Rapid Identification of a Natural Knockout Allele of ARMADILLO REPEAT-CONTAINING KINESIN1

That Causes Root Hair Branching by Mapping-By-Sequencing^{1[C][W]}

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In Arabidopsis (*Arabidopsis thaliana*), branched root hairs are an indicator of defects in root hair tip growth. Among 62 accessions, one accession (Heiligkreuztal2 [HKT2.4]) displayed branched root hairs, suggesting that this accession carries a mutation in a gene of importance for tip growth. We determined 200- to 300-kb mapping intervals using a mapping-by-sequencing approach of F2 pools from crossings of HKT2.4 with three different accessions. The intersection of these mapping intervals was 80 kb in size featuring not more than 36 HKT2.4-specific single nucleotide polymorphisms, only two of which changed the coding potential of genes. Among them, we identified the causative single nucleotide polymorphism changing a splicing site in *ARMADILLO REPEAT-CONTAINING KINESIN1*. The applied strategies have the potential to complement statistical methods in high-throughput phenotyping studies using different natural accessions to identify causative genes for distinct phenotypes represented by only one or a few accessions.

Root hairs are tubular tip outgrowths of single root epidermal cells (trichoblasts). They are an excellent genetic system and serve as a model to study the molecular components regulating tip growth (Carol and Dolan, 2002; Samaj et al., 2004; Lee and Yang, 2008). One of the main regulators of tip growth in root hairs is the small G protein RHO OF PLANTS2 (ROP2; Jones et al., 2002; Payne and Grierson, 2009). ROP2 determines the position of root hairs in incipient epidermal root hair cells and remains localized in the emerging tip during root hair tip growth (Molendijk et al., 2001; Jones et al., 2002). In addition, other factors have been identified to be important for growth and its directionality including phosphoinositides, cytoplasmic [Ca²⁺] gradients and their

oscillation, reactive oxygen species, the RAB GTPase homolog A4B, and the cytoskeleton (Foreman et al., 2003; Preuss et al., 2004, 2006; Carol et al., 2005; Thole et al., 2008; Heilmann, 2009).

Defects in essential processes for the establishment and maintenance of tip growth lead to deviations in root hair morphology such as branching and waviness (Samaj et al., 2004; Lee and Yang, 2008). Both the microtubules (MTs) and actin are important regulators of tip growth with MTs maintaining one growth point (Bibikova et al., 1999; Miller et al., 1999; Baluska et al., 2000). The latter is evident from the finding that artificially induced [Ca²⁺] gradients can induce additional growth tips when MTs are destroyed by drug treatments (Bibikova et al., 1999).

Although the genetic and molecular analysis revealed a well-understood working model for root hair growth, little is known about the natural variation of the underlying processes. Which are the adaptive processes of relevance to specific environmental cues and which have already been selected for in natural accessions? One way to address this question is to link genotype and phenotype by association mapping using various Arabidopsis (Arabidopsis thaliana) accessions. This is greatly facilitated by the 1001 Genomes Project (http://1001genomes.org), providing an increasing number of sequenced accessions. In some cases, phenotypes are only found in one or a few accessions. When the minor allele frequency (AF) is low, the identification of such rare causative alleles with genome-wide association mapping studies is challenging because they cannot be discriminated from false positives (e.g. sequencing errors or synthetic

tip growth with MTs maintaining one growth (Molendijk et al., 2001; Jones et al., 2002).

addition, other factors have been identified to be apportant for growth and its directionality including hosphoinositides, cytoplasmic [Ca²⁺] gradients and their signs when MTs are destroyed by drug

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L.R. and A.S. performed the experiments; H.S. performed the mapping-by-sequencing analysis; A.S. performed the accession-based SNP extraction and data analysis; A.S., L.R., M.H., and K.S. designed the experiments; A.S., M.H., and L.R. wrote the article.

^[C] Some figures in this article are displayed in color online but in black and white in the print edition.

