The Aquilegia JAGGED homolog promotes proliferation of adaxial cell types in both leaves and stems

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Summary

- In order to explore the functional conservation of JAGGED, a key gene involved in the sculpting of lateral organs in several model species, we identified its ortholog AqJAG in the lower eudicot species Aquilegia coerulea.
- We analyzed the expression patterns of AqJAG in various tissues and developmental stages, and used RNAi-based methods to generate knockdown phenotypes of AqJAG.
- AqJAG was strongly expressed in shoot apices, floral meristems, lateral root primordia and all lateral organ primordia. Silencing of AqJAG revealed a wide range of defects in the developing stems, leaves and flowers; strongest phenotypes include severe reduction of leaflet laminae due to a decrease in cell size and number, change of adaxial cell identity, outgrowth of laminar-like tissue on the inflorescence stem, and early arrest of floral meristems and floral organ primordia.
- Our results indicate that AqJAG plays a critical role in controlling primordia initiation and distal growth of floral organs, and laminar development of leaflets. Most strikingly, we demonstrated that AqJAG disproportionally controls the behavior of cells with adaxial identity in vegetative tissues, providing evidence of how cell proliferation is controlled in an identity-specific manner.

Introduction

The seemingly infinite variety of forms of leaves and floral organs is achieved by a complex interplay of cell proliferation and expansion. These processes are controlled by a number of fine-tuned programs that are executed with temporal and spatial precision, making decisions about the rate, orientation and duration of cell division and differentiation. Throughout the process of morphogenesis, many of these programs are regulated based on the adaxial–abaxial axis of the lateral organ. This axis is established soon after the organ primordium becomes distinguishable from the meristem, as cells adjacent to or away from the meristem acquire adaxial and abaxial identity, respectively. The genetic programs determining these polarities are strikingly conserved across plant taxa, in which genes including the class III homeodomain-leucine zipper transcription factors control adaxial identity, whereas genes such as the KANADI (KAN) and ETTIN/ARF3 control abaxial identity (Emery et al., 2003; Harrison et al., 2005; Pekker et al., 2005).

It is widely accepted that laminar expansion in lateral organs is induced by the boundary where the adaxial and abaxial domains meet (Waites & Hudson, 1995). Therefore, modification of this boundary or differential development of adaxial and abaxial tissues is one mechanism to generate diversity in organ form. For instance, modifications of organ polarity have been implicated in a range of morphologies, from the pitchfork leaves of Sarracenia purpurea (Fukushima et al., 2015) to the narrow filaments and broad anthers of rice stamens (Toriba et al., 2010). Nonetheless, even for a simple flat leaf, the developmental patterns of the adaxial and abaxial domains are by no means identical: spongy mesophyll cells stop dividing before the overlying palisade mesophyll cells, and are then separated during the active proliferation of the latter, resulting in loosely packed spongy cells with air space on the abaxial side, and densely packed palisade cells on the adaxial side (Dale, 1988).

Although the precise mechanism of how cell proliferation and expansion are controlled in response to abaxial vs adaxial identities remains elusive, research on model species such as Arabidopsis (Arabidopsis thaliana) and rice (Oryza sativa) have discovered a number of genes that generally promote or restrict organ growth, providing a useful starting point to explore this question. For example, members of the YABBY and WUSCHEL-RELATED HOMEOBOX family are downstream of polarity determinants and are thought to integrate polarity signals to promote lamina outgrowth (Siegfried et al., 1999; Eshed et al., 2004; Vandenbussche et al., 2009; Sarojam et al., 2010; Nakata et al., 2012). Genes such as the BLADE ON PETIOLE (BOP) family have been found to shape the basal portion of lateral organs (e.g. leaf petiole), preventing outgrowth of laminae by interacting with polarity genes (Ha et al., 2007; Jun et al., 2010). As a counterpart to the BOPs, JAGGED (JAG) promotes growth of the distal portion of the lateral organ, maintaining active cell proliferation and expansion until the organ reaches the final size (Dinneny et al., 2004; Ohno et al., 2004).

In this study, we identify the JAG homolog (AgJAG) in the lower eudicot species Aquilegia coerulea and investigate its functions during A. coerulea morphogenesis. JAG encodes a...
C2H2-type zinc-finger transcriptional factor that is broadly expressed in all lateral organs in Arabidopsis. Its null mutant displays reduced development in the distal regions of lateral organs, whereas overexpression of JAG leads to ectopic blade tissue growth and formation of bracts (Dinneny et al., 2004; Ohno et al., 2004). NUBBIN (NUB), a close paralog of JAG, has been identified only in Arabidopsis (Dinneny et al., 2006), and, unlike JAG, is expressed strictly in the adaxial domain of lateral organs. Single nub mutants display no apparent defect, whereas jag nub double mutants show a more severe version of the jag single mutant, and thus JAG and NUB are thought to work redundantly to promote proliferation in the distal regions of lateral organs. Complete loss-of-function of mutants in the JAG homologs LYRATE (LYR) and OPEN BEAK (OPB) also have been characterized in tomato (Solanum lycopersicum) and rice, respectively (David-Schwartz et al., 2009; Horigome et al., 2009). These mutants are similar to jag nub in that they show impaired laminar growth, but also differ in several aspects, including the formation of fewer lateral leaflets in lyr compound leaves, and reduced expression levels of B class floral identity genes in opb flowers.

