**LYRATE** Is a Key Regulator of Leaflet Initiation and Lamina Outgrowth in Tomato

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Development of the flattened laminar structure in plant leaves requires highly regulated cell division and expansion patterns. Although tight regulation of these processes is essential during leaf development, leaf shape is highly diverse across the plant kingdom, implying that patterning of growth must be amenable to evolutionary change. Here, we describe the molecular identification of the classical tomato (*Solanum lycopersicum*) mutant *lyrate*, which is impaired in outgrowth of leaflet primordia and laminar tissues during compound leaf development. We found that the *lyrate* phenotype results from a loss-of-function mutation of the tomato *JAGGED* homolog, a well-described positive regulator of cell division in lateral organs. We demonstrate that **LYRATE** coordinates lateral outgrowth in the compound leaves of tomato by interacting with both the **KNOX** and auxin transcriptional networks and suggest that evolutionary changes in **LYRATE** expression may contribute to the fundamental difference between compound and simple leaves.

**INTRODUCTION**

Leaves are the principal organs produced during vegetative growth and provide a flattened surface allowing for efficient light capture during photosynthesis. Leaves begin at the flanks of the shoot apical meristem (SAM) as peg-like outgrowths that rapidly expand along the proximal/distal axis via cell division. As elongation proceeds, the archetypal dicot simple leaf begins lateral expansion of the blade from a continuous band of meristematic cells at the edge of the leaf primordium termed the marginal blastozone (Hagemann and Glesissberg, 1996). Such leaves are divided into two major developmental domains: the distal blade, where lateral growth is initiated, and the proximal petiole, which lacks blade outgrowth. Blade outgrowth continues until a wave of cell cycle arrest usually begins at the distal tip of the developing leaf and proceeds to the leaf base (Avery, 1933; Poethig, 1984; Nath et al., 2003). The remaining growth required to produce a mature leaf is accommodated by cell expansion. Cell division patterns during early leaf development may, in part, guide subsequent expansion, but it is well established that disruption in cell division patterns can lead to changes in expansion and roughly equivalent final organ shape (Tsukaya, 2006). It is therefore hypothesized that cell division-dependent and -independent pathways may dictate final organ shape. In simple leafed model species, such as *Arabidopsis thaliana* and snapdragon (*Antirrhinum majus*), these processes result in a flattened, continuous lamina subtended by a bladeless petiole.

Although coordination of cell division and expansion are critical factors directing continuous laminar development, simple leaves represent only a fraction of the total diversity in plant leaf morphology. Many plants produce leaves that are deeply lobed or are compound, consisting of multiple blade units termed leaflets. In compound leaves, the marginal blastozone initiates leaflet primordia that are discriminated from adjacent intercalary tissues by rapid cell division (Hagemann and Glesissberg, 1996). As leaf development progresses, laminar outgrowth is tightly regulated such that intercalary tissues and leaflet bases are bladeless, ultimately resulting in discrete leaflets (Kaplan, 1975). This pattern can be reiterated to produce secondary and tertiary leaflets. Thus, in comparison to simple leaves, compound leaves must undergo highly patterned growth during several stages of leaf development. Our understanding of the mechanism by which these subdomains within the compound leaf are specified remains in its infancy.

The plant hormone auxin, which is known to positively influence both cell division and expansion, has recently been shown to be important in patterning differential growth in several compound leaved species (DeMason and Chawla, 2004; Wang et al., 2005; Barkoulas et al., 2008; DeMason and Polowick, 2009; Koenig et al., 2009). In tomato (*Solanum lycopersicum*) and *Cardamine hirsuta*, auxin is transported into incipient leaflets where it activates response pathways leading to primordium outgrowth (Barkoulas et al., 2008; Koenig et al., 2009). Furthermore, this differential auxin distribution is critical to pattern blade outgrowth such that equalization of auxin distribution in the primordium results in ectopic blade outgrowth in intercalary tissues between leaflets (Koenig et al., 2009). This repression of auxin response in bladeless intercalary tissues of wild-type tomato compound leaves is accomplished by the auxin/indole-3-acetic acid (IAA) gene **ENTIRE** (Wang et al., 2005; Zhang et al., 2007). **ENTIRE** mutants initiate ectopic blade formation between leaflets leading to simple leaf development (Koenig et al., 2009). A similar phenotype is observed in the **goblet** mutant in tomato...
The sequence conservation between the JAG and S. lycopersicum JAG proteins was low outside of the conserved motifs, making it crucial to determine the extent to which the two proteins are functionally similar. To resolve this question, we examined the ability of S. lycopersicum JAG to suppress the floral loss-of-function phenotype seen in jag-2 and jag-3 Arabidopsis plants (Ohno et al., 2004) by transforming both mutants with the S. lycopersicum JAG genomic fragment driven by the Arabidopsis JAG promoter (PAtJAG::SlJAG). The floral phenotype of jag-2 was fully suppressed in 36 and partially suppressed in 16 out of 124 jag-2 transgenic plants (Figure 1B). Similar complementation was seen in the jag-3 background. In addition, we compared the effect of S. lycopersicum JAG overexpression in the Arabidopsis wild-type, jag-2, and jag-3 genetic backgrounds with the published effects of Arabidopsis JAG overexpression (Dinneny et al., 2004; Ohno et al., 2004) (see Supplemental Figure 3 online). The 35S:SlJAG transgenics displayed phenotypes highly reminiscent of JAG overexpressing Arabidopsis in all attributes except the elaboration of cryptic bracts. These results demonstrate that, despite substantial divergence in amino acid sequence, Arabidopsis JAG and S. lycopersicum JAG show a high degree of functional homology. This being said, S. lycopersicum JAG overexpression did not result in the expansion of cryptic bracts as seen in experiments using JAG, suggesting that some of this protein sequence divergence may have functional consequences.