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associations; Korte and Farlow, 2013), they are not detectable because of chosen thresholds, or they do not support a statistically significant value (Cantor et al., 2010). Other time-consuming approaches with low mapping resolution, such as quantitative trait loci mapping, need to be followed to identify the causative gene. For this, mapping-by-sequencing, which was originally developed for the identification of mutagen-induced changes in model species (Schneeberger et al., 2009b), can help to rapidly identify causal polymorphisms including nonmodel and nonreference strains (Nordström et al., 2013; Takagi et al., 2013). The resolution of mapping-by-sequencing experiments in Arabidopsis mapping populations is typically between multiple hundreds of kilobase pairs up to a few megabase pairs (James et al., 2013). Although intervals of this size allow the identification of causal mutations in forward genetic screens, they are problematic for the analysis of diverse Arabidopsis accessions because the single nucleotide polymorphism (SNP) density is very high; consequently, hundreds of polymorphisms have to be considered.

Here we report on a modification of the mapping-bysequencing strategy providing a shortcut from distinct, monogenic accession-specific phenotypes to the causative SNP. When studying root hair morphology in 62 accessions for which the genome sequences were released by the 1001 Genomes Project (Cao et al., 2011), we found one accession (Heiligkreuztal2 [HKT2.4]) in which almost all root hairs were branched. To identify the causative gene, we used an approach based on mapping-by-sequencing. Instead of one outcross, we used outcrosses with three different accessions. We selected F2 seedlings exhibiting the distinct, monogenic, recessive root hair branching phenotype for sequencing. Combining the intersection of the three resulting mapping intervals with a selection for accession-specific SNPs revealed two primary candidate genes responsible for the root phenotype. We demonstrate that the causative SNP renders a splicing site in ARMADILLO REPEAT-CONTAINING KINESIN1 (ARK1) inactive and therefore leads to a defective ARK1/MORPHOGENESIS OF ROOT HAIR2 (MRH2) protein that is thought to coordinate actin microfilaments and MTs during tip growth of root hairs (Yang et al., 2007; Yoo and Blancaflor, 2013).

RESULTS AND DISCUSSION

HKT2.4 Has a Root Hair Branching Phenotype

In this study, we analyzed the root hair morphology of 62 Arabidopsis accessions in the root hair patterning zone. Among the analyzed accessions, only HKT2.4 that originated from the area around Tuebingen, Germany, showed branched root hairs (Fig. 1; Supplemental Table S1).

To determine whether root hair branching is caused by one or multiple loci, we performed segregation analyses. We chose the Arabidopsis ecotypes Columbia-0 (Col-0), Landsberg *erecta* (Ler), and Wassilewskija-2 (Ws-2)

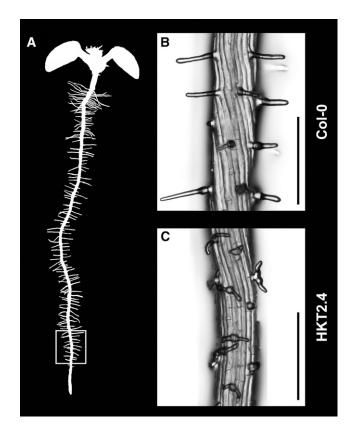


Figure 1. Root hair morphology in Col-0 and HKT2.4 plants. A, Scheme of Arabidopsis roots. The analyzed region is marked with a white square. This region is named the lower region in the text. B, Root hairs of Col-0 plants are tubular. C, Root hairs of HKT2.4 are wavy and branched. Scale bars = $250~\mu m$.

as crossing partners because various root hair mutants have been described in these backgrounds, thereby enabling a direct phenotypic comparison in subsequent experiments. First, we determined the percentage of branched root hairs in all four accessions in the lower zone of the root (Fig. 1A). Whereas almost all root hairs were branched in HKT2.4 (96.8% \pm 4.1%), less than 5% were branched in Col-0 and Ws-2 and 10.9% \pm 4.1% were branched in Ler (Supplemental Table S2). In all F1 seedlings from all three crosses, we observed an intermediate root hair branching phenotype in the upper region of the root (Supplemental Fig. S1). In this region, F2 seedlings segregated in a 1:2:1 ratio (n > 100; Table I) for Col-0/Ler/Ws-2-like, intermediate, and HKT2.4-like phenotypes, indicating semidominance for this phenotype (Col-0, 0.8:2.3:0.9; Ler, 0.9:2.1:1; and Ws-2, 0.9:2.1:1). Because all seedlings with an intermediate root hair branching phenotype in the upper region showed no branching in the lower region, a 3:1 segregation ratio for nonbranching versus branching could be deduced for the lower region.