Our particular interest in Aquilegia JAG homolog function stems from our ongoing analysis of the developmental programs controlling several novel floral organ types. Aquilegia flowers possess elongated nectar spurs on their petals, which previous studies controlling several novel floral organ types, stems from our ongoing analysis of the developmental programs controlling several novel floral organ types controlling several novel floral organ types.

Materials and Methods
Identification of the candidate gene
The complete DNA sequence of AqJAG was obtained via BLAST (Basic Local Alignment Search Tool; Altschul et al., 1990) using the cDNA sequence of JAGGED (JAG) from Arabidopsis against the Aquilegia coerulea (James) genome on PHYTOZOME (http://www.phytozome.net/). A reverse BLAST search of the translated amino acid sequence of the putative AqJAG against an Arabidopsis protein database (www.arabidopsis.com) also identified JAG with the highest score. To confirm the orthology of AqJAG (Aquilegia genome v3.1 locus Aqcoe3G037600, GenBank accession KX879697), a 56-locus alignment containing putative JAG homologs from a variety of land plant taxa was constructed via BLAST searches in GenBank, PHYTOZOME and the DFCI Plant Gene Index (http://compbio.dfci.harvard.edu/cgi-bin/pf/plant.html). Information regarding 14 other related zinc-finger proteins in Arabidopsis was obtained from previous family analyses (Engblom et al., 2004; Ciftci-Yilmaz & Mittler, 2008). All amino acid sequences were aligned using CLUSTAL W (Larkin et al., 2007) and then adjusted by hand using MacVECTOR (Cary, NC, USA). The phylogenetic analysis was performed using the CIPRES web portal (http://www.phylo.org/portal2/login?inputaction). A maximum-likelihood tree was constructed using randomized axelerated maximum likelihood (RAxML; Stamatakis et al., 2008) implemented with the default Jones–Taylor–Thornton (JTT; Jones et al., 1992) model of amino acid substitution. The resultant tree was displayed by software program FIGTREE v.1.3.1 (http://tree.bio.ed.ac.uk/software/figtree/).

In situ hybridization
A 465-bp DNA fragment in a nonconserved region of AqJAG was amplified (forward primer: 5’-TCTACACACCCTATCT TCTCG-3; reverse primer: 5’-TGTCAGTTGTGTTCTTCT ATCTG-3) and cloned into the pCR™4-TOPO® vector. Both sense and anti-sense probes were alkaline-hydrolyzed at 60°C for 41 min to an average length of 150 bp. All in situ hybridization steps were performed as described by Kramer (2005). Results were visualized in the Harvard Center for Biological Imaging on a Zeiss AxioScan microscope using trans-illumination with white light.

Virus-induced gene silencing
Virus-induced gene silencing (VIGS) was performed according to protocols described previously (Gould & Kramer, 2007; Sharma & Kramer, 2013a). The TRV2 vector containing a fragment of A. coerulea ANTHOCYANIDIN SYNTHASE (AqANS-TRV2) as a marker of the target gene silencing has been previously prepared and described (Gould & Kramer, 2007). The same 465-bp AqJAG fragment amplified for in situ hybridization was cloned into the AqANS-TRV2 plasmid to generate the AqJAG-AqANS-TRV2 construct, which was subsequently transformed into GV3101 electrocompetent Agrobacterium cells. Three hundred plants were infiltrated with AqJAG-AqANS-TRV2 construct and 100 plants were treated with AqANS-TRV2 construct as controls. Plants showing knockdown phenotypes were documented and photographed using a Canon X type digital SLR camera (Canon, Melville, NY, USA).

Scanning electron microscopy and histology
Organs of interest were fixed in FAA (50% ethanol, 4% formalin and 5% glacial acetic acid) and stored at 4°C. For scanning electron microscopy (SEM), samples were dehydrated through a graded ethanol series to 100% EtOH at 4°C and critical point-dried before imaging. Materials were imaged using a JEOL
JSM-6010 LC Scanning Electron Microscope. Samples for histology were dehydrated through the same graded ethanol series, stored in 100% Citrasolv, and embedded in Paraplast Plus (Oxford Labware, St Louis, MO, USA). Tissues were sectioned to 8 µm using a rotary microtome and stained with 0.01% Safranin O and 50% Fast Green according to the protocol described in Ruzin (1999). The wild-type (WT) and AqJAG-silenced stems were collected when the terminal flowers were opened, and sections were processed in parallel. Slides were photo-documented in the Harvard Center for Biological Imaging on a Zeiss AxioScan microscope using trans-illumination with white light.