lyrate Is an S. lycopersicum JAG Null Mutant

We next placed JAG on the tomato genetic map using a segregating S. lycopersicum × Solanum pennellii F2 population (Frary et al., 2004) and found that JAG maps to a 2-centimorgan region on the short arm of chromosome 5, between markers T1335 and CD64. Two recessive leaf shape mutants, lyrate (lyr) and trifoliata (tf), roughly map to this region of chromosome 5. We hypothesized that loss of JAG function might be responsible for one of these mutant phenotypes.

To determine if a mutation in S. lycopersicum JAG causes either lyr or tf phenotypes, we analyzed the level of JAG transcript in each mutant. RT-PCR analysis showed complete absence of JAG transcript in lyr relative to VF36 wild-type, tf, and pooled lyr/+ and +/+ segregants (Figure 1C). Confirmation of these results with quantitative RT-PCR analysis suggested that JAG expression in lyr is significantly reduced compared with the wild type (Figure 1D). To determine whether this change of expression was due to mutation of the JAG gene, we preformed DNA gel blot hybridization using a JAG probe on lyr and wild-type segregants. A restriction fragment length polymorphism at the 5’ end of JAG was detected in lyr when compared with wild-type hybridization (see Supplemental Figure 4 online). Sequence analysis of the JAG gene in lyr plants revealed a 509-bp deletion starting at position −265 and ending in intron 2 (Figures 1B and 1E). This deletion greatly reduces JAG transcription and likely interrupts translation of the JAG protein.

The lyr mutant (Figures 2A and 2B) was originally characterized by the fan-like appearance of its leaves. To verify that the mutation in JAG is the cause of the lyr phenotype, we examined JAG expression and sequence from two other lyr alleles, which

et al., 2004). This phenotype is greatly enhanced in jag nub double mutants that display a large number of developmental defects, including deeply lobed leaves (Dinneny et al., 2006). Plants overexpressing JAG display ectopic blade outgrowth on leaf petioles and expansion of cryptic bracts, presumably by overcoming inhibition of cell division in these regions (Dinneny et al., 2004; Ohno et al., 2004). Thus, JAG and NUB are thought to pattern cell cycle activity in young leaf primordia to differentiate the bladed distal tissues and the bladeless proximal tissues. Based on this described role in Arabidopsis, we hypothesized that changes in JAGGED expression might contribute to differential growth in developing compound leaves.

RESULTS

Identification of the Tomato JAG Ortholog

First, we identified the S. lycopersicum JAG homolog and showed it encodes a putative protein composed of 259 amino acids. Three conserved domains were readily identified in the predicted S. lycopersicum JAG sequence when compared with Arabidopsis JAG (Dinneny et al., 2004; Ohno et al., 2004): a putative nuclear localization signal sequence, a single C2H2-type zinc finger motif, and a Pro-rich motif (Figure 1A; see Supplemental Figure 1 online). Another motif, identified previously in Arabidopsis as an EAR motif (Ohno et al., 2004), was identical between Arabidopsis JAG and S. lycopersicum JAG (Figure 1A; see Supplemental Figure 1 online). Outside of these described domains, amino acid sequence conservation between S. lycopersicum JAG and Arabidopsis JAG was low, indicating possible functional dissimilarities (see Supplemental Figure 1 online). Nevertheless, the intron-exon structure of S. lycopersicum JAG (3134 bp) is completely conserved relative to Arabidopsis JAG (Figure 1B). DNA gel blot analysis demonstrated that S. lycopersicum JAG is a single-copy gene (see Supplemental Figure 2 online). In combination, these data demonstrate that S. lycopersicum JAG is a single-copy homolog of the Arabidopsis JAG gene.

(Brand et al., 2007; Blein et al., 2008; Berger et al., 2009). GOB encodes an ortholog of the Arabidopsis CUC2 gene, which specifies organ boundaries in the SAM (Aida et al., 1997). In several compound leafed species, GOB specifies the boundaries between initiating leaflets (Blein et al., 2008; Berger et al., 2009). The relationship between auxin distribution, ENTIRE, and GOB remains unclear at this point, but auxin distribution and response is central to the patterning of growth in compound leaves.

