Twisting development, the birth of a potential new gene

Highlights
The identification of a possible de novo-originated gene TWISTED1 in Brassicaceae

High expression of TWISTED1 causes right-handed helical growth of organs

Not the plant hormone auxin, but altered microtubule orientation causes helical growth

TWISTED1 CRISPR-Cas9 mutations affect plant development
Twisting development, the birth of a potential new gene

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SUMMARY
Evolution has long been considered to be a conservative process in which new genes arise from pre-existing genes through gene duplication, domain shuffling, horizontal transfer, overprinting, retrotransposition, etc. However, this view is changing as new genes originating from non-genic sequences are discovered in different organisms. Still, rather limited functional information is available. Here, we have identified TWISTED1 (TWT1), a possible de novo-originated protein-coding gene that modifies microtubule arrangement and causes helicoidal growth in Arabidopsis thaliana when its expression is increased. Interestingly, even though TWT1 is a likely recent gene, the lack of TWT1 function affects A. thaliana development. TWT1 seems to have originated from a non-genic sequence. If so, it would be one of the few examples to date of how during evolution de novo genes are integrated into developmental cellular and organismal processes.

INTRODUCTION
Genes have long been thought to arise almost exclusively from pre-existing genes through gene duplication, domain shuffling, horizontal transfer, overprinting, retrotransposition, etc.1–3 The genesis of new genes from “scratch,” non-coding, or non-genic sequences was considered extremely unlikely.1,4 However, it is becoming increasingly clear that the de novo birth of genes is more prevalent than was previously thought.5 “De novo” genes are present in a restricted number of related species, and have been found in different organisms including yeast, animals, and plants.6–13 For some such genes, functions have been identified, e.g., modulation of starch metabolism in plants, and adaptation to freezing temperatures in fish.14,15 However, for most de novo genes, their biological functions and the effects of their integration into the cellular machinery or networks remain unknown. Moreover, while the effects of new genes in the development of some animal species are starting to be clarified,16 the effects of these genes in the development of plants have not been explored.

Here, we describe the characterization of the seemingly taxonomically restricted TWISTED1 (TWT1) gene, which likely arose de novo from a non-genic sequence and is involved in plant development. TWT1 conspicuously modifies the growth and development of the plant Arabidopsis thaliana.

RESULTS AND DISCUSSION
Identification and characterization of the TWISTED1 gene in Arabidopsis
TWISTED1 (TWT1; At2g32275) was identified in a mutant (twt1-D) with peculiar, twisted siliques in Arabidopsis thaliana (Figures 1A, 1B, 1G, and 1H). The mutant was discovered in a forward genetic screening of a transposon-based activation-tag population.17 General plant development and reproduction were not severely affected in the mutant, but twt1-D presented a well-organized, prominent, right-handed twisted or helical growth pattern in the petioles, stem, petals, gynoecium, and inflorescences (Figures 1 and S1). Moreover, the epidermal cell files of twt1-D hypocotsyls and roots were also arranged as right-handed helices (Figures 1K and 1M). At early stages of development, twt1-D exhibited shortened primary roots, with the early development of lateral roots (Figures 1I and 1J). The size of the root meristematic region and columella were reduced, though quiescent center (QC) markers such as WOX5 and QC46 were
Figure 1. twt1-D confers a twisted growth pattern in different organs

(A–J) See also Figures S1 and S2. Wild-type (WT) and twt1-D siliques (A, B), stamens and pistils (C, D), and rosettes (E, F). In (F) yellow arrows indicate right-handed twisting. Electron microscopy image of WT and twt1-D siliques (G, H). Seedlings presenting differences in root architecture (I, J).

(K and L) Confocal microscopy image of the upper hypocotyl region of WT (K), and twt1-D (L).

(M) Confocal microscopy image of a twt1-D root.

(N) Subcellular localization of TWT:GFP protein in the plasma membrane region of root cells (in a plant bearing the gtwt1-D construct).

(O) Schematic representation of the twt1-1 genomic region, where the activating I element (AIE) was inserted. Blue lines represent the different regions used for constructs that recapitulated the phenotypes. The four arrows in the AIE represent the enhancer tetramer in the transposon. Dark green boxes represent exons, light green boxes represent UTRs, and triangles indicate the orientation of the gene. The black line represents non-coding genomic regions and introns.

