertions located on 2 different chromosomes. Apparently, both T-DNA integration events were retained after crossing with *sumo1-1*. The first site is 587 bp upstream of the start codon (-578) of the gene *CYP98A3*, while the second site is 2,531 bp downstream of the start codon (+2,531) of the gene *PFK7*.

The 36,6 M reads were then mapped onto the TAIR10 Arabidopsis genome assembly with a similarity of 98% and a length cut-off of 98%. When we visualized the reads onto the Arabidopsis genome using CLC workbench (Figs. 2A and B), we observed a gap in the read coverage (black arrows) exactly at the expected *PFK7* and *proCYP98A3* insertion locations, while the surrounding gDNA is nicely covered by reads. Both insertions are, therefore, present in a homozygous state in the previously reported *sumo1/2^{KD}* line.⁶

The original cross between *sumo1-1* and *SUMO2^{KD}* yielded a F2 population, which included 2 F2 sister plants: B21 (the original line detailed in van den Burg *et al.*⁶) and B22. Whereas selfings of B21 show no segregation for the reported strong ‘*sumo1/2^{KD}* developmental’ phenotype, selfings of the B22 F2 plant (*i.e.*, F3 generation) displayed 3 different phenotypes: normal plants, plants with an intermediate rosette size (*e.g.*, B22#6) and ‘B21-like’ plants (*e.g.*, B22#1) (Fig. 3A). The developmental phenotype of B22#1 proofed to be stable in the next generation. However, the phenotype of B22#6 continued segregating in normal, intermediate, and ‘B21-like’ plants. Using next-generation sequencing, we then also sequenced a pool of the B26#6 progeny that all had the intermediate phenotype. Using our bioinformatics pipeline, we could confirm