To gain insight into how molecular regulation of cell division has been modified to generate differential growth in compound leaves, we cloned and analyzed the function of the JAGGED ortholog in tomato. In the simple leaved species Arabidopsis, JAGGED (JAG) and its paralog NUBBIN (NUB) are important positive regulators of cell division and outgrowth in lateral organs (Dinneny et al., 2004, 2006; Ohno et al., 2004). Both genes encode transcription factors consisting of a single C2H2-type zinc finger and a Pro-rich motif. JAG loss-of-function causes premature cessation of cell proliferation in lateral organs, ultimately resulting in irregular margins (Dinneny et al., 2004; Ohno et al., 2004). This phenotype is greatly enhanced in jag nub double mutants that display a large number of developmental defects, including deeply lobed leaves (Dinneny et al., 2006). Plants overexpressing JAG display ectopic blade outgrowth on leaf petioles and expansion of cryptic bracts, presumably by overcoming inhibition of cell division in these regions (Dinneny et al., 2004; Ohno et al., 2004). Thus, JAG and NUB are thought to pattern cell cycle activity in young leaf primordia to differentiate the bladed distal tissues and the bladeless proximal tissues. Based on this described role in Arabidopsis, we hypothesized that changes in JAGGED expression might contribute to differential growth in developing compound leaves.
show similar phenotypes. lyr² (Figure 2C), a spontaneous allele, showed a normal level of JAG transcription (Figure 1F), but sequence analysis revealed a single nucleotide polymorphism (T to C) that results in substitution of the first highly conserved Cys in the C₂H₂-type zinc finger domain with an Arg (see Supplemental Figure 1 online, red arrowhead). lyr³ (n2731, Figure 2D) is derived from a fast neutron mutagenesis project, (Menda et al., 2004) (Y. Eshed, unpublished data) and showed a strong reduction in JAG transcript levels (Figure 1F). DNA gel blot analysis of lyr² revealed a large insertion at the JAG locus (Figure 1B; see Supplemental Figure 4 online). Finally, to further confirm that reduction in JAG RNA results in a lyr-like phenotype, we generated RNA interference (RNAi) transgenics targeting the JAG locus. One of these transgenic lines exhibited phenotypes highly similar to that of lyr (1/16 transgenic plants; Figure 2E). We therefore named the tomato JAG homolog LYR.

The lyr Phenotype

The naturally occurring mutant lyr was isolated from cultivated tomato in Central America. It is characterized by a broad reduction in outgrowth of lateral organs. The mature leaves of wild-type tomato plants are bipinnate, consisting of a serrated terminal leaflet, three or more lateral leaflet pairs, and a few intercalary leaflets (Figure 2A, Table 1). The base of each of the large primary leaflets will give rise to one or two secondary leaflets. By contrast, the adult lyr leaf consists of a deeply divided terminal leaflet, one or two lateral leaflets, and few intercalary leaflets (Figures 2B to 2D). All of these leaflet types exhibited reduced laminar outgrowth when compared with wild-type plants. Interestingly, secondary leaflet number on the first lateral leaflet pair was increased in all three lyr alleles (Figures 2B to 2D, Table 1). Increased dissection to produce tertiary leaflets was observed in lyr² alleles but not in the lyr³ allele (Figures 2B to 2D, Table 1). lyr² plants were in general more inhibited in blade outgrowth and leaflet outgrowth than the other alleles, suggesting possible differences in allelic strengths. Nevertheless, all three lyr alleles exhibited severely reduced outgrowth of laminar tissues in all leaflets (Figures 2B to 2D, Table 1).

In addition to the dramatic leaf phenotype, lyr floral development is abnormal. lyr flowers open prematurely and have reflexed petals (see Supplemental Figures 5A to 5C online). Wild-type tomato anthers are fused to form a cone surrounding the carpel, but lyr anthers are narrow and unfused curving inwardly toward the carpel (see Supplemental Figures 5D and 5E online). lyr carpels are composed of a thickened style and an enlarged ovary containing disorganized ovules (see

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Figure 1. Molecular Characterization of S. lycopersicum JAG.

(A) Schematic of conserved domains of JAG. The four conserved regions are indicated in gray boxes (not to scale).

(B) JAG gene structure showing five exons (gray boxes) and untranslated regions (UTRs; white boxes). The deletion in the lyr allele is designated as a black line with the flanking nucleic acids. The estimated position of the putative insertion in the lyr³ allele is designated as well as the location of the point mutation in the lyr² allele (not to scale). The arrows a and b indicate primer position for RT-PCR analysis described in (C), and the arrows c and d indicate primer position for quantitative RT-PCR analysis described in (D). Photo on the right shows flowers (from left to right) of wild-type, jag-3, and AtJAG:gSUAG in jag-3, demonstrating rescue of the jag-3 flower phenotype by S. lycopersicum JAG. Bar = 1 mm.

(C) RT-PCR analysis of S. lycopersicum JAG in wild-type VF36, lyr, and a mix of lyr² and wild-type plants. Each sample represents RNA that was isolated from two apices from two different plants. Two samples were tested for each line. Negative control was a PCR reaction without a template (NT), and actin mRNA was amplified as a positive control for RNA integrity.

(D) Quantitative RT-PCR analysis of the expression of JAG in wild-type and lyr tomato plants. Error bars indicate SE over three technical replicates.

(E) PCR on genomic DNA of wild-type and lyr showing smaller JAG PCR product in lyr (numbers on left indicate band size in base pairs).

(F) RT-PCR analysis of SI JAG in lyr² and lyr³. Two samples were tested for lyr² and one for lyr³. Negative control was a PCR reaction without a template (NT), and actin was amplified as a positive control.

[See online article for color version of this figure.]
Supplemental Figures 5F to 5H online). These floral phenotypes are reminiscent of those observed in jag nub mutants (Dinneny et al., 2006). Thus, the function of the LYR gene seems to be conserved with that of JAG and NUB as a broad regulator of aerial lateral organ development. Although the Arabidopsis jag, nub, or jag nub mutants are compromised in leaf development, lyr exhibits an even more dramatic leaf phenotype relative to wild-type tomato. This suggests that tomato compound leaf development is more sensitive to the disruption of LYR/JAG function than Arabidopsis simple leaf development.