(P–S) Phenotypes of stable transgenic lines bearing the constructs depicted in the scheme in (O). Each construct recovers the twt1-D phenotype (right-handed twists indicated by yellow arrows).
The activation tagging population from where the mutant was isolated contains an engineered maize non-autonomous transposable element, Inhibitor (I, or defective Suppressor-Mutator, dSpm), that harbors a tetramer of the CaMV 35S enhancer. This element was named Activating I Element (AIE). The twt1-D phenotype was dominant, resulting from a single activation transposon insertion. The Activating I Element (AIE) was found to be inserted in chromosome 2 between a microRNA (MIR417; At2g32273) and a small peptide-coding region (At2g32275). 334 bp downstream of the transposon enhancer sequences (Figure 1O). miRNA417 has been reported to be involved in the salt stress response, but no twisted phenotype was described. When we tested two recapitulation constructs, including either 6.5- or 2.8-kb-long genomic fragments amplified from the original mutant, comprising the enhancer tetramer and its closest adjacent genomic sequence (Figure 1O), they produced phenotypes similar to those of twt1-D when introduced into wild-type plants (Figures 1P and 1Q). Both contained At2g32275 but not MIR417. Furthermore, At2g32275 expression was also found to be strongly increased in twt1-D compared to wild-type plants (Figure S2). After the increase in expression was confirmed, fragments of either the coding region together with the 3’-UTR of At2g32275 or the coding region alone were overexpressed using the CaMV 35S promoter in wild-type plants (Figure 1O). The increased expression of At2g32275 in these plants also recapitulated the twt1-D twisted leaves and siliques phenotype (Figures 1R, 1S, and S2). Furthermore, when the At2g32275-coding region placed in antisense orientation under the control of the CaMV 35S promoter (35S::antiTWT1) was introduced into twt1-D plants, the expression of At2g32275 was markedly reduced in comparison to the twt1-D mutant (Figure S2) and the characteristic twisting of rosette leaves and inflorescences was suppressed in these plants (Figures 1T–1X). Therefore, we concluded that the twt1-D phenotype was the result of the overexpression of the At2g32275 gene, which was renamed TWISTED1 (TWT1).

The annotation for the At2g32275 gene was “Expressed protein.” It has a relatively short CDS of 279 nucleotides and a single intron in its 3’-UTR (TAIR; arabidopsis.org). Surprisingly, homology searches in the NCBI database using BLASTx and tBLASTn revealed no similar genes or proteins in plants or other organisms besides close relatives of the Brassicaceae family (Arabidopsis lyrata, Capsella rubella, and lower similarity to Arabis alpina). This lack of gene homology with other organisms suggested that TWT1 could be a “new” gene, only present in A. thaliana and closely related species.

To evaluate whether TWT1 is indeed a gene and that its expression was not an artifact created by the nearby enhancer in twt1-D, we searched for transcripts in expression databases. TWT1 transcripts were found at low levels in the expression database ARS and by RT-PCR analysis using different Arabidopsis organs (Figure S2). A fusion of the TWT1 promoter with the β-glucuronidase (GUS) reporter gene was used to explore the pattern of TWT1 expression in more detail. In young plants, GUS activity was mainly observed in the vasculature of cotyledons and leaves, the root meristematic region, and inner tissues of the mature root (Figure S2). Thus, the TWT1 promoter and gene are active in specific tissues. We further explored whether the putative protein coded by TWT1 was stable and whether it had a specific localization in the cell. To visualize this protein, we used a translational fusion to GFP. To obtain enough signal, a genomic fragment cloned from the twt1-D mutant comprising the enhancer tetramer in the transposon and the adjacent region containing the TWT1 gene ending just before its stop codon was fused to GFP. This construct was named gtwt1-D-GFP. The roots of stable Arabidopsis transformants were analyzed, and fluorescence was observed at the edges of the cells, at the membrane (Figures 1N and S2). The construct was also used to perform transient expression assays by agroinfiltration in tobacco leaves, where a similar localization was observed (Figure S2). These findings corroborate previous proteomics studies that have identified TWT1 among plasma membrane proteins of Arabidopsis cells.
**Phenotypes due to increased TWISTED1 expression are related to microtubules**

We then sought to ascertain the physiological processes affected by the increased expression of TWT1 in the mutant plant that caused the peculiar twisted or helicoidal morphological phenotype and root alterations. Auxin is a key plant hormone involved in many plant developmental processes. Alterations in the auxin pathway can cause organ torsions, twisted growth in non-fixed orientations, and root development defects.\(^\text{21,22}\) We analyzed reporter lines for the transcriptional auxin response and auxin transport in the roots of the twt1-D mutant (Figure S3). Furthermore, we evaluated the effects of IAA and NPA treatments (Figure S3). The results suggested that reduced growth of the root in twt1-D is not due to reduced auxin content or obvious differences in auxin transport. Moreover, they could also not explain the observed twisted phenotype.

Helical growth phenotypes can also be caused by alterations in microtubules or microtubule-associated proteins and drugs that affect microtubule polymerization or depolymerization. When the properties of cortical microtubules are compromised, elongated organs can exhibit helical or twisted growth in a fixed orientation.\(^\text{23,24}\) Therefore, we examined the effects of propyzamide, a microtubule-disrupting drug, on twt1-D growth. Propyzamide is a benzamide that binds to β-tubulin, alters microtubule dynamics, and at low doses causes left-handed helical growth.\(^\text{25,26}\) Wild-type seedlings grown for a week on medium supplemented with 1, 3, or 5 μM propyzamide displayed definite left-handed helical growth (Figures 2A, 2E, and S3), while untreated twt1-D plants displayed a right-handed twisted phenotype (Figures 2B, 2H, and S3). At 0.5 μM propyzamide, the rightward twisting of twt1-D cotyledons decreased (Figures 2C and 2D) and was even reversed to leftward twisting at 1 and 3 μM (Figure S3). Moreover, twt1-D mutants grown for 3 or 4 weeks at 5 μM propyzamide were more resistant to this long-term treatment than wild-type plants (Figures 2G–2K). The growth of wild-type plants was severely affected by this treatment. In contrast, twt1-D plants appeared to be less altered, as evidenced by the larger size of the leaves and rosette, and the greener color (Figures 2G–2K). Therefore, it seems that the increased expression of TWT1 in the mutant counteracts the growth inhibition caused by long exposure to 5 μM propyzamide.

