

A cascade of arabinosyltransferases controls shoot meristem size in tomato

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Shoot meristems of plants are composed of stem cells that are continuously replenished through a classical feedback circuit involving the homeobox *WUSCHEL* (*WUS*) gene and the *CLAVATA* (*CLV*) gene signaling pathway. In *CLV* signaling, the *CLV1* receptor complex is bound by *CLV3*, a secreted peptide modified with sugars. However, the pathway responsible for modifying *CLV3* and its relevance for *CLV* signaling are unknown. Here we show that tomato inflorescence branching mutants with extra flower and fruit organs due to enlarged meristems are defective in arabinosyltransferase genes. The most extreme mutant is disrupted in a hydroxyproline O-arabinosyltransferase and can be rescued with arabinosylated *CLV3*. Weaker mutants are defective in arabinosyltransferases that extend arabinose chains, indicating that *CLV3* must be fully arabinosylated to maintain meristem size. Finally, we show that a mutation in *CLV3* increased fruit size during domestication. Our findings uncover a new layer of complexity in the control of plant stem cell proliferation.

A wide range of inflorescence architectures exist among closely related plants¹, and this variation can be explained in part by the rate at which meristems terminate in flowers and initiate new branch meristems from which new flowers will form^{2,3}. However, less appreciated is the role of meristem size in shoot architecture, as well as in flower and fruit production. In maize, for example, variation in meristem size can account for differences in kernel row number between inbred lines⁴, and high row number is a major driver of yield in cultivated hybrids⁵. Meristem size is controlled by a maintenance mechanism that replenishes stem cells lost to lateral organ formation⁶. In *Arabidopsis thaliana*, meristem maintenance is based on a feedback system comprising the stem cell-promoting *WUS* homeodomain transcription factor and *CLV* signaling proteins. *CLV* signaling involves binding of the glycopeptide *CLV3* to cell surface leucine-rich repeat (LRR) receptor complexes that include the receptor kinase *CLV1* and related LRR receptors^{7–10}. These interactions initiate a signaling cascade that restricts *WUS* expression to prevent stem cell overproliferation, and, through negative feedback, *WUS* promotes *CLV3* expression to limit its own activity¹¹. Mutations in *CLV* pathway genes cause meristems to enlarge, which can lead to increased shoot and inflorescence branching, more flowers, and extra organs in flowers and fruits. Studies on mutants with these

phenotypes in maize and rice have shown that the *CLV* pathway is conserved in dicots and monocots¹². However, little is known about the genes and molecular mechanisms underlying stem cell proliferation in other crops, leaving unanswered whether the *CLV* pathway or other pathways are important in controlling meristem size and productivity. Here we investigate the control of meristem size in tomato.

RESULTS

Tomato *fab* and *fin* mutants have enlarged meristems

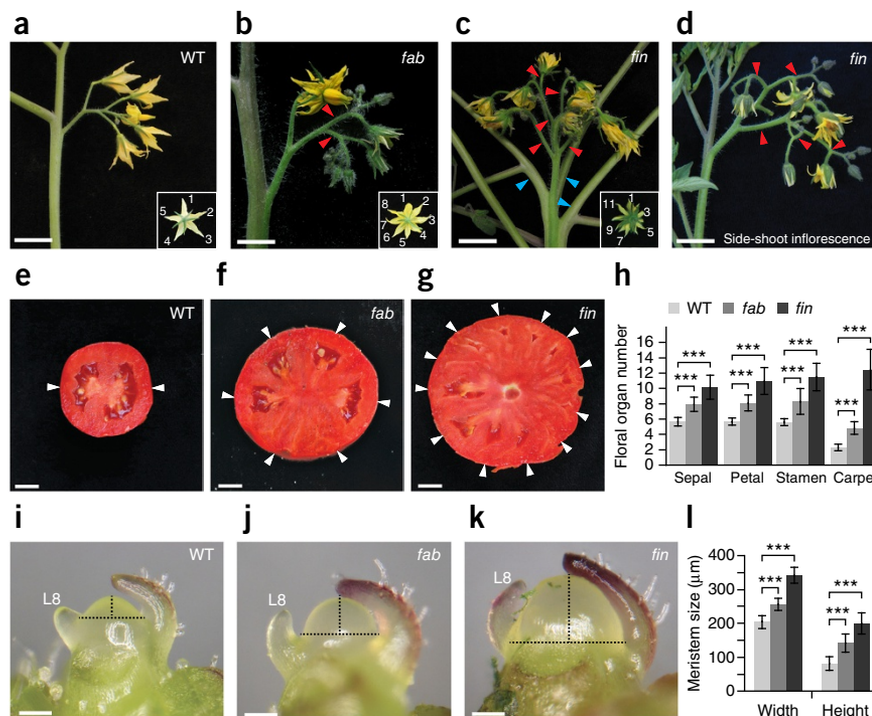
We explored the genetic control of inflorescence architecture in tomato by screening a large ethyl methanesulfonate (EMS) and fast neutron (FN) mutagenesis population for mutants with increased inflorescence branching¹³. In contrast to previously characterized mutants in which unbranched wild-type inflorescences are transformed into highly branched structures with normal flowers as a result of delay in meristem maturation^{3,14}, we identified six new mutants where branching was associated with fasciated flowers having more floral organs than wild-type flowers (Fig. 1a–h). Complementation tests identified two loci that we designated *fasciated* and *branched* (*fab*: one EMS allele) and *fasciated inflorescence* (*fin*: two EMS and three FN alleles).

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Figure 1 The *fab* and *fin* mutants develop branched inflorescences with fasciated flowers as a consequence of enlarged meristems.

(a) An unbranched inflorescence typical of a wild-type (WT) plant and a typical wild-type flower (inset). (b,c) Primary inflorescences from *fab* (b) and *fin* (c) mutants showing branching and fasciated flowers (insets). (d) A branched inflorescence from a *fin* side shoot. Red arrowheads, branches; blue arrowheads, extra side shoots. (e–g) Wild-type fruits produce two locules (e), whereas the fruits of *fab* (f) and *fin* (g) mutants develop extra locules. Arrowheads, locules. (h) Quantification and comparison of floral organ numbers in wild-type, *fab* and *fin* flowers. (i–k) Stereoscope images of the primary SAM from wild-type (i), *fab* (j) and *fin* (k) plants at the transition meristem (TM) stage, before formation of the first flower. Dashed lines mark the width and height used to measure meristem size. L8, leaf 8. (l) Quantification of TM size from wild-type, *fab* and *fin* plants. Data are means \pm s.d.; $n = 25$ (h) and 9–13 (l). A two-tailed, two-sample *t* test was performed, and significant differences are represented by black asterisks: *** $P < 0.0001$. Scale bars: 1 cm (a–g) and 100 μ m (i–k).



Inflorescence branching in *fin* mutants was more severe than in *fab* mutants, and *fin* plants also developed a highly fasciated primary shoot with more side shoots than wild-type plants (Fig. 1a–d). Both mutants produced larger fruits as a consequence of additional carpels, with *fin* mutants bearing the largest fruits (Fig. 1e–g). We designated a strong allele of *fin* as a reference (*fin-e4489*, hereafter *fin* unless otherwise noted), and quantification of floral organs in the corresponding plants confirmed greater fasciation than in *fab* plants (Fig. 1h). Analyses of double mutants of *fab* and *fin* with the meristem maturation mutants *compound inflorescence* (*s*) and *anantha* (*an*), defective in the homolog of *Arabidopsis* WUSCHEL-RELATED HOMEBOX 9 and the floral-identity gene UNUSUAL FLORAL ORGANS, respectively, showed additive genetic relationships, indicating that the genes underlying *fin* and *fab* act separately from the meristem maturation pathway (Supplementary Fig. 1)¹⁴.