Leaf Development in lyr

We examined early leaf development in lyr to determine the origin of its leaf shape alteration. lyr differed significantly from wild-type plants from the earliest stages of leaf development. Wild-type tomato leaves begin as peg-like structures that curve toward the SAM due to increased cell division on the abaxial leaf face relative to the adaxial face. Even in these young leaves (P2-P3, primordia numbered starting from the first visible primordia at the shoot apex) differentiation is apparent at the tip of the leaf. The
In contrast with wild-type plants, the first lateral leaflet primordium just before initiation of blade outgrowth in P4 stage each leaflet occurs acropetally and is first evident in the terminal (Coleman and Greyson, 1976; Dengler, 1984). Lobe formation in P3 or P4 stage, and subsequent leaflets develop basipetally.

Our analysis suggested that differential expression of LYRATE in leaf dissection will give rise to leaflets and blade in tomato compound leaves. The role of LYRATE in leaf dissection results in ectopic spongy mesophyll tissue replacing collenchyma cells just above the epidermis on the abaxial side of the leaf (Figures 2F and 2G, arrow). In addition, collenchyma cells are missing from the center of the adaxial side at the midrib region (Figures 2F and 2G, arrowhead). These results suggest that LYRATE is not a major factor patterning differentiation of internal tissues during leaf development but may play a minor role in specification of collenchyma identity in the midrib.

**LYR Expression Pattern**

To further analyze the role of LYR in leaf development in tomato, we characterized the LYR expression pattern. RT-PCR analysis revealed LYR expression in shoots (including P0-P5), young leaves (4 to 7 mm in length), inflorescences, and flowers but not in the mature leaf, stem, or hypocotyl (Figure 3A). In vegetative apices, the LYR transcript localized to initiating leaf primordia and expanding leaves but was excluded from the SAM (Figures 3B to 3F; see Supplemental Figure 7 online). LYR expression was evident across the adaxial surface of the leaf primordia with more intense expression in distal tissues that give rise to the blade of the terminal leaflet. During these early stages of leaf development, the LYR expression domain resembles that of JAG in Arabidopsis young leaf primordia (Figures 3B and 3C; see Supplemental Figure 7 online) (Ohno et al., 2004). As the leaf transitions to leaflet initiation, LYR is strongly and specifically upregulated in leaflet primordia (Figures 3D to 3F). Expression was low or not detected in tissues adjacent to leaflets. This represents a significant divergence from reported JAG expression. In addition to expression in developing leaves, LYR was also detected in all floral organ primordia (Figure 3G). In summary, the LYR expression domain is consistent with a role in promoting active cell proliferation in developing lateral organs. Interestingly, LYR is strongly expressed in the actively proliferating cells that will give rise to leaflets and blade in tomato compound leaves. This result suggests that LYR expression might regulate the differential outgrowth necessary for dissected leaf development.

**Ectopic Expression of LYR in Tomato Leaves**

We also examined the histology of LYR leaves to determine whether these defects extended to the organization of internal tissues. Paraffin sections through mature LYR blade tissue revealed relatively normal internal patterning, suggesting that LYR does not play a major role in patterning adaxial/abaxial polarity or differentiation (Figures 2F and 2G). Interestingly, histological analysis of the LYR terminal leaflet midrib revealed ectopic spongy mesophyll tissue replacing collenchyma cells just above the epidermis on the abaxial side of the leaf (Figures 2F and 2G, arrow). In addition, collenchyma cells are missing from the center of the adaxial side at the midrib region (Figures 2F and 2G, arrowhead). These results suggest that LYR is not a major factor patterning differentiation of internal tissues during leaf development but may play a minor role in specification of collenchyma identity in the midrib.

### Table 1. Phenotypic Characterization of lyrate Mature Leaves

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Leaf Length (cm)</th>
<th>Width of the First LL (cm)</th>
<th>Number of ILs between the TL and the First LL</th>
<th>TL Lobe Number of LL</th>
<th>Number of SLs on the First LL</th>
<th>Number of TLs on the First LL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>29.6 ± 0.55</td>
<td>15.9 ± 0.52</td>
<td>1.0 ± 0.31</td>
<td>6 ± 0</td>
<td>1.57 ± 0.81</td>
<td>0</td>
</tr>
<tr>
<td>lyr</td>
<td>24.9 ± 1.14a</td>
<td>19.0 ± 0.82</td>
<td>2.5 ± 0.50a</td>
<td>4.7 ± 0.76a</td>
<td>3.33 ± 0.33</td>
<td>0.83 ± 0.40</td>
</tr>
<tr>
<td>Wild type</td>
<td>29.7 ± 0.78</td>
<td>18.5 ± 0.97</td>
<td>3.7 ± 0.37</td>
<td>2.0 ± 0.37</td>
<td>6 ± 0</td>
<td>0.8 ± 0.29</td>
</tr>
<tr>
<td>lyr</td>
<td>22.5 ± 0.73a</td>
<td>22.0 ± 1.07</td>
<td>3.6 ± 0.37</td>
<td>3.5 ± 0.27a</td>
<td>2 ± 0</td>
<td>5.7 ± 0.54</td>
</tr>
<tr>
<td>Wild type</td>
<td>24.0 ± 0.71</td>
<td>18.5 ± 0.89</td>
<td>1.38 ± 0.64</td>
<td>1.5 ± 0.28</td>
<td>6 ± 0</td>
<td>1.75 ± 0.66</td>
</tr>
<tr>
<td>lyr</td>
<td>22.9 ± 0.52a</td>
<td>21.5 ± 1.09</td>
<td>4.30 ± 0.56a</td>
<td>5.3 ± 0.47</td>
<td>3 ± 0.38</td>
<td>8.60 ± 0.37</td>
</tr>
</tbody>
</table>