Taken together, the genetic data indicate that the distinct HKT2.4 root hair branching phenotype in the lower region is monogenetic and therefore accessible to a mapping approach using Col-0, Ler, and Ws-2.

Table 1. Segregation of the root hair branching phenotype in F2 seedlings

For the segregation analysis, root hairs along the whole root were analyzed. The intermediate branching phenotype was only observed in the upper zone (below the embryonic root hairs; Supplemental Fig. S1).

Branching Phenotype	Col-0 × HKT2.4	Ler × HKT2.4	Ws-2 × HKT2.4
Col-0-/Ler-/Ws-2-like	19	29	31
Intermediate	58	51	76
HKT2.4-like	23	26	35
HKT2.4-like (%)	23.0	24.5	24.7

Study Design Aiming at a High Mapping Resolution

Because quantitative trait locus mapping is a powerful but time-consuming method with low mapping resolution, we searched for mapping alternatives to unravel the underlying mutation of the HKT2.4-specific root hair branching phenotype. Mapping-by-sequencing combining bulk segregant analyses with next-generation sequencing is one recently established method for rapid ethyl methanesulfonate mutant identification in Arabidopsis, *Arabis alpina*, and other species (Schneeberger et al., 2009b; Hartwig et al., 2012; Nordström et al., 2013).

Simulations of mapping-by-sequencing experiments using mapping populations generated by crossing Arabidopsis Ler and Col-0 revealed relevant criteria for the experimental design (James et al., 2013). On the basis of these simulations, four main aspects were considered.

First, we decided to use paired-end sequencing because this is expected to result in an increased fraction of informative read alignments (James et al., 2013). Second, because 1% to 2% of false-scored plants can lead to 82% to 145% more candidate SNPs in the simulation (James et al., 2013) and given that roots exhibit high phenotypical plasticity (Müller and Schmidt, 2004), we performed pre-experiments and segregation analysis to ensure precise phenotyping. Third, because the coverage and the number of recombinants are correlated and both have an impact on the size of the mapping interval (James et al., 2013), we aimed for a 25-fold coverage. In addition, we phenotypically selected about 200 seedlings from the F2 mapping population. This strategy is expected to define a 400-kb mapping interval (James et al., 2013). Fourth, we used three populations to take advantage of different recombination hotspots in different accessions (Drouaud et al., 2006, 2007; Kim et al., 2007; Salomé et al., 2012) and to increase the number of recombinants.

Identification of an 80-kb Mapping Interval

In light of the above-mentioned four aspects, we followed the procedure summarized in Supplemental Figure S2, whereby HKT2.4 was crossed to each of Col-0, Ler, and Ws-2, respectively. Because the success of a particular cross can depend on which accession was used as a mother (Autran et al., 2011; Fitz Gerald et al., 2014), reciprocal crosses were performed. F2 seedlings with HKT2.4-specific phenotypes were selected from all three segregating populations and pooled for both directions of the crossings. These seedlings were expected to carry the causative SNP and to vary in other SNPs due to recombination events. This resulted in F2 pools with at least 199 seedlings each (Table II). The parental and bulked segregants' genomes were whole-genome sequenced with a minimum of 25-fold mean coverage, resulting in at least 110.39 million positions with sufficient and unique sequencing information. More than 95% of the reads were mapped (Table II; Supplemental Fig. S3).

For all three crossings, we used the resequencing information of the parental lines to generate genome-wide lists of markers used for the estimation of AFs with SHOREmap software. Average AFs, as assessed in sliding windows, and SHOREmap-derived boost values revealed linkage between a region on chromosome 3 and the phenotype, thereby providing evidence that chromosome 3 harbors the causative locus (Fig. 2, A–C; Supplemental Figs. S4 and S5). The corresponding mapping intervals had sizes of 200 kb (crossings with Col-0 and Ler) and 300 kb (crossing with Ws-2) but their intersection revealed a final mapping interval of not more than 80 kb between 20.30 and 20.38 megabase pairs on chromosome 3 (Fig. 2D).