Quantitative real-time polymerase chain reaction

Total RNA was extracted from flower buds from AqJAG-AqANS-TRV2- and AqANS-TRV2-treated plants using PureLink Plant RNA Reagent (Invitrogen), and treated with Turbo DNase (Ambion). cDNA was synthesized from 1 µg of RNA using Superscript II reverse transcriptase (Invitrogen) and oligo (dT) primers. The resulting cDNA was diluted 1 : 10 as template. Brilliant II SYBR Green QPCR Master Mix, Low ROX (Stratagene, Cedar Creek, TX, USA/La Jolla, CA, USA), was used to carry out the quantitative real-time polymerase chain reactions (qRT-PCR) in the Stratagene Mx3000P QPCR System. QRT primers for AqJAG were designed to span the intron positions (forward primer: 5'-GGTGGAAAAGATGAGTGTGGG-3'; reverse primer: 5'-AAATGAGACCCCTTGTGTC-3'). AqIPP2 (15PENTYL PYROPHOSPHATE:DIMETHYLALLYL PYROPHOSPHATE ISOMERASE2) was used for normalization as it has been shown previously to have little quantitative transcriptional variation across tissues and developmental time points (Sharma et al., 2011). Primer efficiencies were evaluated before the expression analysis using six dilution series. Eight biological replicates for both control and AqJAG VIGS group were examined. Expression for each biological sample was assayed from three replicates per reaction plate. The variability resulting from technical replicates was negligible compared with the variability from biological replicates, so only biological variability is presented here. Relative gene expression levels were calculated using the 2^(-ΔΔCT) method described in Livak & Schmittgen (2001), and formulas were adjusted with the obtained primer efficiencies for both AqJAG and AqIPP2. An unpaired Student’s t-test was used to determine the statistical significance of differences between experimental and control values.

Results

Identification of AqJAG

The putative JAG homolog in A. coerulea genome is a 2690-bp-long single copy gene that is predicted to encode 423 amino acid residues. Consistent with the structure of JAG in Arabidopsis, AqJAG possesses characteristic features of a C2H2 type zinc-finger protein, specifically a single zinc-finger domain, as well as a conserved QALGGH sequence, a proline-rich motif positioned towards the C-terminal and an ERF-associated amphiphilic repression (EAR) motif (Supporting Information Fig. S1). In order to determine the correct orthology of AqJAG, we constructed a maximum-likelihood tree using the amino acid sequences from 42 putative JAG homologs from various land plant taxa, as well as 14 C2H2 zinc-finger proteins with one or two zinc-finger domain(s) from Arabidopsis (Fig. S2). The phylogeny was rooted by using sequences with two zinc-finger domains as an outgroup. This produced a tree in which all putative JAG homologs, including AqJAG, cluster into a large clade with strong bootstrap support (95%), whereas the other seven proteins with a single zinc-finger domain from Arabidopsis resolved into a separate clade. Within the JAG clade, many branches did not show strong support due to the combined facts that the conserved domains are very highly conserved, and therefore have little phylogenetic information, whereas other regions of the genes show relatively low conservation and are therefore too variable to resolve deeper nodes. Nevertheless, although the phylogenetic position of AqJAG within the JAG homolog clade cannot be fully resolved with high confidence, our analysis has sufficient power to show that we have identified the correct JAG ortholog in A. coerulea.

AqJAG is strongly expressed in meristems, lateral organ primordia and regions undergoing rapid growth

A detailed picture of the specific spatial and temporal patterns of AqJAG expression in different tissues and developmental stages in A. coerulea was obtained by in situ hybridization (developmental stages of Aquilegia flowers in Table S1). Strong signal of AqJAG was detected in shoot apical meristems (Fig. 1a), leaf primordia (Fig. 1a), lateral root primordia (Fig. 1b), floral meristems (Fig. 1d–i) and all initiating floral organ primordia (Fig. 1c–i). During stage 7 of flower development, when the stamens became stalked at the base and the anthers started to differentiate, hybridization signal in stamens declined in the filament and became concentrated in the anthers (Fig. 1h). Meanwhile, the staminodia became distinguishable from stamens, with higher concentration of AqJAG expression in their distal tips (Fig. 1g,h). Signal disappeared when staminodia began to flatten in stage 9 (Fig. 1i). AqJAG was highly expressed in the distal region of the carpels from their inception, but moderate expression could also be seen throughout the carpels during all developmental stages (Fig. 1g,h). Subsequently, signals were detected in the initiating ovules in the carpels (Fig. 1i). When the flower transitions into stage 10 and onward, petal primordia continue differentiating and initiate petal spurs, and strong expression can be detected throughout the petals (Fig. 1i; later stage data not shown). No expression was observed with sense probe (Fig. 1c).

VIGS silencing of AqJAG

In order to knock-down the expression of AqJAG, we cloned a 465-bp fragment from a nonconserved region of AqJAG together with a fragment of gene AqANS into the multiple cloning site of TRV2 vector. AqANS has been used previously as an indicator of gene silencing from VIGS in A. coerulea flowers, and it has been demonstrated that plants infiltrated with AqANS-TRV2 show no
morphological change from the WT aside from a loss of anthocyanin in vegetative and floral tissues (Sharma & Kramer, 2013a). The AqANS-TRV2 construct was used as VIGS control.

