

# **AGAMOUS mediates timing of guard cell formation during gynoecium development**

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## Abstract

In *Arabidopsis thaliana*, stomata are composed of two guard cells that control the aperture of a central pore to facilitate gas exchange between the plant and its environment, which is particularly important during photosynthesis. Although leaves are the primary photosynthetic organs of higher plants, floral organs are also photosynthetically active. In the Brassicaceae, evidence suggests that silique photosynthesis is important for optimal seed oil content. A group of transcription factors containing MADS DNA binding domains is necessary and sufficient to confer floral organ identity. Elegant models, such as the ABCE model of flower development and the floral quartet model, have been instrumental in describing the molecular mechanisms by which these floral organ identity proteins govern flower development. However, we lack a complete understanding of how the floral organ identity genes interact with the underlying leaf development program. Here, we show that the MADS domain transcription factor AGAMOUS (AG) represses stomatal development on the gynoecial valves, so that maturation of stomatal complexes coincides with fertilization. We present evidence that this regulation by AG is mediated by direct transcriptional repression of the master regulator of the stomatal lineage, *MUTE*, and that this interaction is conserved among the Brassicaceae. This work extends on our understanding of the mechanisms underlying floral organ formation and provides a framework to decipher the mechanisms that control floral organ photosynthesis.

## Introduction

In eudicots, such as *Arabidopsis thaliana*, flowers are composed of four types of floral organs: sepals, petals, stamens, and carpels. The sepals contain high levels of chlorophyll and bear stomata, making them leaf-like in appearance. In contrast, mature petals and stamens lack substantial concentrations of chlorophyll (Smyth *et al.*, 1990; Pyke and Page, 1998). Stomata are absent from petals but modified stomata are present on the abaxial surfaces of anthers (Smyth *et al.*, 1990; Nadeau and Sack, 2002a). In *A. thaliana*, the ovules are encased within the gynoecium which is formed of two fused carpels and other tissues that arise from the carpels, such as the style, stigma and replum (Smyth *et al.*, 1990; Alvarez and Smyth, 2002). Although the gynoecial valves lack stomata, they are present on the differentiated and elongated silique valve epidermis (Geisler *et al.*, 1998; Gu *et al.*, 1998). Stomata on siliques enable atmospheric carbon fixation to support local photosynthesis and photosynthetic activity of siliques has been demonstrated in several members of the Brassicaceae family including *A. thaliana* (Sheoran *et al.*, 1991; Gammelvind *et al.*, 1996; Wang *et al.*, 2016; Zhu *et al.*, 2018). This photosynthetic activity positively influences seed oil content and is of interest to crop breeders (Hua *et al.*, 2012; Wang *et al.*, 2016; Zhu *et al.*, 2018). Stomata also support transpiration, which drives the movement of nutrients through the plant and simultaneously facilitates cooling (Cramer *et al.*, 2009). However, very little is known about the molecular mechanisms of stomatal development on these organs.

On leaves of *A. thaliana*, the stomatal cell lineage is initiated by the asymmetric cell division of a protodermal cell (or meristemoid mother cell), which is regulated by the basic helix-loop-helix (bHLH) transcription factor SPEECHLESS (SPCH) (MacAlister *et al.*, 2007; Dong and Bergmann, 2010; Torii, 2021) ([Supplemental Figure 1A](#)). This asymmetric division produces a meristemoid and a stomatal lineage ground cell (SLGC). The meristemoid may then undergo further asymmetric cell divisions to renew itself and produce more SLGCs. Alternatively, a meristemoid may transition into a rounded cell termed a guard mother cell (GMC) (Dong and Bergmann, 2010; Torii, 2021). The transition from meristemoid to GMC is coordinated by another bHLH transcription factor, MUTE, which also functions to attenuate asymmetric cell divisions (Pillitteri *et al.*, 2007). GMCs then divide symmetrically to produce two guard cells and ultimately a mature stomatal complex, a process that is mediated by a third bHLH transcription factor, FAMA (Ohashi-Ito and Bergmann, 2006). Two other bHLHs, SCREAM (SCRM) and SCRM2, form heterodimers with SPCH, MUTE, and FAMA to coordinate gene expression (Kanaoka *et al.*, 2008). These division steps are regulated such that stomatal

complexes are separated by at least one cell, which ensures control of the central pore aperture (Dong and Bergmann, 2010; Torii, 2021). Several transmembrane receptors have been implicated in this stomatal patterning, such as TOO MANY MOUTHS (TMM), ERECTA (ER), ER-LIKE1 (ERL1), and ERL2 (Nadeau and Sack, 2002*b*; Shpak *et al.*, 2005; Torii, 2021). The activities of these receptors are modulated by the EPIDERMAL PATTERNING FACTORS (EPFs) and EPF-LIKES (EPFLs) secreted peptide families ([Supplemental Figure 1A](#)) (Hara *et al.*, 2007; Hunt and Gray, 2009; Kondo *et al.*, 2010; Torii, 2021).