The phenotypes of *fab* and *fin* mutants were reminiscent of *CLV* gene mutants in *Arabidopsis*, in which inflorescence and flower fasciation are caused by enlarged shoot apical meristems (SAMs)^{7,8,10,15}. The vegetative SAMs of the *fab* and *fin* mutants, unlike those of the *s* and *an* mutants¹⁴, were larger than wild-type SAMs (Supplementary Fig. 2a–d), and this effect became more pronounced at the reproductive transition stage, just before formation of the first flower (Fig. 1i–l). We investigated meristem development in *fin* mutants using scanning electron microscopy (SEM) and found that the enlarged SAM allowed more leaf primordia and thus axillary branch meristems to develop (Supplementary Fig. 2e–l).

The phenotypic similarities between the *fab* and *fin* mutants and the fact that *fab* and *fin* double-mutant plants were more fasciated than either single mutant alone (Supplementary Fig. 3) suggested a functional link in controlling meristem size. To better understand the relationship between these mutants, we compared RNA sequencing profiles from *fab*, *fin* and wild-type vegetative primary shoot apices (Online Methods). We detected a 2-fold or greater difference in expression for 679 genes in *fab* mutants and 1,194 genes in *fin* mutants in comparison to wild-type apices (Online Methods

and Supplementary Table 1). Notably, there was significant overlap ($P < 0.001$) among the differentially expressed genes (Fig. 2a), suggesting that the *FAB* and *FIN* genes could act in the same pathway. Because *fab* and *fin* meristems were enlarged, we checked for expression changes in the homolog of *WUS* (*SIWUS*, *Solyc02g083950*)^{16,17} and a likely functional homolog of *CLV3* (*SICLV3*, *Solyc11g071380*) (Supplementary Fig. 4a–c)¹⁸. Despite a 25-fold increase in *SICLV3* expression, *SIWUS* expression doubled in *fin* meristems (Fig. 2b and Supplementary Table 1). *CLV*-*WUS* feedback was also mildly affected in *fab* mutants: *SICLV3* expression was upregulated by nearly fivefold, but *SIWUS* expression was maintained, consistent with a minor increase in vegetative meristem size (Supplementary Fig. 2a–d and Supplementary Table 1). Notably, two additional *CLV3*/EMBRYO-SURROUNDING REGION (*CLE*) genes (*SICLE3*, *Solyc02g067550*; *SICLE9*, *Solyc06g074060*) were also upregulated in *fin* mutants (Fig. 2b and Supplementary Fig. 4d,e)¹⁸. We validated the expression changes for *SIWUS* and *SICLV3* using *in situ* hybridization, which further showed substantial expansion of their expression domains in *fin* meristems, consistent with stem cell overproliferation (Fig. 2c).

FAB encodes CLV1, and FIN encodes an arabinosyltransferase

We used a mapping strategy to clone the genomic regions underlying the *fab* and *fin* mutants, using F₂ populations derived from crosses with the wild tomato *Solanum pimpinellifolium* (Fig. 2d,e and Online Methods). *fab* mapped to a 325-kb interval on the south arm of chromosome 4 where the closest homolog of *CLV1* (*Solyc04g081590*) is located (Fig. 2d and Supplementary Fig. 5a). Like *CLV1* (ref. 7), *Solyc04g081590* was expressed in all tissue types, including meristems, and sequencing its coding region in the *fab* mutant identified a missense mutation affecting the kinase domain (Fig. 2d,f and Supplementary Fig. 5b). Notably, the classical, weakly dominant-negative *clv1-9* allele encodes the same amino acid change¹⁹, and plants heterozygous for the *fab* mutation exhibited weak fasciation (Supplementary Fig. 6).

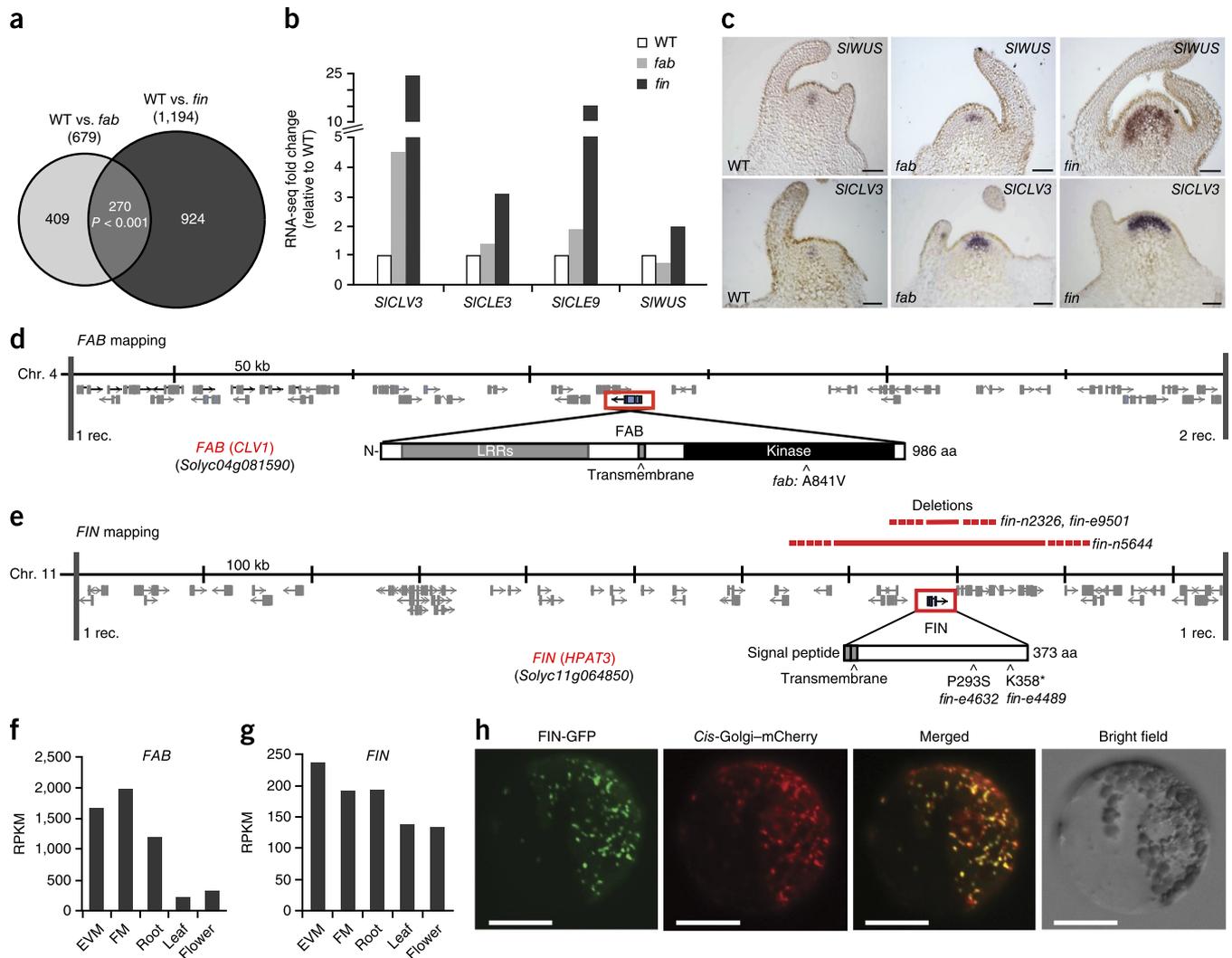


Figure 2 Molecular characterization and positional cloning of the *fin* and *fab* mutants. **(a)** Venn diagram showing significant overlap of differentially expressed genes ($P < 0.001$) in wild-type versus *fab* (light gray) and wild-type versus *fin* (dark gray) vegetative apices by RNA sequencing. **(b)** Fold change in expression of *SICLV3*, *SICLE3*, *SICLE9* and *SIWUS* in *fin* and *fab* vegetative apices relative to wild-type apices by RNA sequencing (RNA-seq). **(c)** mRNA *in situ* hybridization showing substantial expansion in *fin* (right) and mild expansion in *fab* (center) of the *SIWUS* and *SICLV3* expression domains in vegetative meristems in comparison to wild type (left). **(d,e)** The mapping intervals for *FAB* (**d**) and *FIN* (**e**). The *FAB* and *FIN* genes are boxed in red. Protein diagrams are shown with characteristic domains and motifs labeled. Open arrows indicate point mutations and corresponding amino acid changes from *fab-e0497* and two alleles of *fin*. Red horizontal lines in **e** represent three *fin* deletion alleles, which do not produce transcripts (**Supplementary Fig. 5d**). Rec., recombinant. **(f,g)** RNA sequencing read counts (RPKM) for *FAB* (**f**) and *FIN* (**g**) in major tissues, including the early vegetative meristem (EVM) and floral meristem (FM). **(h)** Subcellular colocalization of the transiently expressed FIN-GFP fusion protein with a *cis*-Golgi marker (Man9) in tomato protoplasts (Online Methods). Scale bars: 100 μm (**c**) and 10 μm (**h**).