Next, we tested the genetic interaction with tubulin mutants. twt1-D was crossed with tua5\(^\text{D251N}^{-}\) and tua4\(^{S178D}^{-}\), which present right-handed and left-handed growth, respectively.\(^\text{24}\) (Figures 2M–2O). The tua5\(^{D251N}^{-}\) and tua4\(^{S178D}^{-}\) double mutant displayed a prominent, enhanced, right-handed helical growth phenotype in the leaves and cotyledons (Figures 2M, 2N, 2P, and 2Q). Conversely, the tua4\(^{S178D}^{-}\) and twt1-D double mutant demonstrated a marked reduction or suppression of the left- and right-handed twisting displayed by the respective single mutants (Figures 2M, 2O, 2P, and 2R). The orientation of cortical microtubules is crucial for normal plant cell elongation, and alterations frequently result in helical or twisted growth.\(^\text{24}\)

To analyze the orientation of cortical microtubules in twt1-D, we crossed it with the GFP::TUA6 marker line.\(^\text{27}\) In wild-type plants, cortical microtubules were aligned almost perpendicularly to the long axis of the cell (Figure 2S), whereas, in twt1-D, the microtubules did not show this orientation and appeared to produce left-handed oblique cortical arrays (Figures 2S and 2T). Therefore, the increased expression of TWT1 in the twt1-D mutant appears to affect microtubule orientation, and thereby the cytoskeleton. This was surprising, considering that TWT1 is a recent gene we did not expect it to influence basic cellular structures.

**TWISTED1 likely originated from a non-genic sequence**

Next, we investigated the evolutionary origin of TWT1 by synteny and phylogenetic analyses (Figures 3 and S4), which suggested a recent de novo origin of TWT1 in species related to A. thaliana. The synteny regions of the genomes of these species were analyzed and searched for sequences, using tblastn, that would produce proteins with homology to TWT1. In nine Brassicaceae genomes analyzed, open reading frames (putative in some cases) with homology to TWT1 were identified. Though the identity of these putative TWTs was low (Table S1), at least three regions in the protein sequence appeared to be conserved, to some extent, as revealed by conservation analyses based on the multiple protein sequence alignments (Figure S4). We identified one of these regions as a “conserved TWT1 motif” (WxxPxLxxLxxE). The synteny of the genomic regions where these putative orthologues are is well conserved in most of the analyzed related genomes (Figures 3A, 3C, and S4).

Tarenaya hassleriana is a plant that belongs to Cleomaceae, a sister family of Brassicaceae. The Cleomaceae and Brassicaceae lineages diverged only ~38 million years ago.\(^\text{28}\) Interestingly, though the T. hassleri-ana genome has a very conserved syntenic region, the region that would correspond to TWT1 is an intergenic non-coding region (indicated by a red diamond in Figure 3C). After a careful inspection of putative
Figure 2. Microtubule disruption, mutation, and orientation analyses in twt1-D
See also Figure S3.
(A–F) Effects of propyzamide on helical growth phenotypes in wild-type (WT) (A, C, E) and twt1-D mutants (B, D, F). Seedlings grown on control medium (A, WT; B, twt1-D) or medium containing 0.5 μM (C, WT; D, twt1-D) and 5 μM (E, WT; F, twt1-D) propyzamide for 7 days.
(G–J) WT (G, I) and twt1-D mutant (H, J) plants grown on control agar medium (G, H) or medium containing 5 μM propyzamide (I, J) for three weeks.
(K) WT (left) and twt1-D plants grown in medium containing 5 μM propyzamide for four weeks.
(L) Scheme indicating right- (yellow arrows) and left- (cyan arrows) handed twist.
(M–R) twt1-D interaction with tubulin mutants; (M) WT, (N) right-handed twisted TUA5S251N mutant, (O) left-handed twisted TUA4S178D mutant, (P) twt1-D, (Q) twt1-D TUA5S251N double mutant exhibiting increased right-handed twisting, and (R) twt1-D TUA4S178D double mutant displaying reduced twisting. Cortical microtubules in seedling hypocotyls of GFP:TUA6 (S) and GFP TUA6 twt1-D (T), where microtubules are oriented to the left with respect to the top of the long axis of the cell. Asterisk indicates the basal part of the cell. Cartoons on the right illustrate the microtubule orientation in the cell (S, T). Scale bars: 2 mm (A–F), 5 mm (G–R), 10 μm (S, T).
Figure 3. Synteny and phylogenetic analyses of TWT1

See also Figure S4 and Table S1.

(A) Visualization of genomic regions surrounding the TWT1 gene based on GenBank files and the software clinker and clustermap.js. Syntenic genes are indicated in different colors. Gene percentage identity is presented in gray color scale.

(B) Phylogenetic tree inferred by maximum likelihood (ML) of putative TWT1 genes in Brassicaceae species.