Floral organ identity is controlled by a group of transcription factors that contain MADS DNA-binding domains (Bowman *et al.*, 1989, 1991*b*; O'Maoileidigh *et al.*, 2014*a*; Sablowski, 2015; Thomson and Wellmer, 2019). In the absence of their activities, floral organs are converted into leaf-like organs while ectopic expression of these transcription factors is sufficient to transform leaves into floral organs (Bowman *et al.*, 1991*b*; Pelaz *et al.*, 2000*a*, 2001; Honma and Goto, 2001; Ditta *et al.*, 2004). These observations confirmed a long-standing hypothesis that floral organs are derived from leaves (Goethe, 1790). They also formed the basis of the ABCE model of flower development and the floral quartet model, which largely address organ specification (Bowman *et al.*, 1991*b*; Pelaz *et al.*, 2000*b*, 2001; Honma and Goto, 2001; Ditta *et al.*, 2004; Theißen *et al.*, 2016). However, these MADS domain transcription factors continue to be expressed after the floral organs have been specified and they are known to control the expression of genes required for differentiation (Ito *et al.*, 2004, 2007; O'Maoileidigh *et al.*, 2014*b*; Ó'Maoiléidigh *et al.*, 2018). Identifying the repertoire of differentiation processes that the floral organ identity genes control remains a key challenge (Sablowski, 2015).

The MADS-domain protein AGAMOUS (AG) controls the specification of stamens and carpels and is required for floral meristem termination (Bowman *et al.*, 1989). AG interacts with E-class proteins, SEPALLATA 1-4 (SEP1-4), in heterodimeric or tetrameric complexes to coordinate carpel specification and floral meristem termination, with SEP3 playing an especially important role (Honma and Goto, 2001; Immink *et al.*, 2009; Theißen *et al.*, 2016). More recent data, however, suggests that the AG-SEP3 tetramers are only required for floral meristem termination rather than organogenesis (Hugouvieux *et al.*, 2018). AG activity promotes carpel development in a partially redundant manner with its closest related paralogs *SHATTERPROOF1* (*SHP1*) and *SHP2* (Favaro *et al.*, 2003). AG and the SHP proteins also suppress the formation of epidermal hairs (trichomes) during carpel differentiation, which represents the first

tangible example of how the floral organ identity proteins modify the underlying leaf development program to generate floral organs (Ó'Maoiléidigh *et al.*, 2013, 2018).

Here, we investigate the developmental progression and molecular regulation of stomatal development on gynoecial and silique valves in *A. thaliana*. We describe the normal progression of stomatal development before and after fertilization, and present evidence that AG suppresses this process in *A. thaliana* and other members of the Brassicaceae. The data presented provide further evidence and mechanism that transcription factors conferring floral organ identity directly suppress aspects of leaf development during floral organ formation. They also provide a framework with which to understand the establishment of the silique as a photosynthetic organ in the Brassicaceae.

## Materials and methods

### Plant growth and materials

Plants were grown on soil under cool white light at ~20°C in 16:8 h light:dark conditions. Plant lines used in this study are listed in [Supplemental Table 1](#). Flower development stages are after Smyth *et al.*, 1990.

### Genotyping

Genotyping PCRs were performed with genomic DNA extracted according to Edwards *et al.*, 1991. Tissue was disrupted using a TissueLyser II (Qiagen). Primers used in this study are listed in [Supplemental Table 2](#). The genotyping of *ag-10* and *ful-1* was performed as in Liu *et al.*, 2011 and (Ferrándiz *et al.*, 2000a).

### Confocal microscopy

A Zeiss Laser Scanning Microscope-780 or 880 (Zeiss) was used to generate the fluorescent images. Settings were optimized to visualize GFP, YFP (laser wavelength, 488 nm; detection wavelength, 493 to 598 nm) or chlorophyll (laser wavelength, 545 nm; detection wavelength, 604 to 694 nm). All direct comparisons of images were performed with the same settings. Maximum intensity projections were generated from z-stacks of multiple tiles to visualize the entire apical region of the gynoecium/silique. Samples were mounted on a glass slide with 1% low-melt agarose (Bio-Rad) and submerged in de-ionized water.