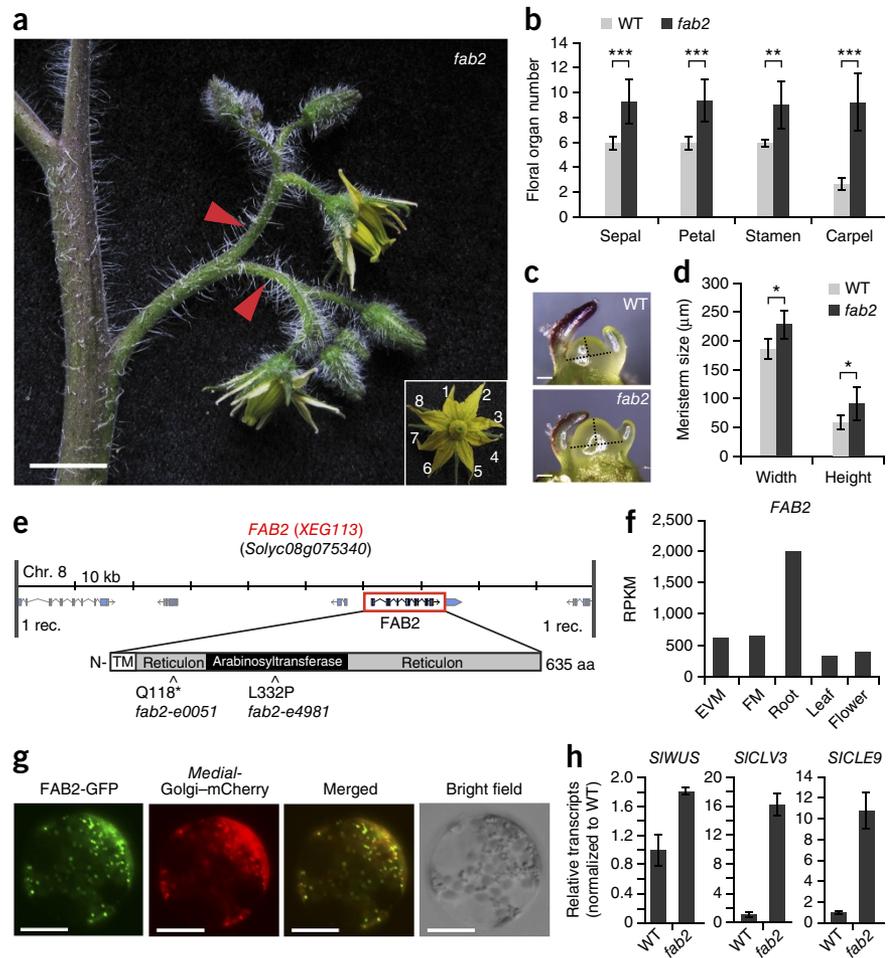
fin mapped to a 1.1-Mb region in the middle of chromosome 11 containing 71 genes. RNA sequencing identified a nonsense mutation in *Solyc11g064850*, encoding a homolog of the recently identified hydroxyproline O-arabinoxyltransferase (HPAT) protein family in *Arabidopsis* (**Fig. 2e** and **Supplementary Fig. 5c**)²⁰. Sanger sequencing of *Solyc11g064850* from four additional *fin* alleles found a missense mutation and deletions (**Fig. 2e**). RT-PCR confirmed that *Solyc11g064850* transcripts were absent in plants with the deletion alleles (**Supplementary Fig. 5d**). Thus, *FAB* encodes the tomato homolog of CLV1, and *FIN* encodes a predicted HPAT protein.

HPATs are Golgi-localized type II transmembrane proteins that are structurally similar to members of the GT8 glycosyltransferase family and catalyze the transfer of L-arabinose to the hydroxyl group of hydroxyproline (Hyp) residues²⁰. Hyp O-arabinoxylation is primarily

found in the form of linear oligoarabinoside chains on extracellular Hyp-rich glycoproteins (HRGPs), such as cell wall-associated extensins^{21–23}. There are three *HPAT* genes in *Arabidopsis*, and extensins are underarabinoxylated in *hpat3* mutants and in *hpat1* and *hpat2* double mutants²⁰. Although cell wall thickness is reduced in *hpat1* and *hpat2* double-mutant plants, perhaps accounting for a range of pleiotropic growth defects, none of the available *HPAT* gene mutants show defects in meristem size²⁰.

FIN is one of four *HPAT* gene homologs in tomato and is most similar to *HPAT3* (**Supplementary Fig. 5c**). *FIN* is expressed in all tissue types (**Fig. 2g**), and transiently expressed FIN-GFP fusion proteins in tomato protoplasts colocalized with a Golgi marker (**Fig. 2h**). To determine whether altered extensin arabinoxylation might explain *fin* mutant phenotypes, we measured the monosaccharide composition

Figure 4 The *fab2* mutant is defective in an arabinosyltransferase gene predicted to extend arabinose chains. (a) Primary inflorescence from a *fab2* mutant and a representative flower (inset). Red arrowheads mark branches. (b) Quantification and comparison of floral organ numbers in *fab2* and wild-type plants. (c) Stereoscope images of the primary SAM of *fab2* plants, showing an enlarged TM stage in comparison to wild type. Dashed lines mark the height and width used for meristem size quantification. (d) Quantification and comparison of TM meristem sizes in *fab2* and wild-type plants. (e) Positional cloning of *FAB2*. The *FAB2* gene is boxed in red, and a protein diagram is shown below. Open arrows mark the nonsense and missense point mutations from two independently derived alleles of *fab2*. (f) RNA sequencing counts (RPKM) of *FAB2* in major tissue types. (g) Subcellular colocalization of transiently expressed *FAB2*-GFP fusion protein with a medial-Golgi marker (*Mur3*) in tomato protoplasts (Online Methods). (h) qRT-PCR showing upregulation of *SIWUS*, *SICLV3* and *SICLE9* in *fab2* vegetative apices (Online Methods). Data are means \pm s.d.; $n = 13$ (b), 8 (d) and 3 (h). A two-tailed, two-sample *t* test was performed, and significant differences are represented by black asterisks: *** $P < 0.0001$, ** $P < 0.001$, * $P < 0.01$. Scale bars: 1 cm (a), 100 μ m (c) and 10 μ m (g).