Figure 3A shows the synteny of the TWT1 gene across different species in the Arabidopsis genus, indicating its conservation across various species. Figure 3B presents a phylogenetic tree constructed using maximum likelihood methods, showing the evolutionary relationships among the putative TWT1 genes in Brassicaceae species.
proteins also contain predicted disordered domains, though of different lengths and with variable secondary structure. These results suggest that coding TWT genes could be originated de novo in the Brassicaceae family. Different possible mechanisms of de novo protein genesis have been suggested, one of which proposes that these proteins arise when reading frames initially encoding disordered structures become expressed; young genes have indeed been found to encode highly disordered proteins. This appears to be the case for TWT1 protein, from aa 19 to 76 (Figures 4 and S4). Despite the low sequence identity percentages of the AtTWT1 acid (aa) TWT1 protein contains a predicted disordered domain that covers a large portion of the complete protein, from aa 19 to 76 (Figures 4 and S4). Despite the low sequence identity percentages of the AtTWT1 protein, a close inspection of one of the frames match, reveals the presence out of six (WxPxLxxIxExxE) amino acids of the conserved TWT1 motif (Figures S4 and S5). The DNA alignment between the A. thaliana TWT1 and the 7. hassleri nata putative ‘TWT1’ ORF is shown in Figure S5. Interestingly, two regions of the sequence, when translated in different open reading frames can give rise to one possibly conserved Cysteine at the beginning in one frame, and to an almost conserved TWT1 motif in the middle, in another frame (Figure S5). This sequence has the potential to become a TWT1 coding gene if various mutations would occur, meaning that this sequence could be ancestral to the Brassicaceae TWT1 coding genes.

Therefore, TWT1 likely originated recently in the Brassicaceae family. Different possible mechanisms of de novo protein genesis have been suggested, one of which proposes that these proteins arise when reading frames initially encoding disordered structures become expressed; young genes have indeed been found to encode highly disordered proteins. This appears to be the case for TWT1. The 92-amino acid (aa) TWT1 protein contains a predicted disordered domain that covers a large portion of the complete protein, from aa 19 to 76 (Figures 4 and S4). Despite the low sequence identity percentages of the AtTWT1 with TWTs from the very close species Arabidopsis lyrata, Capsella rubella, and Arabis nemen-ossis, these proteins also contain predicted disordered domains, though of different lengths and with variable sequences (Figure S4). Contrary to proteins that give rise to specific structures when they fold, an Intrinsically Disordered Protein or Domain does not fold spontaneously into a stable three-dimensional structure. These proteins or domains are disordered, and their structures are dynamic, changing their conformation continuously. Disordered domains were recognized some years ago and are gaining increasing attention due to their roles in different processes, including microtubule organization in animals and plants. It is thus plausible that the disordered domain of TWT1 could interact with microtubules, most likely cortical microtubules, considering the membrane localization of the protein.

TWISTED1 loss-of-function in Arabidopsis affects plant development

Finally, to know whether the loss of TWT1 function would affect plant development, the A. thaliana genome was edited using the CRISPR-Cas9 system. Three RNA guides were used to delete a large part of the coding region of the gene. Different deletion alleles were obtained. In two edited lines, in which 118 bp and 143 bp regions were deleted (Figures 4A and 4B), only the first 14 aa of the protein were maintained, whereas the amino acids that are changed due to frameshifts and premature stop codons in the coding region, eliminating the disordered region of the protein. No obvious differences were observed in microtubule organization in the epidermal cells of homozygous edited plants (Figure S6). However, the gene does not seem to be expressed in these cells, so more detailed analyses of other tissues would be required to discard that microtubules are not affected. Nevertheless, homozygous edited plants displayed abnormalities in their cotyledons such as elongated shape, dented edges, or size differences in the cotyledons of a seedling when grown on soil (Figure 4C). Older plants also exhibited delayed development, resulting in a reduced size phenotype (Figures 4D–4G). Furthermore, during reproductive development, edited plants presented smaller flowers (Figures 4H–4L) and reduced fruit length and seed number (Figures 4M–4P).
Figure 4. Phenotype and structure of edited TWT1 alleles

See also Figure S5.

(A) Genomic structure of two twt1 CRISPR-Cas9 edited alleles.

(B) TWT1 protein structure and comparison to the proteins produced by the edited alleles. In both cases, a deletion results in a frameshift and premature stop codons that produce smaller proteins than TWT1. The 30 and 4 letters in red represent the amino acids that are different from the original TWT1 amino acids in each allele, due to frameshifts. Asterisks represent stop codons.

(C) Ten-day-old twt1-cr2 seedling with dissimilar cotyledons.

(D and E) Seventeen-day-old wild-type (WT) (D) and twt1-cr1 (E) plant.

(F and G) Twenty-six-day-old WT (F) and twt1-cr1 (G) plants.

(H–L) Flower phenotypes of WT and twt1-cr mutants.

(M and N) Comparison between WT and mutant siliques.

(O and P) Comparison between WT and mutant open siliques with exposed seeds. Scale bars: 1 mm (C, H, I, M-P), 5 mm (D, E), 1 cm (F, G), 0.5 mm (J-L).
plants have been reported to be involved in other kinds of processes such as starch accumulation\textsuperscript{14} or pathogen-induced defense\textsuperscript{39}, but there are seldom examples of recent genes involved in plant development.