### Use of publicly available genomics data

M-values for the expression of all genes detected during flower development were previously generated (Ryan *et al.*, 2015). Transcripts per million for each biological replicate and time-point from Kivivirta *et al.*, 2020 were generously shared with us by Dr. Annette Becker and Clemens Rössner (Justus-Liebig Universität Giessen, Germany). ChIP-Seq peaks were visualized using online software at ChIP-Hub (<https://biobigdata.nju.edu.cn/ChIPHub/>) (Fu *et al.*, 2022). Seq-DAP-seq sequences were downloaded from the genome browser (<https://genome.ucsc.edu/s/ArnaudStigliani/MADS>) (Lai *et al.*, 2020).

### Scanning electron microscopy

Samples were fixed and processed for SEM as previously described in Laux *et al.*, 1996. Briefly, samples were harvested in 1 x PBS (P4417-100TAB, Sigma-Aldrich) at room

temperature. The PBS was then replaced with a 4% solution of 50% glutaraldehyde (4995.1, Roth) in 1 x PBS and incubated overnight at 4°C. A dehydration series was then performed using freshly prepared solutions of 30%, 50%, 70%, and 90% ethanol (v/v) at room temperature for 30 min each while being gently rocked. Samples were then incubated in 100% ethanol for 60 min at room temperature. Ethanol was then replaced with 100% ethanol and placed at 4°C. A Leica CPD300 or Quorum K850 were used for critical point drying and samples were coated with gold using an Emitech SC 7640 (Polaron) or Quorum Q150T sputter coater. Samples were imaged using a Hitachi TM 4000Plus or a Supra 40VP scanning electron microscope (Zeiss).

### **Chemical treatments**

A solution containing 10 µM dexamethasone (DEX; Sigma) and 0.015% Silwet L-77 in deionized water was applied liberally to the inflorescences of plants with a Pasteur pipette. Mock treatments contained all the same components without dexamethasone. Ethanol vapor treatments were performed by sealing the plants in containers (18 cm x 32 cm x 50 cm) with clear plastic lids and two 50 mL tubes that each contained 10 mL of 100% ethanol for 6 h.

### **Reverse transcription quantitative PCR**

Samples were harvested on liquid nitrogen and stored at -80°C until processed. Harvesting of gynoecia between stages 10-13 was done under a stereomicroscope from the primary inflorescences of 15-20 plants. Only gynoecia, where the majority of the stigmatic tissue was clear of pollen (i.e. largely unfertilized), were harvested. Harvesting of stage 13 gynoecia from the *AG-amiRNA<sup>i</sup>* line for the time-course (Figure 3C) was done by initially removing all flowers and siliques that were beyond stage 13. Then flowers at anthesis were harvested for the 0 d time-point. Plants were treated with DEX and 1 d later, 2-3 flowers from the same plants at anthesis were harvested on liquid nitrogen. This continued until 7 d after the treatment from 15-20 plants.

Samples were disrupted using a TissueLyser II (Qiagen) with adapters that were kept at -80°C. Total RNA was extracted from the samples using an RNeasy Plant Mini kit (Qiagen) or a Spectrum Plant Total RNA kit (Sigma-Aldrich). Removal of genomic DNA was performed with the Turbo DNA-free Kit (Invitrogen) or the On-Column DNase I Digestion set (Sigma-Aldrich). Reverse transcription was performed with the SuperScript™ IV First-Strand Synthesis System (Invitrogen) or RevertAid H Minus Reverse Transcriptase (ThermoFisher Scientific). Quantitative PCR was performed with

either iQ SYBR green (Bio-Rad) or PowerUp SYBR Green (Thermofisher Scientific) using a Lightcycler 480 (Roche) or Stratagene Mx3005 (Agilent Technologies). At least 2 technical measurements for each biological replicate were made and the mean of these technical replicates was used to represent each biological replicate. The number of biological replicates for each RT-qPCR experiment is indicated in [Supplemental Table 3](#). Melting curves were obtained for the reactions, revealing single peak melting curves for the amplified products. The amplification data were analyzed using the second derivative maximum method, and resulting values were converted into relative expression values using the comparative cycle threshold method (Livak and Schmittgen, 2001). Primers used in this study are listed in [Supplemental Table 2](#).

### **Motif enrichment analysis**

The 1421 sequences identified as bound by AG in Ó'Maoiléidigh *et al.*, 2013 were downloaded from [www.araport.org](http://www.araport.org) (Pasha *et al.*, 2020). These sequences were uploaded to the MEME-ChIP interface on [meme-suite.org](http://meme-suite.org) (Machanick and Bailey, 2011; Bailey *et al.*, 2015). The settings applied and motifs identified are summarized in [Supplemental Table 4](#).