Measurements of leaf length were done from tip to petiole base. Measurements of leaf width were done from margin to margin of the first lateral leaflet pair. TL, terminal leaflet; LL, lateral leaflet; SL, secondary leaflet; TL, tertiary leaflet. Asterisk indicates that the difference between the wild type and lyr is statistically significant at P < 0.05.
in dissected leaves. To determine whether this might be the case, we attempted to alter the LYR expression domain by generating transgenic plants that ubiquitously express LYR (35S:LYR). Analysis of 16 plants expressing 35S:LYR revealed eight plants with ectopic blade growth on petioles, rachis, and petiolules (Figures 4B to 4D). In some cases, ectopic outgrowth was along the entire rachis (Figure 4B), while in weaker events, the ectopic blade only partially covered the main rachis and petiolules (Figures 4C to 4F). Leaflet placement was irregular compared with that of the wild type, suggesting ectopic primordia initiation (Figure 4E). Secondary leaflet formation was not in evidence or was dramatically reduced (Figure 4). Occasionally, the leaves of these plants were bifurcated (see Supplemental Figure 8 online). Flowers of transgenic plants were larger than wild-type flowers (see Supplemental Figure 8 online). These plants rarely produced fruits, and when they did, the fruits were usually irregular in shape and mostly parthenocarpic. The increased expression of LYR, as detected by RT-PCR and quantitative RT-PCR analyses, could be correlated to these phenotypes (Figures 4G and 4H). In contrast with JAG overexpression in Arabidopsis, and in agreement with what was shown for 35S:LYR in Arabidopsis (see Supplemental Figure 3 online), bract formation was not evident in tomato plants expressing 35S:LYR. Together, these results suggest that LYR promotes blade outgrowth in tomato leaves. Furthermore, ubiquitous expression of LYR throughout the developing leaf primordia results in highly simplified leaf development. This result further suggests that punctuated LYR expression is important to specify differential outgrowth in dissected leaves.

LYR Negatively Regulates KNOX Gene Expression

The knotted-like homeobox genes have been shown to positively regulate compound leaf development by promoting a meristematic environment in developing leaves (Hareven et al., 1996; Chen et al., 1997; Parnis et al., 1997; Janssen et al., 1998), and overexpression of several KNOX genes in tomato results in a highly dissected leaf form. We investigated the possibility that the lyr phenotype might result from alteration in expression of members of the KNOX transcription factor gene family. Using quantitative RT-PCR, we examined the expression of two KNOX genes, LeT6 (the STM ortholog, also known as TKN2) and TKN1 (the bp ortholog) in wild-type and mutant apices (Hareven et al., 1996; Chen et al., 1997). Both LeT6 and TKN1 expression levels were upregulated in lyr when compared with the wild type, although only LeT6 upregulation was statistically significant (Figure 5A). These results suggest that LYR negatively regulates KNOX gene expression in developing tomato leaves. Thus, KNOX overexpression may partially explain the increase in leaf complexity seen in lyr.

LYR Is Necessary for a Robust Auxin Response in Tomato Leaves

Auxin signaling has been shown to be important for leaflet initiation and for lamina outgrowth in tomato (DeMason and Chawla, 2004; Wang et al., 2005). Since lyr develops fewer leaflets and displays restricted lamina outgrowth, we examined whether the auxin response is altered in lyr. Using quantitative
RT-PCR, we found that the expression of the auxin-responsive genes PIN1, IAA9, and IAA4 (Wang et al., 2005) was significantly downregulated in lyr when compared with wild-type plants (Figure 5A). IAA3 was similarly downregulated, albeit not to statistically significant levels (Figure 5A). These results suggest that LYR positively regulates the auxin response.

To further address this hypothesis, we examined the activity of the auxin-responsive transgene pPIN1:PIN1:GFP in the lyr mutant background. In agreement with previously published results, pPIN1:PIN1:GFP expression in a wild-type background is highest in initiating leaf and leaflet primordia (Figure 5B) (Koenig et al., 2009). Green fluorescent protein (GFP) was also upregulated in initiating leaves and leaflets of lyr but at much reduced levels (Figures 5C and 5D). This reduction in PIN1:GFP expression supports the quantitative RT-PCR results showing endogenous tomato PIN1 is downregulated in lyr (Figure 5A).
somewhat intermediate phenotype relative to both single mutants (Figure 5H). The increased secondary dissection observed in lyr was eliminated in the e lyr doubles (Figure 5H), but ectopic blade outgrowth in e lyr was restricted distally when compared with e. This intermediate phenotype suggests that E and LYR regulate the auxin response independently.

**DISCUSSION**

We have described the identification of the classical tomato mutant lyr, which results from a mutation in the tomato ortholog of JAG. Like JAG, LYR acts to promote lateral outgrowth in aerial organs. LYR is expressed in a punctuated pattern during leaf development that coincides with the location of auxin response maxima and subsequent leaflet initiation. LYR promotes auxin response in the developing leaflet, first to drive primordia initiation, and later to promote blade outgrowth.

**The Function of LYR in Patterning Dissected Leaves**

During early development, the simple leaves of Arabidopsis are divided into two major domains that are delineated by JAG expression (Ohno et al., 2004). The distal region of the leaf primordium that will give rise to the blade is characterized by high expression of JAG, while the proximal bladeless petiole lacks JAG expression. In compound leaves, the patterning of outgrowth is significantly more complex, and our results reveal LYR as an important molecular component of this process.