In order to assess the downregulation of AqJAG in the VIGS-treated plants, we subsequently performed qRT-PCR (Fig. S3). We collected young axillary buds from inflorescences showing strongly silenced phenotypes (see below). After normalizing the expression levels with the housekeeping gene AqIPP2, the relative expression levels of AqJAG were significantly lower in the AqJAG VIGS samples compared with those of control plants (Student’s t-test, \( P < 0.005 \)), indicating that the observed phenotypes were indeed due to the silencing of AqJAG expression.

AqJAG-silencing results in severe lamina reduction and concave curvature in leaves

A typical WT A. coerulea rosette leaf consists of three parts: a large pair of stipules at the leaf base, a long petiole and a ternate compound lamina in which each leaflet is usually deeply lobed or further segmented (Fig. 2a; Kramer & Hodges, 2010; Pabón-Mora et al., 2013). The petiolules are not as elongated as the petioles and higher order petiolules with leaflets are usually only observed in leaves produced after vernalization (Fig. 2a).

Among our AqJAG-treated plants, there was no change in the overall number of leaflets or segments per leaflet, but c. 31% exhibited a decrease in leaflet area, usually accompanied by reduction in petiolule length (Fig. 2b–h). The degree of laminar reduction varied greatly, reflecting the variation typically seen with VIGS-based silencing. Some leaflets only had small patches of affected lamina, whereas the remaining parts of the leaflet underwent normal cell proliferation and expansion, leading to asymmetric morphologies of the leaflets (e.g. leaflet segments in Fig. 2b–d). Normal leaf margin formation was often disrupted in these leaflets due to the impaired coordination of growth between the silenced and unsilenced regions (Fig. 2b–d). A number of plants also exhibited severe reduction in the entire lamina, resulting in significantly smaller leaflets (Fig. 2b–h). In the strongest silencing phenotypes, the lamina was so reduced that it became almost indistinguishable from the petiolule, appearing as an extremely stunted structure (Fig. 2d,h).

In addition, when looking at the leaves from the adaxial side, all leaflets experiencing laminar reduction were concave, at least in part (Fig. 2b–g), with an apparent association between the degree of growth repression and the severity of curvature (Fig. 2d–h). In some case, the entire lamina was concave, forming a bowl-like structure (Fig. 2c–g).

AqJAG-silenced leaves show a decrease in cell number and a change in adaxial cell identity

We analyzed the strongest phenotype of AqJAG-silenced leaves using SEM, and determined that the stunted structures are, in fact, differentiated ternate leaflet primordia in which the leaflet laminae failed to develop and expand (Fig. 3a). Disruption in the

![Fig. 1 Expression patterns of AqJAG in Aquilegia coerulea (a) vegetative meristems and leaf primordia, (b) lateral root primordia, and (d–i) early stages of flower development. (c) Hybridization with sense probe. (d) Stage 2 floral meristem. (e) Stage 3 floral meristem; only sepal primordia initiated. (f) Stage 6 floral meristem; all organ primordia except carpels initiated. (g) Stage 7 young flower bud with carpels just initiated. (h) Stage 9 young flower bud. (i) Stage 10 flower bud with flattened staminodia. lp, leaf primordia; sam, shoot apical meristem; asterisk, sepal primordia; arrow, petal primordia; arrowhead, staminodium. Bars, 100 µm.](image-url)
development of the laminar margin and extreme curvature of the laminae were both prominent. For leaves exhibiting this severe laminar reduction, no WT marginal cells, trichomes, or vascular strands were observed.

Furthermore, we compared the epidermal cell morphologies between WT/control and *AqJAG*-AqANS-TRV2-treated leaves. No difference in abaxial cell morphology was observed; the abaxial surface in all analyzed leaves from all treatment groups displayed irregular convex cells with stomata scattered across the whole surface (Fig. 3b,d). On the adaxial surface of *AqJAG*-silenced leaves, however, the cobblestone-shaped epidermal cells of WT leaves (Fig. 3c) were replaced entirely by small conical cells (Fig. 3e). These cells do not especially resemble the cell types found in other lateral organs, including floral organs. Given that no change in cell size or shape was found on the abaxial surface, we can infer that laminar reduction is associated primarily with a decrease in cell numbers in that domain. In the adaxial domain, the conical cells in *AqJAG*-silenced leaves are significantly smaller than the WT epidermal cells (up to 80% smaller), but this size difference is not entirely sufficient to explain the degree of lamina reduction, which can be as great as 97%. Based on these observations, we conclude that the reduction in laminar size in *AqJAG*-silenced leaves not only involves reduction in cell number in both the adaxial and abaxial domains, but also includes changes in cell size in the adaxial domain, apparently due to improper differentiation of cell identity.