Loss of an arabinosyltransferase cascade causes fasciation

By rescreening the original mutagenesis population¹³ and a new EMS-mutagenized population, we identified two alleles for *fab2*, a mildly fasciated and branched mutant that closely resembled *fab* mutants (Fig. 4a–d). Positional cloning of *fab2* identified a missense and nonsense mutation in the tomato homolog of *Arabidopsis* *XYLOGLUCANASE 113* (*XEG113*), encoding a predicted arabinosyltransferase in the GT77 family (Fig. 4e and Supplementary Fig. 11). *Arabidopsis* plants with mutations in *XEG113* display predominantly unusual diarabinoside chains on extensins, resulting in longer hypocotyls and shorter root hairs^{31,32}. In tomato, *FAB2* is expressed in all tissues and localizes to Golgi like *FIN* (Fig. 4f,g). Similar to in *fin* mutants, the expression levels of *SIWUS*, *SICLV3* and *SICLE9* were all upregulated in *fab2* SAMs, with *SICLV3* and *SICLE9* showing greater than tenfold upregulation in their expression (Fig. 4h). Thus, *FAB2*, like *FIN*, is a critical regulator of CLV signaling.

Mutations in *Arabidopsis* *REDUCED RESIDUAL ARABINOSE 3* (*RRA3*), also encoding an arabinosyltransferase in the GT77 family³³, result in shorter root hairs as a consequence of predominantly monoarabinoside chains on extensins³². This observation suggested that *RRA3* and *XEG113* act sequentially after HPATs to extend arabinoside chains^{20,32}. To test whether the tomato homolog of *RRA3* functions with *FIN* and *FAB2* in tomato CLV signaling, we mutated the closest homolog of *RRA3* (*Solyc04g080080*) using CRISPR/Cas9 (Fig. 5 and Supplementary Fig. 12)²⁹. We identified two *RRA* genes in tomato; both were most similar to *RRA3*, but only *Solyc04g080080* (hereafter referred to as *SIRRA3a*) encoded a predicted transmembrane domain (Fig. 5a). *SIRRA3a* is expressed in all tissues, and transiently expressed *SIRRA3a*-GFP fusion proteins in tomato protoplasts colocalized with a Golgi marker (Fig. 5b,c). Eight *CR-slr3a* plants

developed branched inflorescences with fasciated flowers, with these phenotypes more severe than those of the *fab2* mutants, and sequencing identified a range of indel mutations for all alleles, including a homozygous small deletion in one plant (Fig. 5d–h). Using quantitative RT-PCR (qRT-PCR), we found that *SIWUS*, *SICLV3* and *SICLE9* were all upregulated, with *SICLV3* and *SICLE9* showing more than 15-fold higher expression than in wild type (Fig. 5i). Thus, *FIN*, *RRA3a* and *FAB2* all function in the tomato CLV pathway.

A natural CLV3 mutation was selected during domestication

Our CLV pathway mutants reminded us of naturally occurring fasciation responsible for the dramatic increase in tomato fruit size during domestication³⁴. Extreme size is based on more carpels, which increases the number of seed compartments (locules). In comparison to the invariant bilocular fruits of wild tomatoes and most small-fruited cultivars, large-fruited varieties can have eight or more locules³⁴. A large portion of this variation is due to *locule number* (*lc*) and *fasciated* (*fas*), two classical mutations that have synergistic effects on locule number and thus fruit size when combined^{35,36}. The effect in *lc* mutants is weaker than in *fas* mutants and is likely due to a regulatory mutation downstream of *SIWUS*¹⁶, which has been proposed to affect binding of the tomato homolog of *AGAMOUS*, a transcription factor that downregulates *WUS* in *Arabidopsis* floral meristems³⁷. The mutant *fas* phenotype was reported to be caused by loss of expression of a *YABBY* transcription factor gene³⁸. However, the *YABBY* gene (*Solyc11g071810*) is located on chromosome 11 close to *SICLV3*, and further dissection of the *fas* locus identified a 294-kb