To summarize, mapping-by-sequencing can be used to map Arabidopsis accession-specific phenotypes. In this study, our strategy outperformed the prediction and reduced the mapping interval to 80 kb. This suggests

 Table II. Summary of genome resequencing

N.A., Not available.							
Sample and Resequencing Information	HKT2.4	Col-0	Ler	Ws-2	HKT2.4 \times Col-0	HKT2.4 \times Ler	HKT2.4 \times Ws-2
No. of pooled F2 seedlings	200	200	200	200	205	214	199
Coverage (mean)	$39 \times$	$25 \times$	$33\times$	36×	$28 \times$	$42\times$	33x
No. of reads	5.68×10^{7}	3.65×10^{7}	4.89×10^{7}	5.31×10^{7}	4.20×10^{7}	6.28×10^{7}	5.00×10^{7}
No. of unique positions	1.11×10^{8}	1.18×10^{8}	1.10×10^{8}	1.12×10^{8}	1.18×10^{8}	1.14×10^{8}	1.15×10^{8}
No. of markers	N.A.	N.A.	N.A.	N.A.	178,042	43,204	21,164

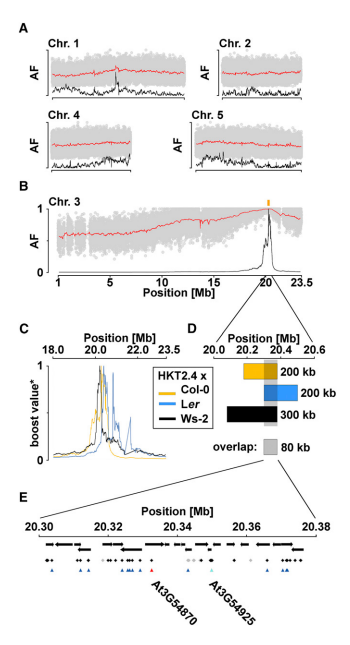


Figure 2. Identification of candidate SNPs through a combined mappingby-sequencing and HKT2.4-specific SNP-selection approach. A, Visualization of AFs at the noncausal chromosomes from the phenotypically selected F2 mapping population of HKT2.4 crossed to Col-0 (scheme in Supplemental Fig. S2). The points in gray are individual AFs at the markers, which were created by using SHOREmap according to the resequencing information of parental lines, namely HKT2.4 and Col-0. The red line shows the average AFs at the markers within 200-kb sliding windows (10-kb step). The black line is the sliding windowbased boost value, a slightly modified version of the r value as introduced with SHOREmap. B, The same visualization reveals that chromosome 3 carries the causal mutation for the selected phenotype. The peak of boost values is expected to be linked to the causal mutations. The yellow rectangle marks the SHOREmap-derived mapping interval. C, Enlargement for the boost values derived from HKT2.4 crossed to Col-0, Ler, and Ws-2 on chromosome 3. D, Intersection of SHOREmap-derived mapping intervals from the three crossings leads to a reduced region of 80 kb (20,300,000-20,379,999), which harbors the causal mutation. E, The 80-kb overlapping mapping interval with all genes annotated by

that, depending on the focal accession, different crossing partners (namely those with the highest SNP density and different recombination landscapes) can further increase the mapping resolution.

Identification of Candidate Genes through HKT2.4-Specific SNP Selection

The identified 80-kb mapping interval contained 21 genes (The Arabidopsis Information Resource release 10 [TAIR10], http://www.arabidopsis.org; Fig. 2E). To further narrow down the number of candidate genes, we made use of the sequencing data and extracted all SNPs that were HKT2.4 specific and therefore did not occur in Col-0, Ws-2, or Ler (Supplemental Table S3). This step resulted in 166 candidate SNPs. Using the Col-0 TAIR10 annotation data, we identified 72 intragenic SNPs, 13 of which change the amino acid sequence of the corresponding proteins, reducing the number of candidate genes to 5 (Supplemental Table S3).

Because none of the 62 accessions showed the HKT2.4-specific phenotype, it is conceivable that they do not have the causative polymorphism. We therefore extracted the sequences of the mapping interval for the analyzed 62 accessions from the 1001 Genomes Project (http://signal.salk.edu/atg1001/3.0/gebrowser.php), called the SNPs, and selected for HKT2.4-specific SNPs (Supplemental Table S3). When selecting only those HKT2.4-specific SNPs that were identified by both approaches (sequencing and an accession-based approach), 36 SNPs were considered, 15 of which are intragenic and two of the intragenic SNPs change the amino acid sequence of two proteins (Fig. 2E; Supplemental Table S3).