*AqJAG*-silenced inflorescences display a number of novel phenotypes

Upon transition to the reproductive phase, *Aquilegia* flowers usually exhibit strong apical dominance and only a primary inflorescence will develop. Among all the *AqJAG*-AqANS-TRV2-treated plants, 43.2% showed more than one inflorescence, whereas only 8% of the VIGS control plants had secondary inflorescences (Fig. 4c), a significant difference (Pearson’s $\chi^2$ test, $P<0.0001$). Moreover, the first internodes of the WT inflorescence axes of *Aquilegia* often extend to the same level as the rosette leaves,
meaning that flowers always bloom above the rosette leaves (Fig. 4b). However, flowers from AqJAG VIGS plants were often observed opening below or at the level of rosette leaves (Fig. 4c). We thus measured the length of the first internode of the inflorescences in all of the VIGS plants after the terminal flowers had opened, which revealed a significant decrease in internode length among the AqJAG VIGS plants (Fig. 4a).

In addition to these shortened inflorescences, we observed peculiar outgrowths on the inflorescence stem in c. 9% of AqJAG VIGS plants (Fig. 4d–f). The outgrowths resemble thin laminae with little pigmentation. In most cases, the outgrowth occurred along just one side of the stem (Fig. 4e,f), but rarely there were growths on both sides (Fig. 4d). These outgrowths were observed only on the first internodes of secondary inflorescence axes.

In order to further characterize the structural changes that led to the lamina-like outgrowths, we investigated the stem anatomy using histology. The WT inflorescence of A. coerulea possesses the features of a typical dicot stem lacking secondary growth (Fig. 5a): moving from the center to the outermost layer, it consists of central pith, a single ring of collateral vascular bundles, collenchyma cells for mechanical support, and epidermis. Each vascular bundle contains a pole of heavily lignified xylem oriented towards the pith and a pole of phloem positioned towards the epidermis (Fig. 5b,c).

Cross-sections of the AqJAG-silenced inflorescence stems all exhibited a reduced amount of collenchyma cells and a clear decrease in the amount of differentiated xylem (Fig. 5d–f). Further, the existing xylem was much less lignified compared with xylem of WT (Fig. 5c vs 5f). Most strikingly, we observed distinct differences in the morphology of vascular bundles in the region adjacent to the outgrowth as compared to within the outgrowth itself (Fig. 5e,g). Near the site of the outgrowth, the vascular bundles were clustered with no normal pattern and appeared largely to lack xylem (Fig. 5d,g). By contrast, the vascular bundles present in the outgrowths possessed well-developed xylem, but their arrangement more closely resembles that seen in a leaf (Fig. 5g).

AqJAG is essential for floral organ primordium initiation and expansion

A WT A. coerulea flower possesses five petaloid sepals in the outermost whorl, a whorl of five petals with pronounced nectar spurs, five to ten whorls of stamens, a whorl of ten staminodia, and five carpels in the innermost whorl (Fig. 6a–d). The petal spurs elongate very rapidly during stage 11 and reach a final length of 4–5 cm (Fig. 6a–c).

About 40% of AqJAG VIGS-treated plants displayed significant floral defects, which fall into four categories: defects in the distal growth of lateral organs, including jagged petal blades (Fig. 6e), jagged sepal margins (Fig. 6f), abolished and/or underdeveloped anthers (Fig. 6k–m), and petal spurs with reductions in length and width (Fig. 6f,g,l,n); mosaic organs, for which the most commonly observed were petal–sepal (Fig. 6h) or petal–stamen chimeras (Fig. 6i); missing floral organs, (Fig. 6j–o), for which missing carpels (Fig. 6k–o), staminodia (Fig. 6k–o), and

Fig. 3 Scanning electron microscopy of wild-type (WT) Aquilegia coerulea and AqJAG-silenced leaves. (a) Close-up of leaflet with the most severe silencing phenotype (inset shows lateral leaflet before fixation, also boxed in Fig. 2d). (b, c) Cells on the (b) abaxial and (c) adaxial surface of WT leaves. (d, e) Cells on the (d) abaxial and (e) adaxial surface of AqJAG-silenced leaves. s, segments of leaflets. Bars: (a) 200 μm; (b, d) 100 μm; (c, e) 50 μm.
stamens (Fig. 6k–o) were observed more often than missing petals (Fig. 6j,n,o) and sepals (Fig. 6n); a small number of ‘flowers’ failed to develop any floral organs, usually resulting in a small bulge of tissues with undefined identity (Fig. 6p,q). We considered this to be the most severe silencing phenotype. We further analyzed a number of flowers with missing reproductive organs using SEM to investigate whether this phenotype was resulted from initiated but underdeveloped primordia, but no initiated primordium was found in any sample and the center meristem was always completely consumed (e.g. Fig. 6m).