In summary, TWT1 is likely a de novo-originated lineage-specific gene able to produce a strong, yet exquisite, morphological phenotype in Arabidopsis. As occurs with most de novo genes\textsuperscript{8}, it is a small gene that is expressed in many tissues at the low level. It encodes a 92 aa protein consisting mostly of a disordered domain that can be found at the plasma membrane. It appears to have integrated into processes involving basic cellular structures such as microtubules. Determining whether TWT1 binds to cortical microtubules, possibly affecting their dynamics and cellulose deposition and therefore affecting plant morphology, will help further clarify the role of this possibly “new” Brassicaceae protein. Moreover, since it seems to have originated recently from a non-genic sequence, it would be very interesting to reconstruct the ancient gene or putative transitory proto-gene(s), if they existed in more genomes\textsuperscript{8}.

Limitations of the study
The identification of de novo-originated genes is not trivial. The erroneous conclusion of the identification of a lineage-specific gene that originated de novo could be the result of a homology detection failure\textsuperscript{40}. Furthermore, disordered domain protein-coding genes may evolve rapidly, and identification of homologs may be difficult. The putative ORF identified in the outgroup T. hassleri\textae may be ancestral to TWT1. Another scenario would be that the sequence in T. hassleri\textae was originally a TWT gene that was lost due to mutations. That would mean that the gene originated earlier. In the same line, we cannot exclude the possibility of horizontal gene transfer, though we did not find any evidence of similar proteins with the complete conserved TWT1 motif outside plant species. Only one hit to a bacterial sequence was found (MP285508.1), where a match between one amino acid (proline) matched the proline of the conserved TWT1 motif and other some less conserved residues. Furthermore, the bacterial sequence lacks the conserved Cysteine pair in the N-terminal part of TWT1, therefore we favor the hypothesis that the found hit to a bacterial sequence is not due to common ancestry but convergent evolution. Related to cortical microtubules, we cannot exclude that the natural function of TWT1 is not related to them. The affected microtubule arrangement, and other striking specific phenotypes such as helical growth of the fruit, were observed in an activation tag mutant and in overexpression lines, and therefore, care should be taken with drawing conclusions on the natural function of the gene.

DATA AND MATERIALS AVAILABILITY
All data and materials used in this study are available.

STAR METHODS
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SUPPLEMENTAL INFORMATION
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AUTHOR CONTRIBUTIONS
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DECLARATION OF INTERESTS
The authors declare no competing interests.

INCLUSION AND DIVERSITY
We support inclusive, diverse, and equitable conduct of research.

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## STAR+METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| anti-a-tubulin (mouse, monoclonal) | Sigma-Aldrich | Cat#T5168 |
| Alexa Fluor 488 (donkey anti-mouse IgG) | Invitrogen | Cat#A-21202, RRID: AB_141607 |
| Chemicals, peptides, and recombinant proteins |        |            |
| Indole-3-Acetic Acid (IAA) | Duchefa Biochemie | Cat#I0901, CAS 87-51-4 |
| Naphthylphthalamic Acid (NPA) | Chemical Service | Cat#N-12507, CAS 132-66-1 |
| Propyzamide | Sigma-Aldrich | Cat#45645, CAS 23950-58-5 |
| X-gluc | Gold Biotechnology Inc | Cat#G1281C, CAS 114162-64-0 |
| Propidium iodide | FLUKA/Sigma-Aldrich | Cat#81845, CAS 25535-16-4 |
| Cellulase Onozuka R-10 | Duchefa Biochemie | Cat#C8001, CAS 9012-54-8 |
| Driselase | Sigma-Aldrich | Cat#D8037, CAS 85186-71-6 |
| Mazerozyme R-10 | Yakult | N/A |

**Experimental models: Organisms/strains**

- *Arabidopsis thaliana*: twt1-D
- *Arabidopsis thaliana*: DR5rev::GFP
- *Arabidopsis thaliana*: PIN1::PIN1-GFP
- *Arabidopsis thaliana*: GFP-TUA6
- *Arabidopsis thaliana*: PIN3::PIN3-GFP
- *Arabidopsis thaliana*: DR5::GUS
- *Arabidopsis thaliana*: PIN2::PIN2-GFP
- *Arabidopsis thaliana*: PIN7::PIN7-GFP
- *Arabidopsis thaliana*: WOX5::GFP
- *Arabidopsis thaliana*: QC46::GUS
- *Arabidopsis thaliana*: tua4
- *Arabidopsis thaliana*: tua5
- *Arabidopsis thaliana*: pARC170 (twt1-D genomic long) This study
- *Arabidopsis thaliana*: pGD625STR (twt1-D genomic short) This study
- *Arabidopsis thaliana*: pGD625GES (35S::TWT1) This study
- *Arabidopsis thaliana*: pGD121GE (35S::cTWT1) This study
- *Arabidopsis thaliana*: pTWT1::GUS
- *Arabidopsis thaliana*: gtwt1-D-GFP
- *Arabidopsis thaliana*: 35S::antiTWT1 (pGD121ANT) This study

**Oligonucleotides**

- See Table S2

**Recombinant DNA**

- pCR2.1 Invitrogen Cat#K202020
- pGEM-T easy Promega Cat#A1360
- pDONR207 Invitrogen N/A
- pENTR/D-TOPO Invitrogen Cat#K240020
- pGD121 de Folter et al. 50 N/A
- pGD625 de Folter et al. 50 N/A

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Stefan de Folter (stefan.defolter@cinvestav.mx).