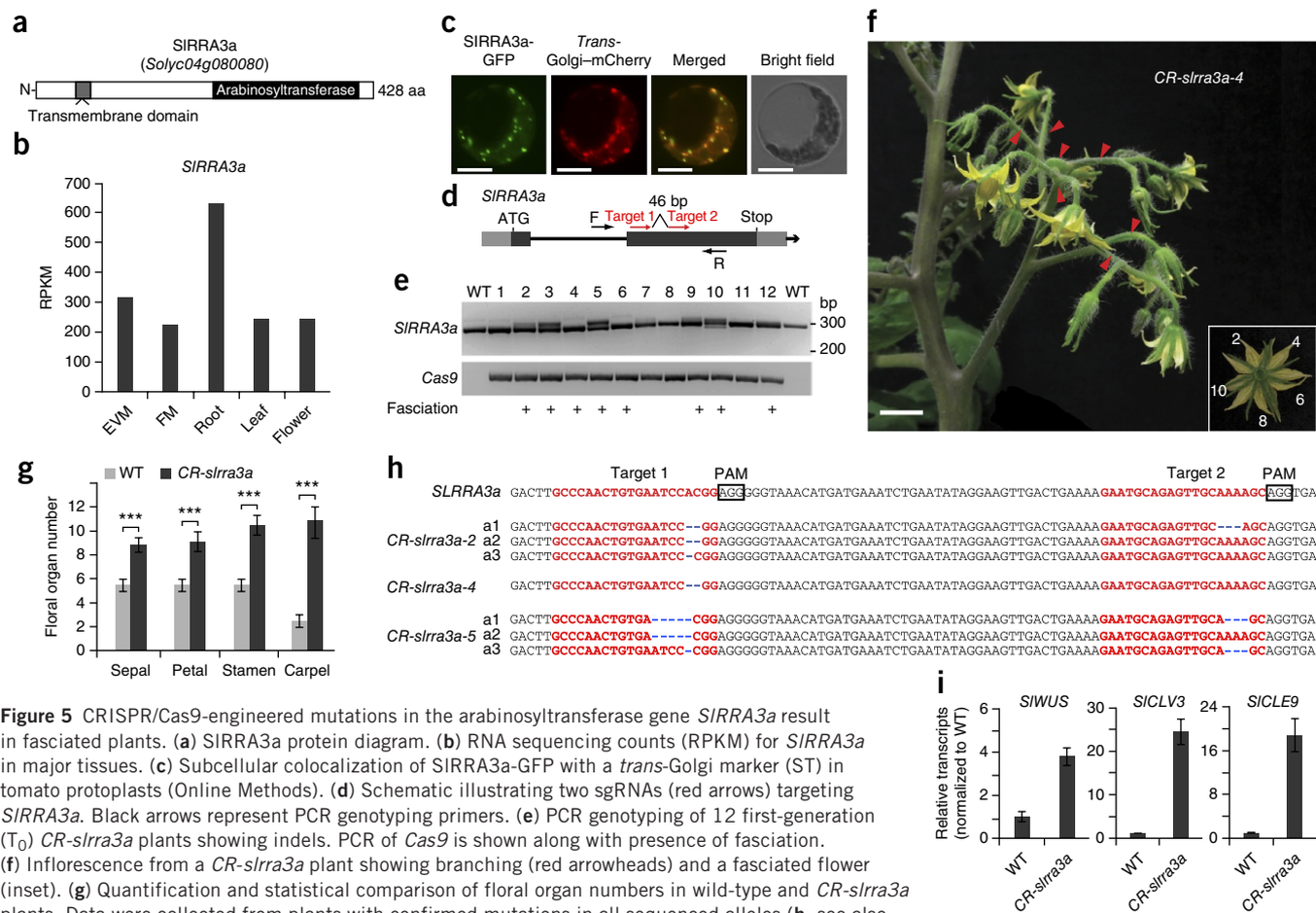


Figure 5 CRISPR/Cas9-engineered mutations in the arabinosyltransferase gene *SIRRA3a* result in fasciated plants. (a) *SIRRA3a* protein diagram. (b) RNA sequencing counts (RPKM) for *SIRRA3a* in major tissues. (c) Subcellular colocalization of *SIRRA3a*-GFP with a *trans*-Golgi marker (ST) in tomato protoplasts (Online Methods). (d) Schematic illustrating two sgRNAs (red arrows) targeting *SIRRA3a*. Black arrows represent PCR genotyping primers. (e) PCR genotyping of 12 first-generation (T_0) *CR-slr3a* plants showing indels. PCR of *Cas9* is shown along with presence of fasciation. (f) Inflorescence from a *CR-slr3a-4* plant showing branching (red arrowheads) and a fasciated flower (inset). (g) Quantification and statistical comparison of floral organ numbers in wild-type and *CR-slr3a* plants. Data were collected from plants with confirmed mutations in all sequenced alleles (h; see also Online Methods). (h) *CR-slr3a* alleles identified from three T_0 plants. *CR-slr3a-2* and *CR-slr3a-5* plants were chimeric, with multiple indels (blue dashed lines and letters). One of each representative allele identified by sequencing is shown. The *CR-slr3a-4* plant was homozygous for a 2-nt deletion. Red font highlights sgRNA targets, and black boxes indicate protospacer-adjacent motif (PAM) sequences; a, allele. (i) qRT-PCR showing upregulation of *SIWUS*, *SICLV3* and *SICLE9* in *CR-slr3a* plants. RNA was extracted from the side-shoot apices of wild-type and *CR-slr3a* T_0 plants (Online Methods). Data are means \pm s.d.; $n = 11$ (g) and 3 (i). A two-tailed, two-sample *t* test was performed, and significant differences are represented by black asterisks: *** $P < 0.0001$. Scale bars: 10 μ m (c) and 1 cm (f).

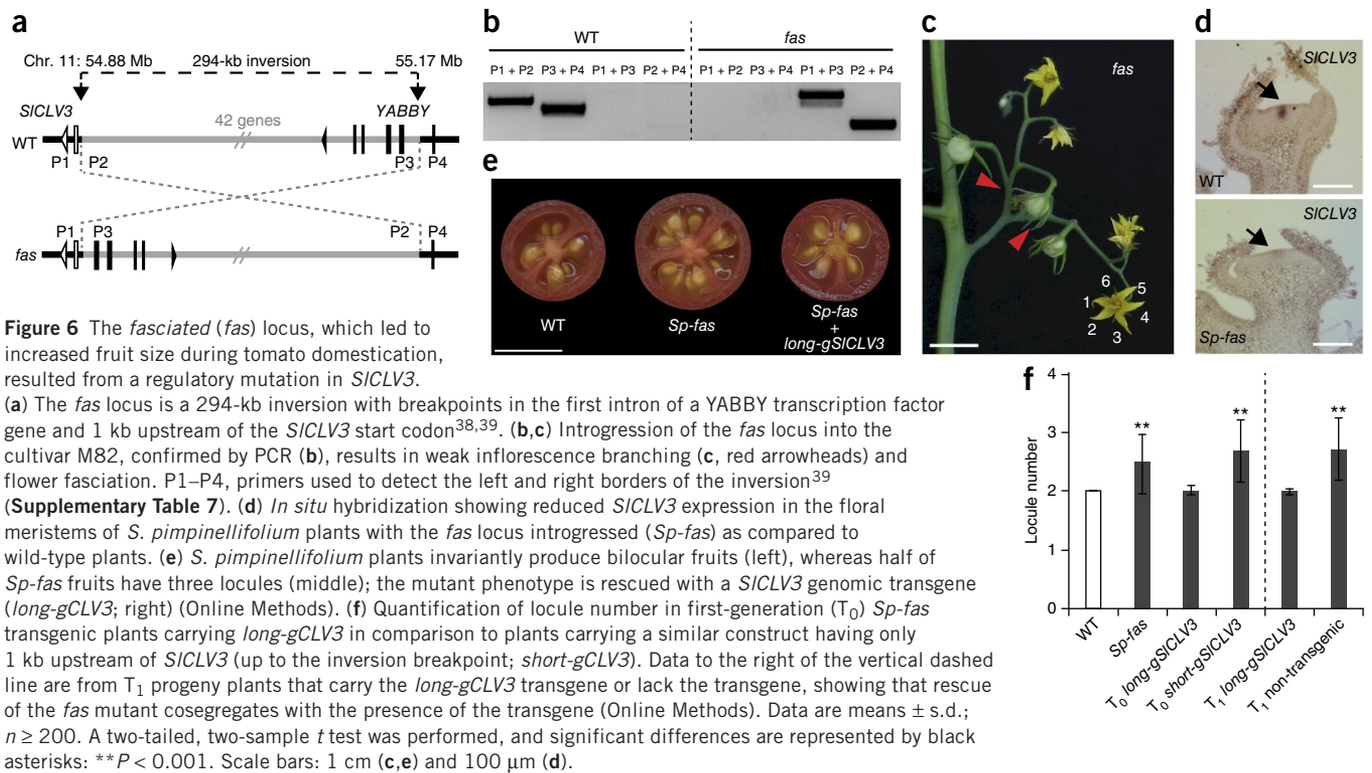
inversion with breakpoints in intron 1 of *YABBY* and 1 kb upstream of *SICLV3* (Fig. 6a,b)³⁹. This finding, in combination with weak rescue of the *fas* mutant upon introduction of functional *YABBY*³⁸, led us to ask whether *SICLV3* might also contribute to naturally occurring fasciation.

Introgression of the *fas* locus into our standard cultivated background resulted in weak branching and flower and fruit fasciation resembling the phenotype for *CR-clv2* plants (Fig. 6c and Supplementary Fig. 10). Introgression of *fas* into *S. pimpinellifolium* (*Sp-fas*) produced even weaker fasciation, with little branching and half of all fruits producing three locules, likely owing to the absence of modifier loci in the wild species background³⁷. Such a weak effect would be consistent with a change in *SICLV3* expression³⁸, and *in situ* hybridization showed reduced expression of *SICLV3* in the floral meristems of *Sp-fas* plants (Fig. 6d). To determine whether *SICLV3* underlies the *fas* mutant, we carried out transgenic complementation experiments in the *Sp-fas* genotype (Online Methods). We generated 18 plants carrying a construct comprising the *SICLV3* gene with 5.5 kb of upstream and 3.5 kb of downstream regions (*long-gCLV3*), and all plants were rescued to bilocular, smaller fruits (Fig. 6e). In contrast, 16 plants transformed with a similar construct containing 1 kb upstream of *SICLV3* (up to the inversion breakpoint; *short-gCLV3*)

did not show reduced locule number (Fig. 6f). Notably, we validated these effects in T_1 *long-gCLV3* progeny plants and found that rescue of the *fas* phenotype cosegregated with the transgene (Fig. 6f). We therefore conclude that a weak regulatory mutation in *SICLV3* underlies the *fas* mutant.

DISCUSSION

By uniting forward genetics with CRISPR/Cas9 technology²⁷, we exposed arabinosyltransferase genes as critical new components of the *CLV* pathway that are essential to control meristem size. As members of the plant and animal glycosyltransferase superfamily^{40–42}, several putative arabinosyltransferases have recently been found to function in diverse processes, including pollen tube growth by HPATs²⁰, root hair elongation by RRAs and XEG113 (ref. 32), and Rhizobia-induced nodulation by FIN homologs in pea and clover⁴³. Yet, no meristem phenotypes for any available arabinosyltransferase mutants have been reported. Our findings that arabinosyltransferase genes are critical for stem cell homeostasis suggest that members of different arabinosyltransferase families were coopted to target distinct sets of proteins and signaling peptides in different species and developmental contexts. In support of this hypothesis, unlike the predominant fasciation phenotype of *fin* mutants, we found that *fab2* and *CR-rra3a* plants



also suffer from sterility defects (data not shown). In the context of stem cell maintenance, our data imply that addition of complete arabinose chains on CLV3 and related CLE peptides by a cascade of arabinosyltransferases is required to fully activate the conserved CLV-WUS circuit. Notably, in this respect, fasciation in *fab2* mutants is weaker than in *fin* mutants and *CR-rra3a* plants appear intermediate (Supplementary Table 5), suggesting that the potency of CLE peptide signaling *in planta* increases with arabinose chain length²⁶. Thus, developmentally programmed or environmentally induced changes in arabinosyltransferase activity on CLV3 and related CLE peptides could be a mechanism to quantitatively control stem cell proliferation and meristem size.