Fig. 4 AqJAG-AqANS-TRV2-treated Aquilegia coerulea plants shows a significant decrease in the length of the first inflorescence internode and peculiar outgrowths on the stem. (a) First internode lengths were measured from the primary inflorescence of all virus-induced gene silencing (VIGS) control plants (con), and the primary inflorescence (1st) and secondary inflorescence (2nd) of all AqJAG-AqANS-TRV2-treated plants. When there were multiple secondary inflorescences on an AqJAG-silenced plant, the average of the first internode lengths of all secondary inflorescences was taken. Error bars indicate ± SE and asterisks indicate the significance levels of Student’s t-test with the control group: *, P < 0.05; ***, P < 0.0005. (b) A VIGS control plant. (c) An AqJAG VIGS plant showing reduced internode lengths and secondary inflorescence. (d) An AqJAG VIGS inflorescence with two regions of stem exhibiting outgrowth. (e) Outgrowth on the stem sometimes was torn apart into segments due to the elongation of the internode. (f) Inflorescence was bent downwards due to the outgrowth on the stem. Brackets, first internode of inflorescences; arrows, outgrowths; arrowheads, adjacent inflorescence stems without outgrowths. Bars: (b, c) 5 cm; (d–f) 1 cm.
Discussion

In the present study, we investigated the developmental roles of the *Aquilegia coerulea* JAGGED (*AqJAG*) ortholog, allowing us to further explore its functional conservation across various taxa. Strong expression of *AqJAG* is detected in the shoot apex, floral meristem, all initiating organ primordia, and regions undergoing rapid growth, such as the developing petal spur and leaflet. Silencing of *AqJAG* led to widespread defects in both meristems and lateral organs. Leaflets are unable to expand when lacking *AqJAG* and are

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**Fig. 5** Histology of (a–c) wild-type (WT) and (d–g) *AqJAG*-silenced *Aquilegia coerulea* inflorescence stems. (a) Cross section of a WT inflorescence stem. (b) False-colored overlay of (a) with xylem in magenta and phloem in blue. (c) Close-up of a WT vascular bundle. Xylem cells are on the left in the panel, discernable by pinkish staining in their lignified cell walls. (d) Cross section of an *AqJAG*-silenced inflorescence stem with outgrowth. (e) False-colored overlay of (d) with xylem in magenta and phloem in blue. (f) Close-up of a vascular bundle in *AqJAG*-silenced plants. Note the almost complete absence of lignified xylem. (g) Close-up of the outgrowth of an *AqJAG*-silenced stem. Arrowheads point at the disorganized bundles with no discernable xylem in the region adjacent to the outgrowth. Bars: (a, d, g) 200 μm; (c, f) 100 μm.
Fig. 6 Phenotypes of AqJAG virus-induced gene silencing (VIGS) flowers of Aquilegia coerulea. (a–d) Wild-type and VIGS control flowers. (a) Front view of a fully opened wild-type (WT) flower. (b) Side view of a fully opened control flower. (c) A developing pre-anthesis WT flower; during this stage, red pigmentation is just starting to appear in petals and sepals. (d) A pre-anthesis stage WT flower with perianth removed. (e–q) Phenotypes of AqJAG-AqANS-TRV2-treated flowers. (e) Flower with jagged petal margins. (f) Pre-anthesis flower with reduced sepal size, jagged sepal margins, reduced petal spur. (g) A fully opened flower with petal spurs reduced to different lengths (sepals removed). (h) Sepal–petal mosaic organ. (i) Petal–stamen mosaic organ. (j) Flower with two petals completely missing. (k) Flower with abnormal stamens (arrows) and missing carpels (perianth removed). (l) SEM of a mature flower (one sepal removed). Petal spurs failed to elongate; only few stamens developed but were abnormal; carpels were missing. (m) SEM of a mature flower (perianth removed). Only five stamens developed in the inner whorls and anthers were abnormal. (n) A mature flower having only three petals and one sepal (which was removed in the picture). (o) A terminal flower with only leaf-like organs developed. (p) A mature terminal ‘flower’ with no established floral organs (arrowhead). (q) An aborted axillary ‘flower’ (arrowhead). Bars: (a–g, j) 1 cm; (h–i, n–q) 5 mm; (k) 1 mm; (l, m) 500 μm.
concave due to changes in cell number and adaxial cell differentiation. Peculiar lamina-like outgrowths were observed on inflorescence stems, as well as reduction in the expansion of distal regions of floral organs. Some reproductive meristems terminated early or failed to produce primordia at all. Based on these results, we conclude that AqJAG is involved in multiple key aspects in A. coerulea morphogenesis, including initiation of primordia and promotion of cell proliferation and differentiation, with a disproportionate effect on the behavior of cells with adaxial identity.

**AqJAG is critical to floral primordium initiation in A. coerulea**

During the earliest stages of A. coerulea flower development, AqJAG is expressed across the floral meristem and in all initiating floral organ primordia (Fig. 2c–f). Consistent with this, many AqJAG VIGS flowers had missing organs, and the strongest silencing phenotypes failed to establish any floral organs at all. In less severe phenotypes, we observed mosaic floral organs, which have also been reported from the opb mutant in rice (Horigome et al., 2009). The opb phenotype is considered to reflect a role as a positive regulator of the floral organ identity genes, but we do not believe that the chimeric organs in our study were due to specific interactions between AqJAG and the identity genes. When recovered, the combination of identities present in the mosaic organs was highly variable, as was the number of mosaic organs in each flower. We believe that this is most likely due to a mispositioning of primordia on the floral meristem, which may be caused, in turn, by differential rates of proliferation across meristems that are partially silenced for AqJAG. Analogous phenotypes have been observed in shoot meristemless-2 mutants in Arabidopsis that similarly show uncoordinated proliferation in their floral meristems (Clark et al., 1996).