Materials availability

Plant lines generated in this study will be made available on request, but we may require a completed Materials Transfer Agreement if there is potential for commercial application.

Data and code availability

- All data reported in this paper will be shared by the lead contact upon request.
- This study did not generate new code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Plant materials and growth in soil

Seeds of the DR5rev::GFP (CS9361),41,42 PIN1::PIN1::GFP (CS9362),41 GFP.TUA6 (CS6551)27 lines were obtained from the Arabidopsis Biological Resource Center (https://abrc.osu.edu; ABRC, Ohio State University, Columbus). The PIN3::PIN3::GFP31 was obtained from Lars Østergaard. The DR5::GUS,44,45 PIN2::PIN2::GFP,16 PIN7::PIN7::GFP,47 WOX5::GFP,48 and QC46::GUS49 were donated by Luis Herrera-Estrella.
The tua5D251N and tua4s178d mutants were supplied by Takashi Hashimoto. Arabidopsis plants in soil were grown under normal greenhouse or growth chamber conditions (22°C, long day light regime).

METHOD DETAILS

Activation tagging mutant screen
The stable I transposon (based on the En-I or Spm-dSpm transposon system) population of Arabidopsis thaliana plants, ecotype Wassilewskija (Ws-3) was visually screened for pistil/silique aberrations under normal greenhouse conditions. From the selected twt1-D mutant, the F1 segregating generation was grown to examine the dominance of the mutation and copy number of the I element.

Southern blot analysis
Genomic DNA from 24 segregating plants and the parental mutant was isolated and approximately 300 ng of DNA was digested with the restriction EcoRI enzyme. Equal loading of the DNA was checked by Ethidium bromide staining. DNA was electrophoresed in a 1.0% (w/v) agarose gel in 1x TBE (1.0 M Tris, 0.9 M boric Acid, 0.01 M EDTA) blotted onto a Hybond N+ membrane (Amersham Pharmacy Biotech) following the normal manufacturer’s instructions. A 1.3 kb PCR fragment amplified from the 5' end of the BAR gene and from the 3' end of the right transposon junction, was labelled by random oligonucleotide priming (Gibco BRL) and used as a probe.

Localization of the activating I element in the Arabidopsis genome
To identify the putative activated gene, genomic DNA was used to isolate flanking fragments of the Activating I element (AIE) by modified thermal asymmetric interlaced–PCR (TAIL-PCR) as described previously. Flanking sequences were compared to the Arabidopsis database using BLASTN in the TAIR database. The insertion was located in a genomic region between two genes, At2g32273 and At2g32275 (TWT1). The promoter of gene At2g32275 was found to be the closest to the 4 x 35S enhancer cluster located at the left border of the I transposable element.

Gene expression analysis
The expression of gene At2g32275 (TWT1) was analyzed by semiquantitative RT-PCR. RNA was isolated from rosette leaves, stem, and flower buds using the using TRIzol method (Invitrogen). After DNase I treatment, cDNA was prepared using M-MLV (Invitrogen) according to the manufacturer’s instructions. Primers corresponding to the single coding exon (TWT1-exon), and the different splicing versions of the transcript (TWT1-splicing) were used.

Expression analysis of TWT1 was also performed using transcriptome data different tissues of Arabidopsis thaliana. Data was obtained from the ARS database; a collection of 20,000 RNA-seq libraries (http://ipf.sustech.edu.cn/pub/athrna/).

Recapitulation and reporter constructs
For the genomic recapitulation construct pARC170, a two-step cloning strategy was used to clone a 6.5 kb genomic fragment of the twt1-D mutant, starting from the NOS terminator within the AIE element to 446 bp downstream of the predicted stop codon. This 6.5 kb twt1-D genomic fragment was divided into two fragments to facilitate cloning. In all PCR amplification steps, TAQ plus precision polymerase enzyme (Stratagene) was used. Fragment “A” was amplified using the primers PRO142 and PRO143, resulting in a 4.5 kb fragment spanning the NOS terminator to 1.2 kb upstream of the putative ATG. The “B” fragment was amplified with the primers PRO097 and PRO176. This 3.2 kb fragment contains the region 2.0 kb upstream of the ATG to 446 bp upstream of the predicted stop codon. Each fragment was cloned independently into pGEM-T Easy (Promega), digested with the restriction enzymes SstI and BsmBI and ligated to each other resulting in the full 6.5 kb twt1-D genomic sequence in pGEM8-T Easy. Primers PRO142 and PRO176 contain additional GATEWAY™ sequences allowing recombination into the pDONR207 vector (Gibco BRL), creating pARC169; followed by LR recombination step into the vector pKGW, establishing pARC170 (twt1-D genomic long). The primer sequences with GATEWAY™ sites underlined are included in Table S2.