Our findings show that disrupting the tomato CLV-WUS circuit has dramatic consequences for inflorescence architecture, flower production and fruit size. Increased fruit size was likely the first stage of tomato domestication, and, although many loci were involved³⁴, the synergistic effect of *fas* and *lc* mutations on flower and fruit fasciation may have largely been responsible for fruits that exceed 500g (refs. 35,36). In support of this idea, combining *fas* with *lc* in *S. pimpinellifolium* enhances fasciation (Supplementary Fig. 13). Our findings that *fas* is due to partial loss of *S1CLV3* expression and that a regulatory change in *SIWUS* is the likely cause of *lc*¹⁶ indicate that tomato domestication relied on subtle changes in the activity of the CLV-WUS circuit. Although the CLV-WUS circuit in *Arabidopsis* seems to be buffered against large changes in *CLV3* expression⁴⁴, *fas* mutants demonstrate that even a modest decrease in expression can cause a substantial increase in fruit size, without compromising plant growth and architecture. In this respect, it is notable that our collection of *CLV* pathway mutations provided a range of fasciation, indicating that there are multiple ways to manipulate the *CLV*-WUS circuit to quantitatively control meristem size and, thus, inflorescence architecture, flower production and fruit size. We predict that the set

of genes that can be targeted to achieve this control will extend to other LRR receptor-like genes and CLE genes operating in parallel and compensatory pathways^{45–50} (Fig. 3). In this respect, the upregulation of multiple potentially functionally redundant *SICLE* genes when *FIN* is mutated might explain the extreme fasciation in *fab* and *fin* double mutants, if higher levels of unmodified *SICLE* peptides provide some activity through *S1CLV1* (Fig. 2b and Supplementary Fig. 3). Notably, our results suggest that the ability to fine-tune *CLV* signaling could potentially be exploited to improve crop productivity. For example, in parallel to the improvement of tomato fruit size by *fas*, induced weak alleles of a *CLV2* homolog in maize can increase kernel row number⁴, and a naturally occurring *CLV3* mutation in mustard (*Brassica rapa*) could provide increased seed production⁵¹. With the widespread success of the CRISPR/Cas system in plants, the entire *CLV* pathway, now including arabinosyltransferase genes, can be targeted in many crops to test whether customized alleles can benefit breeding.

URLs. Solanaceae Genomics Network, <http://solgenomics.net/>; Solanaceae Genomics Network FTP site for genomic data deposition and access, ftp://ftp.solgenomics.net/transcript_sequences/by_experiment/lippman_lab/Liberatore_et al/; tomato tissue RNA sequencing database, http://tomatolab.cshl.edu/~lippmanlab2/allexp_query.html; ImageJ software, <http://rsb.info.nih.gov/ij/download.html>.

METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. RNA sequencing data have been deposited to the Solanaceae Genomics Network (SGN; <http://solgenomics.net/>). Data are available from the following ftp link: ftp://ftp.solgenomics.net/transcript_sequences/by_experiment/lippman_lab/Liberatore_et al/.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

C.X., K.L.L., C.A.M., Z.H., Y.-H.C., M.P., J.V.E., Y.M., E.v.d.K. and Z.B.L. designed and planned experiments. C.X., K.L.L., C.A.M., Z.H., Y.-H.C., C.B., M.O.-O., G.X. and Z.B.L. performed experiments and collected the data. C.X., K.L.L., C.A.M., Z.H., Y.-H.C., K.J., C.B., M.O.-O., G.X., M.P., Y.M., E.v.d.K. and Z.B.L. analyzed the data. K.L.L., C.X., Y.M., E.v.d.K. and Z.B.L. designed the research. C.X., K.L.L. and Z.B.L. wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare competing financial interests: details are available in the online version of the paper.

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ONLINE METHODS

Plant materials and growth conditions. Six branched and fasciated inflorescence mutants forming two complementation groups were isolated from a saturated mutagenesis population in the standard genetic background cultivar M82 (ref. 13). One complementation group, comprising a single EMS mutant, *e0497*, was designated *fasciated and branched (fab)*. The other five mutants (three EMS and two FN alleles: *fin-e4489*, *fin-e4643*, *fin-e9501*, *fin-n2326* and *fin-n5644*) were designated *fasciated inflorescence (fin)*. We rescreened the mutagenesis population and, using the same EMS mutagenesis protocol, developed and screened a new population of 3,000 M₂ families. We identified two alleles of a new mutant phenotypically similar but not complementary to *fab* that was designated *fab2 (fab2-e4891 and fab2-e0051)*. Three reference alleles, *fab-e0497*, *fin-e4489* and *fab2-e4891*, were backcrossed to the M82 parental line at least three times before generating F₂ mapping populations or double mutants. F₂ mapping populations were generated by crossing reference alleles to *S. pimpinellifolium* and self-fertilizing the F₁ plants. *fas* was backcrossed to M82 six times and to *S. pimpinellifolium* ten times, using PCR genotyping markers that identify the left and right breakpoints of the inversion³⁹. *lc* was also backcrossed to *S. pimpinellifolium* ten times. All double mutants were generated using standard crossing schemes and were confirmed by genotyping. Seeds used for meristem measurements, RNA sequencing and RNA *in situ* hybridization were pregerminated in petri dishes on Whatman paper moistened with sterile water. Seedlings with similar germination staging were transferred to soil in 72-cell plastic flats and grown in the greenhouse. Seeds used for transient expression were directly sown and germinated in soil in 72-cell plastic flats and grown in the greenhouse. Greenhouse plants were grown under natural light with supplementation from high-pressure sodium bulbs (50 mM/m²/s) on a 16-h light/8-h dark photoperiod. Daytime and nighttime temperatures were 26–28 °C and 18–20 °C, respectively.

CRISPR/Cas9 gene editing and genotyping, and phenotyping of the resulting mutants. CRISPR/Cas9 mutagenesis, plant regeneration and greenhouse care were performed as described²⁹. Briefly, constructs were designed to produce defined deletions within each target gene-coding sequence using two sgRNAs alongside the *Cas9* endonuclease gene (see **Supplementary Table 6** for a list of the sgRNAs used in this study). For genotyping of each first-generation (T₀) transgenic line, three different leaf samples were collected to capture all possible induced mutant alleles due to sectoring (chimerism), and genomic DNA was extracted using a standard cetrimonium bromide (CTAB) protocol. Each plant was genotyped by PCR for the presence of the *Cas9*-sgRNA1-sgRNA2 construct with primers designed to amplify a region spanning the 3' end of the 35S promoter and the 5' end of *Cas9*. The CRISPR/Cas9 T-DNA-positive lines were further genotyped for indel mutations using a forward primer to the left of sgRNA1 and a reverse primer to the right of sgRNA2 (**Supplementary Table 6**). PCR products from selected plants were purified for cloning into the pSC-A-amp/kan vector (Stratagene). A minimum of eight clones per PCR product were sequenced. Homozygosity for the desired deletion occurred but was rare. We typically obtained biallelic or chimeric plants in which indels occurred at one or both sgRNA target sites. Only plants in which all sequenced alleles were mutated were phenotyped, to ensure that quantification and comparison of fasciation were based on effectively null mutants. For quantification of the floral organs of the genotyped mutants, we randomly collected about 4 flowers from each mutant and counted floral organs; we then pooled the data from 3–5 different mutants to compare with the floral organs of wild-type plants grown under the same condition.

Meristem imaging. Stereomicroscope imaging of meristems was performed as described previously⁵². Briefly, hand-dissected tomato meristems were captured on a Nikon SMZ1500 microscope. For SEM, meristems were dissected by hand and passed through a dehydrating ethanol series, dried using a critical-point dryer (Tousimis), coated in gold particles and imaged under a Hitachi S-3500N microscope.