Interestingly, these effects on meristem function seem to be more severe in the reproductive phase than the vegetative phase. Although we only recovered clearly aborted meristems in inflorescences, we recognize that it may have been too difficult to recognize a comparable phenotype in the vegetative meristem, which is completely obscured by previously produced leaves. Regardless, in all other previously examined JAG homologs, expression was limited to primordia themselves rather than the meristem and, in null mutants, the initiation of lateral organs seemed unaffected (Dinneny et al., 2004; Ohno et al., 2004; David-Schwartz et al., 2009; Horigome et al., 2009; Schiessl et al., 2012). This suggests that the function of AqJAG in promoting cell proliferation may be expanded in Aquilegia relative to what is seen in other model systems.

**AqJAG controls laminar expansion in Aquilegia coerulea**

Consistent with earlier studies, laminar reduction is observed in AqJAG-silenced leaves. However, the degree of lamina reduction we recovered is on the severe end of the phenotypic spectrum when compared with the jag, nub, lyr or opb mutant phenotypes. SEM showed that the extremely stunted structures from the strongest AqJAG-silencing phenotypes were, in fact, established leaflet segments with distinct petioles, but laminar expansion is entirely lacking (Fig. 7a). During normal A. coerulea rosette leaf

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**Fig. 7** Schematic that represents the relative growth of abaxial and adaxial tissues in both stems and leaves of a variety of Aquilegia coerulea plants. In wild-type plants, stems and leaves show clear abaxial–adaxial polarity as reflected by phloem and xylem differentiation (cross-hatching and circles, respectively), as well as differentiation of other polar cell types (blue abaxial, cells; dark pink, adaxial cells). Strong AqJAG silencing results in reduction in the proliferation of adaxial cell types along with improper differentiation, causing the formation of unexpanded stems and laminae. When the silencing of AqJAG is uneven, patches of unaffected adaxial tissue proliferate unevenly within the stem, which produces abaxial–adaxial interface zones that form ectopic laminar outgrowths along the stem. In leaves, uneven silencing result in distorted leaflets with localized patches of reduced, concave laminae. The AqJAG phenotypes are in distinct contrast to what is observed in true abaxialized mutants, in which vasculature and lateral organs are strongly radialized, something never observed in AqJAG-silenced plants.
morphogenesis, the segments of the leaflets become recognizable very soon after leaflet primordium initiation, whereas petiole elongation and the progressive expansion of laminar cells occurs much later in development (Pabón-Mora et al., 2013). Within strongly silenced leaves, the most severe phenotypes are observed in lateral leaflets and lobes (Fig. 2d,h). Therefore, our observed defects are not just restricted to the distal region of growing lamina as seen in other jag mutants; cell proliferation in the entire leaflet lamina appears to be dramatically reduced when lacking AqJAG.

On the other hand, leaflet initiation itself does not appear to be dependent on AqJAG. Although there was no change in leaflet numbers in AqJAG-silenced plants, the lyr mutant in tomato exhibits reduced primary leaflet numbers, especially for the basal lateral pairs (David-Schwartz et al., 2009). This difference could be due to a number of factors, including the fact that Aquilegia and tomato compound leaves are independently derived, and potential differences in the genetic programs of trifoliate vs pinnately compound leaves (Bharathan et al., 2002; Harrison et al., 2005). Examining JAG homolog functions in additional taxa bearing different types of compound leaves will be needed to determine which pattern of JAG function is more common.

Another difference between silencing of JAG in A. coerulea and what has been observed in other taxa is that our laminar expansion phenotypes were much more severe in vegetative organs than in floral organs. In fact, the weaker aspect of floral phenotypes for AqJAG-silenced flowers are much more reminiscent of jag or jag nub mutants (Dinneny et al., 2004; 2006; Ohno et al., 2004). Laminar expansion is primarily affected in distal regions, leading to jagged margins or failure to develop anthers. It is intriguing to see reduced nectar spur development, although not especially surprising because we have previously established that localized cell divisions are critical to formation of the spur pocket (Puzey et al., 2012; Yant et al., 2015). Further study will be required to understand why laminar expansion in vegetative organs of Aquilegia appears to be so much more dependent on AqJAG function than that of floral organs.