A major difference between early compound and simple leaf development is the initiation of leaflet primordia. Simple leaves initiate outgrowth throughout the distal margin to promote development of a continuous blade. By contrast, initiation and delineation of leaflets in compound leaves requires unequal promotion of outgrowth along the margin. In accordance with this requirement, we find that LYR/JAG expression has diverged from the bipartite pattern that delineates blade and petiole in simple leaves to a punctuated pattern accommodating localized leaflet outgrowth. Additionally, the lyr mutant suppresses leaflet initiation along the primary axis of development. Loss of leaflet initiation is most evident in the lyr2 allele that results from a mutation of the LYR DNA binding domain. It is possible that the more severe phenotype of this allele results from a dominant-negative effect, while lyr1 and lyr3 are more representative of LYR loss of function. This hypothesis remains to be tested directly in future experiments. Regardless, all alleles reduced primary leaflet number to some extent, demonstrating that LYR is an important regulator of leaflet initiation.

The reduction in the number of leaflets in lyr plants seems to result from loss of one or two basal lateral pairs, suggesting that the distal pair may be less sensitive to LYR loss of function. This may result from the activity of a redundant factor in the distal leaf primordium, and in the absence of a tomato genome sequence, we cannot exclude the possibility of the existence of a NUB ortholog. It is worth noting that a common compound leaf morphology is trifoliate (as seen in several legumes), suggesting that production of a single pair of lateral leaflets may be fundamentally different from reiteration of this process to form additional lateral pairs. This hypothesis is supported by the tomato combination, these results suggest that LYR is necessary to condition high levels of auxin response in developing compound leaves.

We additionally examined the role of LYR in promoting the auxin response in leaflets by generating double mutants with the auxin response mutant *entire* (e) (Figures 5E to 5H). e mutants fail to inhibit the auxin response between developing leaflets, resulting in ectopic blade outgrowth along the rachis (Figure 5G) (Koenig et al., 2009). e lyr double mutants exhibited a
monogenic mutant \( tf \), which produces trifoliate leaves that are otherwise similar to those of wild-type plants. Identification of the molecular nature of the \( tf \) mutant and examination of the genetic interaction between \( lyr \) and \( tf \) should add to our understanding of the process that regulates initiation of the first pair of leaflets.

The role of \( LYR \) in tomato compound leaf development bears a striking resemblance to that seen for auxin (Koenig et al., 2009). Both \( LYR \) and auxin accumulate in initiating leaflets and are absent in the bladeless tissues separating leaflets. Furthermore, exogenous application of auxin to leaf primordia results in ectopic blade production on the rachis, reminiscent of the \( LYR \) overexpression phenotype. This phenotype is also mimicked by ectopic activation of auxin response in the simple leafed tomato mutant \( e \). Our results show that these similarities are not a coincidence; \( LYR \) promotes a strong auxin response, suggesting that the \( lyr \) phenotype might result from a reduction in the magnitude of transcriptional response to auxin. The intermediate phenotype of \( e lyr \) suggests that the two genes regulate auxin response independently. An additional patterning factor, \( GOB \), has also been shown to act in at least partial independence of \( E \). Thus, it is difficult at this time to establish a linear relationship between the initial auxin signal and the downstream patterning elements required for leaf dissection in tomato. Although auxin distribution can influence \( E \) activity directly via degradation (Koenig et al., 2009), the relationship between \( GOB \) and \( LYR \) expression and auxin distribution remains unclear. Future experiments testing the phenotypic outcome of auxin treatment in \( lyr \) and the effect of auxin treatment on \( LYR \) and \( GOB \) transcript levels will help resolve this issue.

Function of \( LYR \) in Leaflet Indeterminacy

\( KNOX \) genes are key regulators in maintaining indeterminacy in the SAM and in the developing compound leaf (Bharathathan et al., 2002). Increases in both leaflet number and the order of leaf complexity are seen in tomato mutants or transgenics that overexpress \( KNOX \) genes (Hareven et al., 1996; Chen et al., 1997). The \( lyr \) loss-of-function mutants also showed an increase in leaflet number, and expression analysis showed a modest increase in \( KNOX \) transcript levels (Le76 and \( TKV1 \)) in \( lyr \) (Figure 5). It is possible that \( lyr \) produces additional leaflets that are due to this increase in \( KNOX \) expression. Furthermore, both \( KNOX \) overexpression and \( lyr \) loss of function result in delayed trichome differentiation during leaf development. In the absence of loss-of-function mutations in tomato \( KNOX \) genes, it is difficult to definitively establish their contribution to the \( lyr \) phenotype. However, it is clear that even moderate \( KNOX \) overexpression in tomato transgenics and naturally occurring mutants results in highly ramified leaves along all leaf axes (Janssen et al., 1998). It is therefore likely that upregulation of \( KNOX \) gene expression in \( lyr \) contributes in part to the production of additional leaflets.

Secondary Morphogenesis and the Relationship between Simple and Compound Leaves

The homology between compound and simple leaves has long been a subject of botanical debate. The resemblance of the leaflets of compound leaves to an entire simple leaf has led some classical botanists to conclude that compound leaves possess a shoot-like identity (Arber, 1950). An alternative hypothesis suggests that compound leaves are simply the result of dissection of the simple leaf blade into smaller units (Kaplan, 1975). Although such mutually exclusive arguments are unlikely to accurately represent biological reality, they serve as useful starting points to interpret comparative developmental analyses.