For the latter two SNPs, we extracted the nucleotide at the corresponding position for all available accessions from the 1001 Genomes Project. For the first SNP at position 20,332,575 on chromosome 3, none of the accessions shared the SNP with HKT2.4 (Supplemental Table S4). For the second SNP, only one accession (Kastel-1) shared the same SNP allele with HKT2.4 at position 20,349,799 on chromosome 3. Moreover, 11 additional accessions featured a third allele at the same position (Supplemental Table S5). Neither Kastel-1 nor one of the other accessions (Taynuilt) showed root hair branching (Supplemental Fig. S6). The SNP at position 20,332,575 on chromosome 3

arrows according to TAIR10 (ftp://ftp.arabidopsis.org/home/tair/Genes/TAIR10_genome_release/TAIR10_gff3/). Black and gray diamonds are HKT2.4-specific intragenic SNPs identified from the sequencing approach. Gray indicates SNPs with a coverage below 10. Triangles indicate SNPs that are also HKT2.4 specific for the set of 62 initially analyzed accessions. Only two of these change the coding DNA sequence as annotated by TAIR10. Cyan indicates nonsynonymous SNPs. Red indicates change of splicing site. On the basis of this selection, the two most likely causal genes are given. See Supplemental Figures S4 and S5 for the crossings with Ler and Ws-2. The HKT2.4-specific SNP identification from the sequencing and from the accession-based approach is documented in Supplemental Table S3. Chr, Chromosome; Mb, megabase pair.

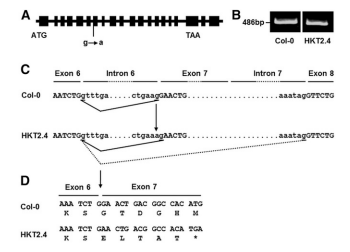


Figure 3. *ARK1* gene structure and splicing site in Col-0 and HKT2.4. A, A scheme of the Col-0 *ARK1* gene structure (annotated according to TAIR10, http://www.arabidopsis.org). Boxes represent exons. HKT2.4 has a G-to-A nucleotide exchange in the splicing acceptor site of intron 6. B, Real-time PCR for the region surrounding the mutated splicing site on *ARK1* showing that Col-0 and HKT2.4 both have a transcript of similar size. C, A scheme showing the actual splicing sites based on sequencing (black lines) and prediction based on NetGene2 (dashed lines). Note that the splicing site in HKT2.4 is shifted only by one nucleotide compared with Col-0. D, Sequencing revealed that the differing splicing sites in *ARK1* of HKT2.4 causes a frame shift resulting in a premature stop codon in exon 7 (asterisk).

changes a splicing site in At3g54870 according to the TAIR10 annotation. The gene encodes *ARK1/MRH2*. Root hair branching has been reported for mutants of *ARK1* (Jones et al., 2006).

Without additional information, we could identify the causative SNP exclusively by the analysis of wild accessions and their genomes.

Interestingly, no other accession carries the causative SNP in the worldwide collection of more than 800 sequenced accessions, although several accessions around Tuebingen are among these. There might have been either selection against this SNP or the SNP has recently arisen.

A Modification in an ARK1 Splicing Site Causes the HKT2.4-Specific Root Hair Branching Phenotype

Compared with the Col-0 TAIR10 annotation, the splicing acceptor site in intron 6 of ARK1 differs in HKT2.4 (Fig. 3A). We generated complementary DNA from the relevant root tips including the lower zone analyzed in this study. Using primers located in exon 6 and exon 7, we obtained a PCR product for Col-0 and HKT2.4 of similar size (Fig. 3B). A control PCR using primers for the translation elongation factor EF1 α A4 (Kirik et al., 2007) verified the absence of genomic DNA. According to splicing site prediction by NetGene2 (Brunak et al., 1991; Hebsgaard et al., 1996), no PCR product was expected for HKT2.4 because the splicing acceptor site of intron 7 was predicted

to be used instead of the splicing acceptor site in intron 6 (Fig. 3C). This prompted us to sequence the HKT2.4 complementary DNA. As a result, the splicing site in intron 6 of HKT2.4 was found to be shifted by one nucleotide leading to a frame shift and premature stop codon in exon 7 (Fig. 3D).