For the genomic recapitulation construct pGD625TR (twt1-D genomic short), a 2.8 kb genomic fragment was amplified from genomic DNA isolated from the twt1-D mutant. The fragment was cloned from the
NOS terminator within the AIE element to 500 bp downstream the predicted stop codon of At2g32275, using Pfx polymerase (Invitrogen). It was introduced into pENTR/D-TOPO (Invitrogen). For the overexpression recapitulation construct pGD625GES (35S::TWT1), a 795 bp genomic fragment was amplified from genomic DNA isolated from wild-type Ws-3 ecotype plants, the amplified region started from the ATG start codon to 500 bp downstream of the predicted stop codon. It was cloned in pENTR/D-TOPO. For the overexpression recapitulation construct pGD121GE (35S::cTWT1), the coding region of the At2g32275 gene was amplified with HiFi taq-polymerase (Invitrogen) using DNA from the same wild-type ecotype Ws-3 as the previous fragment and cloned into pCR2.1 (Invitrogen). All amplified fragments were verified by sequencing. The fragment used for pGD625STR was recombined with the binary Gateway vector pGD625-35S (the pGD625 vector without the 35S promoter). The fragments cloned for pGD625GES and pGD121GE were recombined or cloned into the binary overexpression vector pGD625 or pGD121 50, respectively.

To obtain a GUS reporter line, an 824 bp TWT1 (At2g32275) promoter fragment was amplified from genomic DNA isolated from wild-type Ws-3 plants using the HiFi taq-polymerase (Invitrogen). The fragment corresponds to the sequence directly upstream of the ATG start codon till the MIR417 gene and was cloned in pCR2.1 (Invitrogen). The amplified fragment was verified by sequencing. Finally, the fragment was cloned into the binary vector pBI12151, generating a transcriptional fusion with the GUS reporter gene (pTWT1::GUS).

A translational fusion with the Green Fluorescent Protein (GFP) was made by cloning a 2.3 kb genomic fragment amplified from twt1-D mutant genomic DNA using Pfx polymerase (Invitrogen) in pENTR/D-TOPO (Invitrogen). This fragment includes the enhancer of the AIE element and the coding region minus the stop codon. This construct was recombined with the binary Gateway vector pMDC204 (gtwt1-D::GFP).53 To perform transient expression analysis to visualize protein localization, the gtwt1-D::GFP construct was agroinfiltrated in young Nicotiana tabacum leaves as previously described in69 and stably transformed in Arabidopsis plants.

The antisense construct 35S::antiTWT1 (pGD121ANT) was made by introducing the reverse sequence of the At2g32275 gene (cTWT1) into the pGD121 binary vector,50 and was stably transformed in twt1-D plants.

Plant transformations and transformant selection
The constructs were used to transform Agrobacterium tumefaciens strain GV3101 and introduced into wild-type Arabidopsis thaliana, ecotype Col-0, Ws-3, or the twt1-D mutant using a modified floral dip method.70 For selection, the harvested seeds were vapor phase surface-sterilized (http://plantpath.wisc.edu/% abf/vapster.html) or sterilized by rinsing with 70% ethanol, leaving them for 10 to 15 min in a 20% bleach solution, and rinsing at least 3 times with sterile water. Depending on the construct, the selection of first generation (T1) seedlings was done by germinating seed on medium containing Arabidopsis 0.5X MS nutrient solution (PhytoTechnology labs), 1% sucrose, and 0.6% or 0.8% agar. The medium was supplemented with the corresponding selective agent. They were stratified for 2 days at 4°C, and grown at 22°C, under a long day light regime. Resistant plants were transferred to soil for further analysis. Alternatively, seeds were sown directly on soil, and resistant plants were selected after spraying twice with 100 mg L−1 Finale®, 7 days after germination.

In vitro phenotypic analyses
Seeds were handled and sown on plates containing medium as for transformant selection, with some differences. The medium contained 0.5 or 2% sugar, depending on the experiment, and 1.5% agar. Plates were placed either horizontally or in a near vertical position. Day 0 of growth was defined as the time when plates were transferred to the growth chamber, after stratification. Indole-3-Acetic Acid (IAA) (Duchefa Biochemie, Amsterdam, The Netherlands) and Naphthylphthalamic Acid (NPA) (Chem Service, Westchester, PA) stock solutions were prepared in ethanol. The final concentration used for the experiments was 5 μM. IAA treatment experiments were repeated twice, with a total n of at least 25 plants per treatment. The data is presented as a graph of the average length and the standard deviation.

The Propyzamide (PZ; Sigma-Aldrich) stock solution was prepared by dissolving it first in dimethyl sulfoxide (DMSO). Propyzamide was then used at a final concentration of 0.5–5 μM. The final concentration of DMSO
in the medium was lower than 0.3% and did not appear to affect the growth of seedlings. The experiment was repeated twice with an n of at least 10 plants per treatment.

**Leaf morphology and vasculature analyses**

For the analysis of leaf morphology and vasculature, the tissue was first fixed in 90% acetone at −20°C for one hour, followed by ethanol treatments at 80% and 40% (v/v) ethanol for a 24 h period each and fixed in 50% (v/v) glycerol. Cleared plants were imaged using an Olympus stereoscope SZH10.