Compound leaf morphogenesis is characterized by two basic developmental processes; the elaboration of leaflet primordia during early leaf development and restriction of blade outgrowth during late leaf development that maintains the independence of these leaflets. The \( KNOX \) genes are known to be important for the former process, and \( KNOX \) gene expression in young leaf primordia is closely correlated with initiation of leaflet primordia across most of flowering plants (however, see Champagne et al. [2007] for an exception). Since \( KNOX \) genes are important regulators of meristem identity, it is generally thought that this result provides evidence for homology between shoots and compound leaf primordia. Although the expression of \( KNOX \) genes in leaves and the subsequent formation of leaflet primordia may be prerequisites for compound leaf development in most species, it is clear that they are not sufficient. Many species show both of these characteristics, but still produce simple leaves by initiating blade outgrowth throughout the leaf margin, resulting in a continuous lamina (Bharathathan et al., 2002). This process, by which blade outgrowth contributes to leaf shape, is termed secondary morphogenesis.

The mere existence of leaf simplification during secondary morphogenesis suggests that compound leaves are generated in part by dissecting the simple leaf blade, but several other lines of evidence support this idea. Specification of a complex pattern of growth such as that found during dissected leaf development requires the asymmetric distribution of molecular or physiological signals. Auxin appears to be one such signal, as its distribution during compound leaf development is in part responsible for delineating blade outgrowth and leaflet initiation (Barkoulas et al., 2008; Koenig et al., 2009). Our results demonstrate a close association between the function of \( LYR \) and auxin during compound leaf development. Much like the proposed function of auxin, \( LYR \) promotes outgrowth of both leaflets and blade, and its ubiquitous expression results in the formation of ectopic blade on petiolules and rachis. When considering the homology of compound and simple leaves, it is important to note that auxin peaks also form in the margins of simple leaves (guiding vascular and serration development) (Hay et al., 2006; Scarpella et al., 2006), while no such peaks have been reported for \( JAG \) or \( NUB \) expression. Thus, changes of \( JAG/LYR \) expression pattern represent a clear difference between simple and compound leaf development. It is therefore possible that alteration in auxin distribution, response pathways, or \( LYR \) expression pattern might be responsible for the natural occurrence of leaf simplification through secondary morphogenesis. Interestingly, both currently known promotive factors (auxin and \( LYR \)) in this network regulate both leaflet initiation and blade outgrowth. This suggests that all outgrowths from the marginal blastozone have some homology at the genetic and molecular levels and calls into question the idea that leaflet outgrowth is inherently different from blade outgrowth. It may be that the difference between
these two fates is determined purely by the localization and/or intensity of the promotive signal during initiation. Previous experiments with auxin have suggested that this is the case (Koenig et al., 2009), and the fact that Lyr overexpression can result in ectopic blade outgrowth and additional leaflet initiation also supports this hypothesis. Combined, these results reveal an intricate network of molecular and physiological regulators of growth that determine leaf shape in tomato, and it will be interesting to determine how these factors may influence the patterning of the myriad of leaf morphologies yet to be explored.

**METHODS**

**Plant Material and Growth Conditions**

*Solanum lycopersicum* cv VF36 was used for all experiments described in this article. The tomato mutants l4r (L/A0763), lyr (L/A0792), and tf (L/A0579) were kindly provided by Tim Willis and Roger Chetelat from the C.M. Rick Tomato Genetics Resource Center (htp://tgrc.ucdavis.edu/). The tomato mutant lyr3 was kindly provided by Yuval Eshed (Weizmann Institute of Science, Israel) and the Genes that Make Tomatoes mutant database (htp://zamir.sgn.cornell.edu/mutants/).

The tomato transgenic line expressing pPIN1:PIN1:GFP was kindly provided by Cris Kuhlemeier (University of Bern). All tomato plants were grown in a growth chamber at 22°C with 75% relative humidity and a daylength of 16 h.

*Arabidopsis thaliana* plants were grown in a growth chamber at 20°C and a daylength of 18 h. All plants were in the Landsberg erecta background. Seeds of jag-2 and jag-3 (Ohno et al., 2004) were kindly provided by Carolyn Ohno and Elliot Meyerowitz (California Institute of Technology).

**Cloning S. lycopersicum JAG**

The *Arabidopsis* JAG sequence was used to search the SOL Genomics Network (htp://www.sgn.cornell.edu) identifying a single EST sequence using the Vector NTI Advance 9 Suite (Invitrogen). The tomato mutant LYRATE genomic DNA was extracted from mature leaf tissue using the CTAB protocol. The LR reaction was performed between the entry vector and the destination vector pTKO2 as described in the Gateway cloning technology instruction manual. The pTKO2:LYRATE plasmid was introduced into A. tumefaciens ASEX.

**3SS:LYR**

The cDNA fragment of LYR was amplified from a cDNA library, which was made from SAM, by PCR using primers LYRATE_F, 5’-ATCATCAAAAGTCGTCGTCACCAGCTTCCGCGTAATGAG-3’, and LYRATE_R, 5’-TGGAAGAGGTAACCTCTGTAATGAG-3’. The resulting cDNA was cloned into the pCR8/GW/TOPO entry vector (Gateway; Invitrogen) following the manufacturer’s protocol. The LR reaction was performed between the entry vector and the destination vector pTKO2a as described in the Gateway cloning technology instruction manual. The pTKO2:LYRATE plasmid was introduced into A. tumefaciens ASEX.