The *ark1-1* line is a transfer DNA mutant in the Col-0 background for which a branching phenotype has been reported (Jones et al., 2006). We determined the percentage of branched root hairs in *ark1-1* in the lower root region and found no significant difference from HKT2.4 (Supplemental Table S2). Then the causative nature of the *ARK1* allele from HKT2.4 was verified by crossing *ark1-1* with HKT2.4. F1 seedlings showed

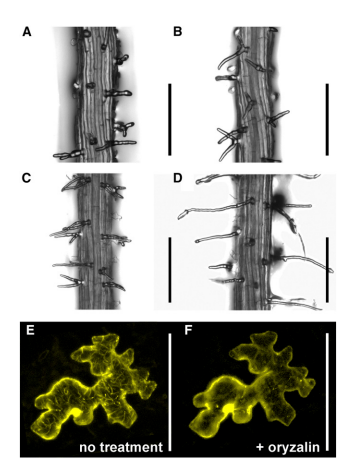


Figure 4. *ARK1*^{Col-0} complements root hair branching in HKT2.4 and localizes to MTs. A and B, The ark1-1 mutant (A) and HKT2.4 (B) have branched root hairs in contrast with Col-0 (Fig. 1 shows Col-0). C, F1 plants of HKT2.4 crossed to ark1-1 have branched root hairs. D, Overexpression of the genomic region of *ARK1* from Col-0 rescues the HKT2.4 branched root hair phenotype in T1 plants. The plant with the strongest rescue is shown . See Supplemental Table S6 for the analysis of additional seedlings. For all bright-field pictures, z-stacks of the upper zone (Supplemental Fig. S1) of 7-d-old seedlings were used. E and F, Localization of YFP-ARK1^{Col-0} before (E) and after (F) oryzalin treatment. Transient expression using the YFP-ARK1^{Col-0} rescue construct in Arabidopsis leaf epidermal cells. CLSM z-stacks (step size, 2 μm) are shown. Scale bars = 250 μm in A to D and 100 μm in E and F. [See online article for color version of this figure.]

branched root hairs similar to HKT2.4 and *ark1-1* confirming the allelism (Fig. 4; Supplemental Table S2).

To test whether ARK1 from Col-0 (ARK1^{Col-0}) can rescue the HKT2.4 root hair branching phenotype, yellow fluorescent protein (YFP)-ARK1^{Col-0} and ARK1^{Col-0} were overexpressed under the Cauliflower mosaic virus 35S promoter (Pro35S) in HKT2.4 background. The transformation efficiency of HKT2.4 was low. We obtained four Pro35S:YFP-ARK1^{Col-0} (HKT2.4) lines and eight Pro35S:ARK1^{Col-0} (HKT2.4) lines, respectively, which showed partial rescue (Fig. 4; Supplemental Table S6). The *Pro35S:YFP-ARK1*^{Col-O} (HKT2.4) rescue lines did not show a detectable YFP signal; therefore, we tested the construct in a transient expression assay in Arabidopsis (Col-0) rosette leaf epidermal cells. As previously shown in infiltrated tobacco (Nicotiana tabacum) leaves (Yoo et al., 2008) and supported by oryzalin treatment, YFP-ARK1^{Col-0} can decorate MTs in Arabidopsis (Fig. 4, E and F). Whereas oryzalin abolished YFP-ARK1 Col-0 localization to cytoskeletal structures, this was not seen upon latrunculin B treatment (Supplemental Fig. S7). ARK1 function has been well characterized in root hairs of the Arabidopsis Col-0 ecotype (Sakai et al., 2008; Yoo et al., 2008). The ARK1 protein was recently shown to be a MT plus-end directed catastrophe factor (Eng and Wasteneys, 2014). ARK1 is additionally proposed to specify root hair polarity through interactions with actin and proper targeting of ROP2 and RAB GTPase homolog A4B to the root hair tip (Yang et al., 2007; Yoo and Blancaflor, 2013).

Together, these data show that the candidate SNP in *ARK1*, as identified by the combined mapping-by-sequencing and HKT2.4-specific SNP-selection approach, leads to a splicing site shift causing the root hair branching phenotype of HKT2.4.