**GUS reporter staining**

3- to 7-day-old seedlings were collected and placed in tubes containing GUS substrate solution composed of 50 mM sodium phosphate pH 7, 5 mM K3/K4 FeCN, 0.1% (w/v) Triton X-100, and 2 mM 5-bromo-4-chloro-3-indolyl-beta-GlcUA (Gold BioTechnology Inc). They were left in the solution for 2 to 6 days. After application of vacuum for 5 min, samples were incubated at 37°C for 24 h, processed to eliminate chlorophyll, and imaged.

**Tissue preparation and confocal analysis**

To observe the GFP fluorescence signal, hypocotyls and roots were collected and observed using a Zeiss LSM 510 META inverted confocal microscope (Carl Zeiss, Germany) with either a 20X or 40X air objective. 20 μM Propidium iodide (Fluka), was used as counterstain. GFP was excited with the 488 nm line of an Argon laser, and propidium iodide (PI) with a 514 nm laser line. GFP emission was filtered with a BP 500–520 nm filter and PI emission was filtered with a LP 575 nm filter.

**Scanning electron microscope analysis**

Fresh leaf and silique samples were visualized using a scanning electron microscope EVO40 (Carl Zeiss) using the VPSE G3 or the BSD detector with a 15–20 kV beam.

**Immunolocalization of microtubules**

Alpha-tubulin immunolabeling experiments were conducted following the protocol by with some modifications. Hypocotyls were dissected from six-days old Arabidopsis plants and transferred to 24-well plates to perform fixation, permeabilization, antibody incubation and washing steps. Cell permeabilization was done using 0.2% cellulase, 0.2% driselase, 0.2% mazerozyme, and 1% triton in 2 mM MES pH 5.0, during 1 h at room temperature. For primary antibody incubation an anti-a-tubulin primary antibody (T5168; mouse, monoclonal, Sigma) was diluted 1:800 and incubated for 16 h at 4°C in the dark; the secondary antibody was Alexa Fluor 488 (A-21202; donkey anti-mouse IgG, Invitrogen) diluted 1:1000 and incubated for 3 h at 37°C in the dark. Samples were mounted in 50% glycerol; images were acquired with a LSM800 confocal microscope using a 63X oil objective (Carl Zeiss).

**Phylogenetic and sequence analyses**

The protein coding sequence of TWT1 (AT2G32275.1) was used as a query in tblastn and blastp to search the nucleotide collection (nr/nt) and non-redundant (nr) databases at NCBI (https://blast.ncbi.nlm.nih.gov/Blast.cgi). To improve sensibility, the search was initially restricted to Brassicales (taxid:3699). Subsequent blast searches excluding the Brassicales were performed to look for TWT1 homologs outside this clade (none were found). Homolog sequences in Fasta and Genbank formats were downloaded from NCBI. Genbank files included neighboring genes for synteny analyses.

Multiple sequence alignments were performed with MUSCLE software as implemented in MEGA for macOS software. MEGA software was also used for phylogenetic inference based on Maximum Likelihood (ML) principle. Multiple sequence alignment was visualized with Jalview and with MAUVE (http://darlinglab.org/mauve/mauve.html) and genoPlotR.

**Generation of edited loss-of-function mutant alleles**

The CRISPR-Cas9 system was used to edit TWT1. The CRISPR construct was generated using GoldenGate cloning and the MoClo toolkit. The plasmids were obtained from Addgene (https://www.addgene.org/). Possible off-targets were analyzed at http://crispor.org. Three guide RNAs were designed inside the coding region of the gene, and ordered as the primers indicated in Table S2, fused to the U6-26 promoter (pICSL01009::AtU6-26p) and ligated into level 1 vector pICH47751 or pICH47761.
RPSSa promoter\textsuperscript{76} was cloned into pICH47742 and used to drive an Arabidopsis-optimized Cas9 (aCas9). The FAST seed marker pOle::TagRFP\textsuperscript{77} obtained in the MoClo kit as pICSL70008 \textsuperscript{55}, was cloned into pICH47732 and used as selection marker. These level 1 modules were subsequently combined into the binary level 2 vector pICSL4723. The resulting construct was transformed into Agrobacterium strain CS8C1 and introduced in Arabidopsis Ws-3 and Col-0 using floral dip. Transformed seed was selected by visual inspection for the RFP marker. First transformants were evaluated by PCR (twt1-cr primers in Table S2) and seed was obtained from those that presented clear deletions.

To evaluate segregation of the deletion and the morphological phenotype, around sixty seeds (sown as fifteen seeds per tray, in 4 trays) of 4 Ws-3 and 5 Col-0 twt1 CRISPR mutant alleles (twt1-cr) were sown in soil (peat moss, perlite and vermiculite 3:1:1). They were stratified at 4°C for 3 nights and moved to a growth chamber at 22°C. Two weeks after germination, the plants were transferred to a greenhouse with a temperature range from 22 to 28°C, and natural light conditions. Day length varied in different seasons. Genomic DNA from plants that were homozygous for the deletion was used as template to obtain a PCR fragment that was subsequently sequenced (using the same twt1-cr primers indicated in Table S2). The fragment was sequenced from both sides and provided the same, but complementary, sequence.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

For all the experiments, details of statistical tests used and error-bars in graphs are indicated in the relevant figure legends.