**Transgenic Plants**

3SS:LYR and PAlJAG:glLYRATE constructs were transformed into jag-2 and jag-3 (Landsberg erecta) plants by the floral dipping method (Clough and Bent, 1998). 3SS:LYR and LYR RNAi were transformed into tomato (S. lycopersicon cv VF36) at the Ralph M. Parsons Foundation Plant Transformation Facility (University of California, Davis).

**Mapping S. lycopersicum JAG**

JAG was mapped using a cleaved amplified polymorphic sequence marker in intron 2, using an F2 mapping population of tomato (*S. lycopersicon × Solanum pennellii*) that was provided by S. Tanksley (Cornell University, Ithaca, NY).

**RT-PCR Analysis**

Total RNA (1 μg) was extracted from tomato tissue using the Plant RNeasy mini kit (Qiagen) followed by RQ1 DNase (Promega) treatment and reverse transcription using random primer hexamers of the SuperScript III first-strand synthesis system for RT-PCR (Invitrogen). In order to obtain results within the linear range, PCR conditions were 94°C for 20 s, 53°C for 5 min followed by 30 cycles of 94°C for 20 s, 53°C for 20 s, and 72°C for 45 s using primers LYRATE RTPCR_F (5’-GAGTCGTCGTCACAGAGCGACTCCTGGAAGAGGTAATGAG-3’) and LYRATE RTPCR_R (5’-CCTCCCGCAGTTCCGCGTAATGAG-3’) for LYR; tomato actin primers (5’-GCTTCGCGTCGCGTCAMAAGAGCGACTCCTGGAAGAGGTAATGAG-3’) and (5’-GCGACTCCTGGAAGAGGTAATGAG-3’) were used as internal control for all RT-PCR reactions.
Quantitative Real-Time PCR Analysis

Quantitative real-time PCR analysis was performed using IQ SYBR Green Supermix (Bio-Rad) with a Bio-Rad iCycler iQ. cDNA was made as described above and was diluted 20 times before being subjected to quantitative RT-PCR analysis. Two independent biological experiments were used for each sample, while quantifications were performed in triplicates. Specific gene expression was normalized to the internal control gene glyceraldehyde 3-phosphate dehydrogenase (GAPDH), and a negative control without cDNA was also performed for each primer set. The gene expression value of the wild type was used as a control and set at 1.0. For primer sequences, see Supplemental Table 1 online.

Scanning Electron Microscopy

Tomato apices were fixed as described by Bharathan et al. (2002). Electronic images were obtained with a Hitachi S-3500 N scanning electron microscope (Hitachi Science Systems).

Histology

Tissues were fixed in FAA (formaldehyde 1.85%, glacial acetic acid 5%, and ethanol 63%), dehydrated through an ethanol series (70, 80, 90, and 100%, 30 min each), embedded in paraffin, sectioned using a Microtome HM 340 E (MICROM International), and stained with 0.1% toluidine blue O. The sectioned material was observed using a NIKON Eclipse E600 microscope, and digital images were taken using a SPOT RT camera (Diagnostic Instruments).

In Situ Hybridization Analysis

In situ analysis was performed as previously described (Long et al., 1996). A 560-bp fragment that was amplified from In situ analysis was performed as previously described (Long et al., 1996). (Diagnostic Instruments).

Tomato apices were fixed using a NIKON Eclipse E600 microscope, and digital images were taken using a SPOT RT camera (Diagnostic Instruments).

Leaf Mesophyll Cell Density Measurement

Leaf 1 was collected from wild-type and lyr plants with three expanded leaves. Photographs of all leaves were taken using an Olympus E600 camera. Measurement of leaf area was performed using the Image J software package. Terminal leaflet blade tissue was cleared by boiling in 80% ethanol until most chlorophyll was removed. The tissue was then transferred to 5% sodium hydroxide for 12 h, followed by chloral hydrate treatment overnight. The tissue was mounted and examined under a Nikon Eclipse SP-500UZ microscope. Each tissue sample was randomly photographed three times for technical replication from which a mean cell density was calculated for each biological replicate. Biological variation was then used to compare the genotypes.

Double Mutant Construction

The e homozygous plants were used as males and crossed to lyr heterozygotes to generate F1 plants. Plants were genotyped for lyr to identify heterozygotes, which were then selfed to generate segregating progeny. The novel e lyr double phenotype could be identified in these F2 plants but was confirmed by PCR for lyr and direct sequencing for e.

Accession Numbers

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number EU490614 for LYR and AAR30036 for Arabidopsis JAG.

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure 1. Protein Alignment of JAG and SI JAG.
Supplemental Figure 2. DNA Gel Blot Analysis of SI JAG in Tomato.
Supplemental Figure 3. Function of SI JAG in Arabidopsis Transgenic Plants.
Supplemental Figure 4. DNA Gel Blot Analysis Revealing Altered SI JAG gene in lyr and lyr Tomato Mutants.
Supplemental Figure 5. lyr Flower Phenotype.
Supplemental Figure 6. lyr Leaf Mesophyll Cell Density.
Supplemental Figure 7. Lyr Expression Pattern in an Apex of Wild-Type Tomato.
Supplemental Figure 8. Additional 3SS:LYR Phenotypes.
Supplemental Table 1. Primers Used in This Study.

Supplemental References.

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