The presented pipeline can also be applied for causative SNPs that change regulatory regions. In this case, the SNP-selection process starting with Supplemental Table S3 would be conducted as described for SNPs in the coding DNA sequence but focus on SNPs in the regulatory regions. Subsequent verification can be done through expression analysis.

This study describes a pipeline for the identification of causal SNPs for phenotypes occurring only in one or a few accessions. We present strategies that narrowed down the final mapping interval for the unique root hair branching phenotype of HKT2.4. Application of an unbiased accession-specific SNP-selection approach led to the identification of the causative SNP in *ARK1*.

MATERIALS AND METHODS

Plant Material and Growth Conditions

The following Arabidopsis (*Arabidopsis thaliana*) lines were used in this study: Col-0, Ws-2, Ler, HKT2.4 (CS76404), Kastel-1 (CS22807), Taynuilt (CS1572), ark1-1 (Salk_035063C; Alonso et al., 2003; Jones et al., 2006), $Pro35S:ARK1^{Col-0}$ (HKT2.4), and $Pro35S:YFP-ARK1^{Col-0}$ (HKT2.4). Plant transformation (Clough and Bent, 1998) and growth conditions were previously described (Rishmawi et al., 2014).

Plasmid Construction

For $ARKI^{Col-0}$, the genomic region (from the start codon to stop codon with the introns) was amplified from Col-0 (primers in Supplemental Table S7),

cloned into pDONR20,1 and subsequently transferred to pFAST-R02 (Shimada et al., 2010) and pENSG-YFP using Gateway technology (Invitrogen). pENSG-YFP originates from N. Medina-Escobar (unpublished data). A version for C-terminal fusions is published (Feys et al., 2005).

Material for Sequencing

Reciprocal crosses between Arabidopsis ecotypes were done to obtain the F2 mapping populations. About 200 F2 seedlings of the reciprocal crosses showing HKT2.4-like branched root hairs were pooled. DNA was extracted using an DNeasy Plant Mini Kit (Qiagen). Samples of at least 2 μ g of total genomic DNA with high purity (260:280 ratio \geq 1.8) were sequenced by the Cologne Center for Genomics using one lane for paired-end Illumina HiSeq sequencing.

Resequencing and SHOREmap Analysis

Resequencing was performed with the SHORE short-read analysis pipeline (Ossowski et al., 2008). Raw reads of all seven samples were aligned against the Col-0 reference genome (ftp://ftp.arabidopsis.org/home/tair/Genes/ TAIR10_genome_release/TAIR10_chromosome_files/) using GenomeMapper (Schneeberger et al., 2009a). Short-read alignments were corrected for readpair information. Consensus calling was performed by using SHORE consensus. SHOREmap v2.1 (http://shoremap.org) was used to predict mapping intervals for each of the three bulked mapping populations (Schneeberger et al., 2009b). SNP variants identified in the resequencing analyses of the parental strains were filtered by SHOREmap create and those passing the filtering were used as markers. SHOREmap outcross calculates the AF estimates at each marker position (i.e. the ratio of the reads of mutant alleles divided by all reads at that marker locus) and determines mapping intervals according to the average AFs within sliding windows and the corresponding coefficients of variation. In addition to AFs, SHOREmap visualizes a different metric for the AFs (boost value $[B_v]$). If $m_{\rm obs}$ is the observed mean of AFs within a sliding window and t is the AF at which the causal mutation segregates, then $B_{\rm v} = 1/|1 - \max(t, 1 - t)/\max(m_{\rm obs}/1 - m_{\rm obs})|$. Polymorphisms in the 80-kb intersection of the three mapping intervals (i.e. chromosome 3:20,300,000-20,379,999) were functionally annotated based on TAIR10 gene annotations (ftp:// ftp.arabidopsis.org/home/tair/Genes/TAIR10_genome_release/TAIR10_gff3/).