AqJAG disproportionally influences cells with adaxial identity

Perhaps the most interesting aspect of the AqJAG-silencing phenotype is that there is a disproportionate impact on cells with adaxial identity in both leaves and stems. Among all AqJAG-silenced leaflets, a concave laminar surface was always associated with lamina reduction, which appears to result from a change in both adaxial cell number and differentiation (Fig. 3). It is hard to tell at this point whether the reduction in abaxial cell number is also directly due to loss of AqJAG or the change is indirect and due to compensatory interactions between the two domains (Hisanaga et al., 2015). Histological sectioning of affected stems revealed a substantial reduction of xylem but not phloem, which is consistent with reduced proliferation/differentiation of adaxial cell types (Emery et al., 2003). However, these vascular bundles are collateral rather than amphicribral, suggesting that the tissue is not truly abaxialized (Waites & Hudson, 1995). We, therefore, conclude that AqJAG differentially influences the behavior of adaxial cells rather than determining their identity; without its proper expression, adaxial cells fail to proliferate and differentiate in the vegetative tissues of A. coerulea.

The lamina-like outgrowth in AqJAG-silenced inflorescence axes is a rather peculiar phenotype, and to our knowledge, similar defects in stems have not been characterized previously in any JAG homolog mutants. We propose the following model to explain the phenomenon (Fig. 7). Wild-type stems are radially patterned with adaxial identity positioned internally and abaxial positioned externally. When the stem experiences broad and strong silencing of AqJAG, we would predict that vascular bundles remain collateral but with severely under-proliferated adaxial cells, including the xylem. However, due to the nature of VIGS, silencing is often patchy or temporally variable, which can lead to un silenced or weakly silenced patches in the stem where adaxial cells undergo relatively normal proliferation. The interaction between this kind of patch of adaxial identity and the surrounding abaxial tissue appears to create a margin that leads to laminar outgrowth, similar to what is normally seen in lateral organs (Waites & Hudson, 1995).

Although no previous study of JAG homologs has shown adaxial-specific defects as severe as ours, the phenotypes we obtained are not entirely surprising. Several lines of evidence suggest the involvement of JAG in polarity-related pathways and raise the possibility of an ancestral role for JAG homologs in adaxial cell development. For instance, both Lyr from tomato and NUB from Arabidopsis show adaxial-specific expression in lateral organs, with the NUB expression being consistently adaxial across many organ types (David-Schwartz et al., 2009; Dinneny et al., 2006). Although lyr mutants do not have dramatic adaxial defects, there is some loss or reduction of adaxial cell types combined with expansion of abaxial cell types. Ectopic expression of NUB promotes development of trichomes on the abaxial surface of leaves and the double nub jag mutant exhibits reproductive organ phenotypes that appear to be due to underproliferation/differentiation of cells with adaxial identity (Dinneny et al., 2006). The authors of that study hypothesized that JAG and NUB may promote the competence of tissues to respond to the polarity pathways, a theory that is consistent with our results. It is also interesting to consider the connections between JAG, organ polarity, and the BOP genes. The bop1-1 mutant in Arabidopsis displays laminar outgrowths on the petiole, somewhat analogous to our stem outgrowths (Ha et al., 2003). BOPs are known to antagonistically interact with JAG but also to differentially regulate the expression of adaxial determinants PHABULOSA and PHAVOLUTA and abaxial determinants KANI (Norberg et al., 2005; Ha et al., 2007). Further studies are necessary to understand the potential interactions between JAG/NUB and other polarity determinants in the major model systems, as well as A. coerulea.

Although researchers have achieved a fairly sophisticated understanding of how the interaction of adaxial–abaxial identity promotes lateral organ growth, we are still in the early stages of determining how these identities are fully realized in terms of
cell proliferation and development. Even when we consider a simple flat leaf, we observe complex differential regulation of cell growth and cell types in each domain. Our current results, therefore, provide evidence that cell proliferation could be controlled in an identity-specific manner such that adaxial cell division acts primarily through JAG homologs whereas abaxial cells rely on other pathways. Interestingly, this dependence on AqJAG in Aquilegia for laminar expansion may be more pronounced in vegetative organs than reproductive. Given that there are several genetic pathways to control cell proliferation (Doonan, 2000; Gutierrez, 2005; Hepworth & Lenhard, 2014) as well as adaxial identity (Fukushima & Hasebe, 2014), it is perhaps not surprising to find that over the course of evolutionary time, developmental system drift has led to different patterns of redundancy and functional dominance among these different genetic modules.

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Author contributions
The study was conceived of and designed by Y.M. and E.M.K. All experiments and data collection were conducted by Y.M. with training and oversight provided by E.M.K. The manuscript was written and figures prepared by Y.M. with revisions by E.M.K.

References


Supporting Information

Additional Supporting Information may be found online in the Supporting Information tab for this article:

Fig. S1 Predicted amino acid alignment of AqJAG, AtJAG, AtNUB, AtSUP and OsOPB.

Fig. S2 Maximum likelihood tree showing the relationships among amino acid sequences encoded by 42 AtJAG homologs and 14 Arabidopsis C2H2 zinc-finger proteins.

Fig. S3 Quantitative RT-PCR showing the downregulation of AqJAG.

Table S1 Stages of Aquilegia floral development.

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