Transmission electron microscopy. Tomato vegetative meristems from 9-d-old seedlings were fixed in 2% glutaraldehyde and 2% paraformaldehyde in 0.1 M PBS overnight and then post-fixed in 1% osmium tetroxide in 1.5%

potassium ferrocyanide. Samples were dehydrated in a graded alcohol series followed by 100% dry acetone. Samples were infiltrated with agitation in 50% Epon Araldite resin in 100% acetone for 2 h and then overnight in 100% resin. Samples were flat embedded in resin in BEEM capsules from which the conical ends had been removed and were capped with the flat lid of the capsule. Plastic blocks were carefully trimmed and positioned for sectioning to obtain longitudinal sections through the meristem region. Thin sections were collected on Butvar-coated 1 × 2 mm nickel slot grids (Veco, Electron Microscopy Sciences) and counterstained in lead citrate stain. Samples were examined with a Hitachi H700 transmission electron microscope at 75 kV and were recorded on Kodak 4489 film that was then scanned at 2,400 dpi using an Epson Perfection V750 Pro scanner.

Meristem transcriptome profiling. Vegetative meristem collection, RNA extraction and library preparation for wild-type, *fin* and *fab* plants were performed as described⁵². Briefly, total RNA was extracted from 30–50 meristems per biological replicate using a PicoPure RNA Extraction kit (Arcturus). mRNA was purified from 1–5 µg of total RNA for library construction using the ScriptSeq v2 RNA library preparation kit (Epicentre) with barcodes. The quantity and size distribution of each individual RNA sequencing library were detected on a Bioanalyzer 2100 (Agilent Technologies). Two biological replicates per genotype were generated. Five barcoded libraries were pooled with relatively equal concentrations (determined by qRT-PCR quantification) for one lane of Illumina paired-end 100-bp sequencing on an Illumina HiSeq sequencing machine. See methods of data analysis in the **Supplementary Note**.

Tissue-specific expression patterns for *SICLV3*, *SICLE3*, *SICLE9*, *FAB*, *FIN*, *FAB2* and *SIRRA3a* were obtained from the tomato tissue RNA sequencing database. RNA sequencing data from different tissues (for example, in **Fig. 2f,g**) were mined from the tomato genome project transcriptome profiling data sets deposited in the Sequence Read Archive (SRA) under accession [SRP010775](https://www.ncbi.nlm.nih.gov/sra/SRP010775) (ref. 53) and from our meristem maturation atlas⁵².

In situ hybridization. RNA *in situ* hybridization was performed using standard protocols⁵⁴, with slight modifications. Briefly, full-length coding sequences for tomato *SIWUS (Solyc02g083950)*, *SICLV3 (Solyc11g071380)* and *FAB (Solyc04g081590)* were amplified from M82 cDNA using Phusion Taq (Invitrogen), and the resulting products were ligated into the StrataClone pSC-A-amp/kan vector (Agilent Technologies). Plasmids were linearized and, depending on the orientation of the gene insert, T7 or T3 RNA polymerase was used for *in vitro* transcription. Full-length probes were used for hybridization. SAM stages were defined as described previously⁵². For fixation, meristems were dissected by hand and fixed in 4% paraformaldehyde with 0.3% Triton X-100 under vacuum for 20 min at 400 mm Hg.

Map-based cloning. Markers for recombinant mapping and genotyping were generated using SNPs and indels identified in comparison of *Solanum lycopersicum* and *S. pimpinellifolium*⁵³. By using 76 F₂ mutants and molecular markers derived from the *S. pimpinellifolium* genome sequence, we mapped the locus for *fab-e0497* to a 325-kb interval containing 45 genes on the bottom of chromosome 4, including *Solyc04g081590*, the closest homolog of *CLV1*. Using a similar approach with 624 F₂ mutants, we mapped the locus for the *fin-e4489* mutant to a 1-Mb region in the middle of chromosome 11 containing 71 genes. As a lack of recombination prevented further fine mapping, meristem-enriched mRNA was extracted from *fin-e4489* for deep cDNA sequencing (mRNA-seq). A total of 42 million paired-end 100-bp Illumina sequencing reads were aligned to the 1-Mb interval, and a single nonsense mutation was detected in *Solyc11g064850*. Sanger sequencing of *fin-e4632* identified a missense mutation in the same gene. The other *fin* alleles (*fin-n2326*, *fin-e9501* and *fin-n5644*) could not be amplified by PCR, indicating that there were large deletions encompassing the *Solyc11g064850* gene location, which was confirmed by RT-PCR. Using 42 F₂ mutants, we mapped the locus for the *fab2-e4981* mutant to a 100-kb region on the bottom of chromosome 8 containing 7 genes. Sanger sequencing identified a missense mutation in *Solyc08g075340*. Sequencing of the second allele of *fab2*, *fab2-e0051*, identified a nonsense mutation in *Solyc08g075340*.

Transient expression in tomato protoplasts. To generate the transient expression constructs for analysis of subcellular localization, eGFP was fused to the C termini of FIN, FAB2 and SIRRA3a. To produce a series of Golgi markers, mCherry was fused to the C termini of Man49 (*Arabidopsis* α -1,2 mannosidase I-49)⁵⁵, MUR3 (*Arabidopsis* β -1,2-galactosyltransferase)⁵⁶ and ST (*Rattus norvegicus* α -2,6-sialyltransferase)⁵⁷ to create *cis*-, *medial*- and *trans*-Golgi markers, respectively. Fusion protein expression was driven by the cauliflower mosaic virus (CaMV) 35S promoter in transient expression vectors, as described previously⁵⁸. The transient expression protocol was modified from the *Arabidopsis* mesophyll protoplast system⁵⁹. Well-expanded leaves from 10- to 14-d-old tomato plants were cut into 0.5- to 1-mm leaf strips and digested with enzyme solution (1.5% cellulase R10 (Yakult Honsha), 0.4% macerozyme (Yakult Honsha), 20 mM MES (pH 5.7), 0.4 M mannitol, 20 mM KCl, 10 mM CaCl₂ and 0.1% BSA) in a 90-mm Petri dish for 4 h in the dark with gentle shaking at 40–50 rpm. The digestion products were filtered using 75- μ m nylon mesh (Fisher) in a 50-ml Falcon tube. Cells were collected by centrifugation at 100g for 2 min. The cells were washed by resuspending them in precooled W5 solution (2 mM MES, pH 5.7, 125 mM CaCl₂, 154 mM NaCl, 0.1 M glucose and 5 mM KCl), and cells were finally resuspended in MMG solution (4 mM MES, pH 5.7, 0.5 M mannitol and 15 mM MgCl₂) to a concentration at 2×10^5 cells/ml for immediate transfection. For transfection, 5–10 μ g of plasmid of 5–10 kb in size was added to 100 μ l of protoplast cell suspension in a 2-ml microfuge tube, and samples were then gently mixed well with an equal volume of PEG solution (40% PEG in 0.2 M mannitol and 100 mM CaCl₂). The transfection mixtures were incubated at room temperature for 10 min, and cells were then washed with W5 solution. Cells were resuspended in W5 solution and incubated at 28 °C for 12 h before collection for microscopy or other experiments. Imaging was performed on a Zeiss Axioplan 2 fluorescence microscope with an AxioCam camera.