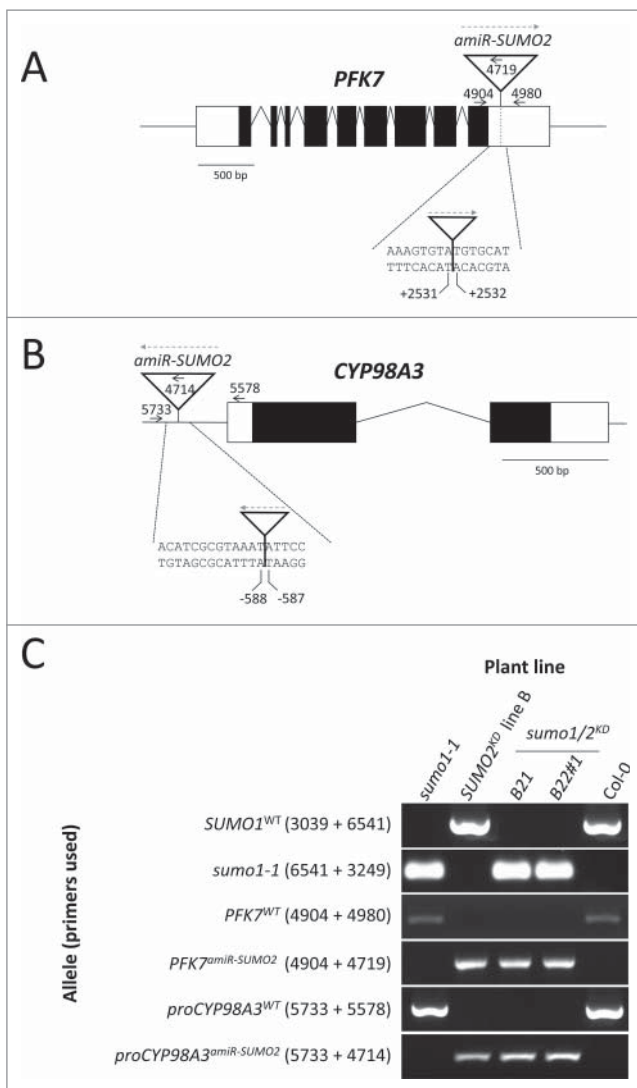


Figure 4. PCR-based genotyping of the *PFK7* and *proCYP98A3* alleles in the *sumo1/2^{KD}* line. A and B. True to scale diagrams of the *PFK7* and *CYP98A3* genes. The *amiR-SUMO2* integration sites (arrowheads) are located in (A) *PFK7* at +2,531 bp and (B) in *CYP98A3* -587 bp, calculated from the start codon (+1). The exons and introns are presented by boxes and broken lines, respectively. The white boxes reflect the 5'- and 3'-untranslated regions, while the black boxes refer to the coding regions. The primers used for genotyping are indicated by their ID numbers given (not to scale; see also Table 1). The orientation of the *amiR-SUMO2* constructs is indicated using gray-dash arrows (from the Left Border to the Right Border). (C) PCR-based genotyping using the primers represented in (A) and (B) of the *SUMO2^{KD}* line B and *sumo1/2^{KD}* lines B21 and B22#1, 2 lines obtain from the same cross between *sumo1-1* and *SUMO2^{KD}* line B. See also Table 1 for primer sequences.

the presence of the silencing construct at both integration sites (*PFK7* and *proCYP98A3*), meaning that both insertions were still present in the parental plant B22#6. The obtained sequencing reads were then mapped onto the TAIR10 Arabidopsis genome assembly with a similarity of 98% and a length cut-off of 98%. Upon visualization of the reads onto the Arabidopsis genome, we found individual reads that span across either of the 2 insertion sites, meaning that both insertions were still heterozygous in B22#6 (Fig. 3B). Combined with the result of outcrossing the B21 line, we conclude that both *amiR-SUMO2* integration events need to be present in a homozygous state to

obtain a strong developmental phenotype as seen with the *SUMO1/2^{KD}* B21 line.

Subsequently, we developed primers to genotype for both insertion sites. These primer pairs either (i) amplify the genomic region surrounding the T-DNA integration site or (ii) amplify a fragment that encompasses both the *amiR-SUMO2* T-DNA and the flanking genomic region (Fig. 4A; B and Table 1). Using the primer pairs 4904+4980, 4904+4719, 5733+5578 and 5733+4714, we could genotype for the *PFK7^{WT}*, *PFK7^{amiR-SUMO2}*, *proCYP98A3^{WT}* and *proCYP98A3^{amiR-SUMO2}* alleles. Using these primers, we then confirmed our next generation sequencing result that both insertion sites are homozygous in the 2 *sumo1/2^{KD}* lines: *sumo1/2^{KD}* B21 and *sumo1/2^{KD}* B22#1 (Fig. 4C).

Discussion

Traditionally, Southern-blotting is used to reveal the number of T-DNA insertions, while TAIL-PCR (Thermal asymmetric interlaced-PCR) is used to identify the integration sites.⁹ However, TAIL-PCR does not guarantee identification of all integration sites. Here, we identified by whole genome re-sequencing followed by the mapping of sequencing reads using bioinformatics that the *sumo1/2^{KD}* line contains 2 *amiR-SUMO2* constructs, and we identified their exact genomic locations. Finally, we established a PCR-based genotyping approach for both *amiR-SUMO2* integration sites. Knowing that the cost of whole genome (re-)sequencing has dramatically decreased over the last years, this constitutes a powerful and rapid method to localize and genotype T-DNA insertions in transgenic lines of e.g., Arabidopsis.

Prior this study, the genotyping of the *sumo1/2^{KD}* line relied on the assessment of the presence of the *amiR-SUMO2* construct by PCR, and on the segregation for the kanamycin-resistance, which was co-integrated with the *amiR-SUMO2* construct. Repeatedly, we observed intermediate phenotypes (similar to the intermediate phenotype observed in the line B22#6) for the *sumo1/2^{KD}* transgenic plants (curled leaves, reduced rosette size) when we outcrossed this mutant to other genetic backgrounds,⁶ despite the fact that the F2 plants were found to be homozygous for both *sumo1-1* and kanamycin-resistance and that they contained the *amiR-SUMO2* construct. Hence, the silencing of *SUMO2* does not only show semi-dominance, but we also noted that lines homozygous for kanamycin-resistance segregated for the morphological phenotype. The here presented data indicates that both *amiR-SUMO2* insertions contribute to the original *sumo1/2^{KD}* phenotype. Genetics with the *sumo1/2^{KD}* line must consequently take into consideration that both *amiR-SUMO2* integrations (at the *PFK7* and *proCYP98A3* loci) are needed for phenotypic comparisons when using this line.

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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