HKT2.4-Specific SNP Selection

Sequences of the accession (Supplemental Table S1) were extracted from the Arabidopsis 1001 Genomes GEBrowser (http://signal.salk.edu/atg1001/3.0/gebrowser.php) for the region of the mapping interval on chromosome 3-covering positions 20,300,000 to 20,379,999 (Cao et al., 2011) and were aligned with R software (http://www.r-project.org/) using the adegenet and seqinr package (Charif and Lobry, 2007; Jombart, 2008; R Core Team, 2013). Single-candidate SNPs were similarly extracted from the GEBbrowser for all available accessions. Sequences were available from the U.S. Department of Energy Joint Genome Institute, the Gregor Mendel Institute of Molecular Plant Biology, Institute for Genome Research and Systems Biology of the University of Bielefeld Center for Biotechnology, Max Planck Institute for Developmental Biology, Monsanto Company, the Salk Institute Genomic Analysis Laboratory, or published sources, respectively (Cao et al., 2011; Gan et al., 2011; Long et al., 2013; Schmitz et al., 2013).

Splicing Site Determination

RNA was extracted from the root using an RNeasy Plant Mini Kit. The fragment flanking the splicing site was amplified using Pf-splicing MRH2 and Pr-splicing MRH2 (Supplemental Table S7). NetGene2 was used for prediction of splicing sites (Brunak et al., 1991; Hebsgaard et al., 1996).

Microscopy, Root Hair Branching Analysis, and Transient Assay

Root hairs were analyzed in the elongation zone (lower zone) by bright-field microscopy with a Leica DM5000B microscope equipped with a Leica DFC360FX camera and Leica Application Software AF (Leica Microsystems). Pictures were acquired as previously described (Failmezger et al., 2013). For transient expression, 3-week-old Arabidopsis (Col-0) rosette leaf epidermal cells were biolistically

transformed with $ARK1^{Col·0}$ -pENSG-YFP or the F-actin marker Lifeact-eGFP (Era et al., 2009) as previously described (Steffens et al., 2014) and analyzed with confocal laser-scanning microscopy (CLSM) after 19 to 21 h. CLSM was performed using a Leica TCS-SPE confocal microscope (DM5500Q) equipped with Leica Lite 2.05 software (LCS; Leica Microsystems). Z-stacks (step size: 2 μ m) were projected with the software's max projection. Drug treatments with oryzalin (40 μ M, 2 h) and latrunculin B (100 μ M, 1.5–2 h) were previously described (Steffens et al., 2014).

Statistical Analysis and Software Used

Statistical analysis, except for the resequencing and mapping, was conducted using Origin (OriginLab) applying the Anderson-Darling test (normal distribution) and the Mann-Whitney *U* test (significance). Images were processed using ImageJ software (http://imagej.nih.gov/ij/, 1997–2012; U.S. National Institutes of Health) and Photoshop Elements software (version 7.0.1; Adobe Systems using brightness and contrast adjustments for z-stacked pictures only. Z-stacking was done using CombineZM (Alan Hadley, http://web.archive.org/web/20090122072134/http://hadleyweb.pwp.blueyonder.co.uk/CZM/News.htm).

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number ARK1 (AT3G54870; Gene ID 824652).

Supplemental Data

- The following materials are available in the online version of this article.
- **Supplemental Figure S1.** Selection of seedlings for F2 pools used for mapping-by-sequencing.
- Supplemental Figure S2. Scheme of mapping-by-sequencing using accessions to generate a mapping interval.
- **Supplemental Figure S3.** Alignment statistics of short reads of parental lines and the mapping populations.
- **Supplemental Figure S4.** AF plots from the phenotypically selected F2 mapping population of HKT2.4 × Ler.
- Supplemental Figure S5. AF plots from the phenotypically selected F2 mapping population of HKT2.4 imes Ws-2.
- Supplemental Figure S6. Root hair phenotypes of accessions with different candidate SNP alleles in AT3G54925.
- Supplemental Figure S7. Latrunculin B treatment cannot abolish YFP-ARK1^{Col-0} cytoskeletal localization.
- Supplemental Table S1. List of accessions analyzed for root hair branching phenotype in the lower zone.
- Supplemental Table S2. Root hair branching percentages.
- Supplemental Table S3. HKT2.4-specific SNPs and their sequence quality and annotation.
- Supplemental Table S4. Candidate SNP in AT3G54870 extracted for 855 Arabidopsis accessions.
- Supplemental Table S5. Candidate SNP in AT3G54925 extracted for 855 Arabidopsis accessions.
- Supplemental Table S6. Root hair branching in rescued T1 seedlings.
- Supplemental Table S7. List of primers.

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