Monosaccharide analysis of the cell wall. Preparation and analysis of cell wall material (alcohol-insoluble residue, or AIR) was performed as described⁶⁰. Neutral sugars and uronic acids from non-cellulosic polysaccharides in AIR were released by hydrolysis with trifluoroacetic acid (TFA). Approximately 1 mg of AIR was hydrolyzed with 250 μ l of 2 M TFA at 121 °C for 90 min. The TFA-soluble material was split into two equal parts, one for uronic acid determination and one for neutral sugar determination. Uronic acid content was determined by subjecting the hydrolyzate to a CarboPac PA200 anion-exchange column with an ICS-3000 Dionex chromatography system. The elution profile consisted of a linear gradient of 50–200 mM sodium acetate in 0.1 M NaOH in 10 min at 0.4 ml/min. Neutral sugar composition was determined by derivatization with alditol acetate as described⁶¹. In brief, monosaccharides in the hydrolyzate were reduced with NaBH₄ (20 mg/ml dissolved in 1 M NH₄OH) at room temperature for 1 h and then peracetylated with acetic anhydride and pyridine at 121 °C for 20 min. The alditol acetates generated were separated on a gas chromatograph equipped with a Supelco SP2380 column (Sigma-Aldrich) and analyzed by a flame ionization detector (FID) (Agilent Technologies, 7890A).

Sequential extraction of cell wall polymers. Cell wall polymers were sequentially extracted as described previously³¹. Freeze-dried plant materials were ground to fine powder. About 5 mg of ground material was extracted overnight in 50 mM ammonium formate buffer, pH 4.5, at 37 °C with shaking at 200 rpm. After incubation, the samples were centrifuged for 10 min at 20,800g, and the supernatant was collected. The remaining pellet was washed three times with 1 ml of water per wash and centrifuged, and the resulting supernatants were pooled with the supernatant from the first spin. The supernatant was precipitated overnight in 80% ethanol and centrifuged for 10 min at 20,800g. The precipitated fraction was freeze dried, representing the buffer extract. The buffer-insoluble pellet was digested with 0.02 U pectin methylesterase (EC3.1.1.11, Novozymes) and endopolygalacturonanase M2 (EC3.2.1.15, Megazyme) in 1 ml of 50 mM ammonium formate buffer (pH 4.5) overnight at 37 °C. After digestion, the samples were treated for 10 min at 99 °C and centrifuged for 10 min at 20,800g, and the supernatant was collected as the EPG/PME fraction (pectin extract). The remaining pellet was digested with 10 U endoglucanase II (EGII, Megazyme) in 1 ml of 50 mM ammonium

formate buffer (pH 4.5) overnight at 37 °C, heat inactivated for 10 min at 99 °C and centrifuged for 10 min at 20,800g; the supernatant was collected as a hemicellulosic/cellulosic extract. The remaining pellet was treated with 1 ml of saturated Ba(OH)₂ (~0.22 M) solution for 6 h at 105 °C. The pellet resulting from centrifugation was washed three times with 1 ml of water per wash and centrifuged. The resulting supernatants (extensin fraction) were pooled and neutralized with 1 M sulfuric acid, and the precipitating salts were removed by centrifugation. The remaining pellet (residue) and all fractions were freeze dried and analyzed for monosaccharide composition as described above.

In planta peptide treatment of tomato apices. Peptide treatment of tomato apical meristems was modified from previous methods^{26,62}. Sterilized tomato seeds were distributed onto two layers of Whatman paper moistened with sterile water in a 180-mm petri dish. The seeds were pregerminated at 28 °C for 1–2 d. Seeds with similar staging were selected for further growth in a 50-ml Falcon tube (10 seeds/tube) with 10 ml of liquid medium containing half-strength Murashige and Skoog salt mixture (Sigma-Aldrich), 1% sucrose and 0.5 g/l MES (pH 5.8) and supplemented with different concentrations of peptides. The Falcon tubes were incubated on a roller bank with a 16-h light/8-h dark photoperiod at 28 °C for 10 d. Chemical synthesis of arabinosylated SICLE peptides was performed as described previously²⁶.

Tomato shoot apical meristem imaging and size measurement of treated meristems. After peptide treatment, SAMs were cleared for imaging and size quantification. Individual SAMs from each treatment were dissected and fixed in FAA (1% formaldehyde, 0.5% acetic acid and 50% ethanol) overnight at 4 °C. The fixed SAMs were sequentially washed with 70%, 85% and 100% ethanol for 30 min each wash and dehydrated for 60 min in 100% ethanol; they were then immersed in an ethanol:methyl salicylate solution (1:1) for an additional 60 min. Fixed tissue was transferred to methyl salicylate (100%) for storage before microscopy. Slides were sealed with Vaseline to keep the SAMs immersed in methyl salicylate. Imaging by Normarsky illumination was conducted using a Leica DMRB microscope (Leica Microsystems). Micrographs were acquired with a Leica MicroPublisher 5.0 RTV digital camera system using Q-IMAGING software. All images were taken at the focal plane corresponding to the median optical section of the SAM. The size of each SAM was determined by outlining and calculating the meristem area between the basal edges of two opposite leaf primordia using ‘freehand selections’ and ‘measure’, respectively, in ImageJ software. The peptide treatment assays were performed three times with similar results.

Quantitative RT-PCR. qRT-PCR was performed as described previously⁵². Briefly, total RNA from vegetative meristems and shoot apices was extracted with the PicoPure RNA Extraction kit and the RNeasy mini kit (Qiagen), respectively. One microgram of total RNA was treated with DNase I and used for cDNA synthesis with a SuperScript III reverse-transcriptase kit (Invitrogen). qRT-PCR was performed using gene-specific primers in the iQ SYBR Green SuperMix (Bio-Rad) reaction system on the CFX96 Real-Time system (Bio-Rad), following the manufacturer’s instructions (**Supplementary Table 7**). The tomato *UBIQUITIN* gene was used as an internal control.

Constructs for transformation and complementation of *Sp-fas*. The binary vector (pHaoNM) used in transgenic complementation of *Sp-fas* was modified from the standard vector pCAMBIA1300. The two constructs used for complementation tests were generated through the following steps. First, fosmid SL_FOS0151E08 from tomato accession Heinz 1706 was digested at MfeI and SacI sites to obtain a 4,320-bp fragment containing the *SICLV3* gene. This fragment was then ligated into pHaoNM at the EcoRI and SacI sites (pHC1). A 5,233-bp fragment upstream of the *SICLV3* gene was isolated from SL_FOS0151E08 by digesting at SacI and NsiI sites. This fragment was ligated into pHC1 at the SacI and SbfI sites to create the complementation construct named *long-gCLV3*, containing the *SICLV3* gene with 5,465 bp upstream of the start codon and 3,445 bp downstream of the stop codon. For the control construct *short-gCLV3*, a PCR product was amplified with primers 12EP315 and 12EP389 (**Supplementary Table 7**). The PCR product was then digested with SacI and PstI and inserted into pHC1 at the SacI and SbfI sites. The resulting

short-gCLV3 construct contained the *SICLV3* gene, 1,008 bp upstream of the start codon of *SICLV3* (up to the *fas* inversion breakpoint) and 3,445 bp downstream of the stop codon.

To test whether *SICLV3* was responsible for the *fas* phenotype, *long-gCLV3* and *short-gCLV3* were introduced separately into *Sp-fas* using *Agrobacterium*-mediated transformation (*Agrobacterium tumefaciens* strain LBA4404). We generated 18 first-generation (T_0) transgenic plants, which all carried 1 or 2 insertions of *long-gCLV3*. All 18 plants were used for phenotypic analysis of fruit locule number ($n = 200$). We also generated 18 T_0 plants carrying *short-gCLV3*, and we phenotyped 16 plants that carried 1 or 2 insertions. Two plants that were found by Southern blot to carry four and six copies of the transgene showed evidence of cosuppression and were therefore excluded from analysis. A more detailed analysis of complementation of *fas* with *long-gCLV3* was performed in T_1 progeny plants derived from a T_0 plant with a single transgene insertion. Rescue of bilocular fruit cosegregated with the presence of the *long-gCLV3* transgene, whereas *short-gCLV3* had no effect (Fig. 6). Genotyping for the presence of the transgene in T_1 progeny plants was performed using PCR primers that amplified a fragment of the pCAMBIA1300 backbone.

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