### **Genomic sequence alignments**

The genomic and inferred amino acid sequences of *MUTE* were downloaded from The Arabidopsis Information Resource ([www.arabidopsis.org](http://www.arabidopsis.org)) (Berardini *et al.*, 2015). Orthologs of *AtMUTE* were previously identified in *Arabidopsis lyrata*, *Capsella rubella*, and *Eutrema salsugineum* (formerly *Thellungiella halophila*) (Ran *et al.*, 2013). We identified putative orthologs in *Capsella grandiflora*, *Arabidopsis halleri*, and *Boechera stricta* by using the TBLASTN functionality in Phytozome (<https://phytozome-next.jgi.doe.gov/>) with the *AtMUTE* amino acid sequence against the above genomes (Goodstein *et al.*, 2012). We found one clear candidate in each genome, which included the same orthologs identified in Ran *et al.*, 2013. The TBLASTN search results are presented in [Supplemental Table 5](#). Sequence alignments were performed with mVISTA (<https://genome.lbl.gov/vista/mvista/submit.shtml>) using standard settings (Frazer *et al.*, 2004).

### **Statistical analysis**

Some recommendations from Wasserstein *et al.*, 2019 are implemented here. The term “statistically significant” has been avoided and *p*-values have been reported as continuous



where possible. Although grouping data based on statistical thresholds is not recommended (Wasserstein *et al.*, 2019), a grouping threshold is provided in supplemental tables as a helpful guide, but results are not interpreted on the basis of these groupings. Data were analyzed using the R studio software (Team, 2022). For multiple comparisons, data were analyzed using a one-way analysis of variance (ANOVA). Post hoc tests were employed if a significant result was obtained from the above tests ( $p < 0.01$ ). The base function `pairwise.t.test` was used following an ANOVA test, with analysis being paired or unpaired depending on the experimental design. Adjustments for multiple corrections were performed in each case using the Benjamini–Hochberg method (Benjamini and Hochberg, 1995). For the box plots, the box encapsulates the 25th to 75th percentile, and the error bars encapsulate the 10th and 90th percentiles. The horizontal line running through the box indicates the median, and each point represents an individual plant.

### **Gel shift assays**

Electrophoretic mobility shift assays were performed essentially as described by Melzer *et al.*, 2009. DNA probes were ordered as single stranded oligo, annealed, and radioactively labelled with [ $\alpha$ - $P^{32}$ ] dATP by a Klenow fill-in reaction of 5'-overhangs. Probe sequences are shown in [Supplementary Table 6](#). Proteins were produced *in vitro* via the TNT SP6 Quick Coupled Transcription/Translation System (Promega) following manufacturer's instructions. Plasmids for *in vitro* transcription/translation of SEP3 and SEP3 $\Delta$ C have been generated previously (Melzer, et al. 2009). For generation of the plasmids for *in vitro* transcription/translation of AG, SHP1, and SHP2, coding sequences of AG (X53579.1), SHP1 (NM\_115740.3), and SHP2 (NM\_129844.5) were synthesized via the GeneArt gene synthesis service (Thermo Fisher Scientific) and cloned into pTNT (Promega) using *Eco*RI and *Sal*I restriction sites. The composition of the protein-DNA binding reaction buffer was essentially as described by Egea-Cortines *et al.*, 1999 with final concentrations of 1.6 mM EDTA, 10.3 mM HEPES, 1mM DTT, 1.3 mM spermidine hydrochloride, 33.3 ng  $\mu$ l<sup>-1</sup> Poly dI/dC, 2.5 % CHAPS, 4.3 % glycerol, and 6  $\mu$ g  $\mu$ l<sup>-1</sup> BSA. For each binding reaction 2  $\mu$ l of *in vitro* transcribed/translated protein were co-incubated with 0.1 ng of radioactively labelled DNA probe in a total volume 12  $\mu$ l. Binding reactions were incubated overnight at 4°C and loaded on a polyacrylamide (5% acrylamide, 0.1725% bisacrylamid) 0.5 $\times$  TBE gel. The gel was run at room temperature with 0.5 $\times$  TBE buffer for 3 h at 7.5 V cm<sup>-1</sup>, and afterwards vacuum-dried and exposed onto a phosphorimaging screen to quantify signal intensities. For size comparison a

radioactively labelled DNA ladder (100 bp DNA ladder, New England Biolabs) was applied.

## Results

### Stomatal development on gynoecial valves

As mentioned above, the photosynthetic activity of siliques of the Brassicaceae family contributes significantly to the carbon requirements of seeds (Hua *et al.*, 2012; Wang *et al.*, 2016; Zhu *et al.*, 2018). Siliques assimilate atmospheric CO<sub>2</sub> (Sheoran *et al.*, 1991; Gammelvind *et al.*, 1996; Hua *et al.*, 2012), probably through the stomata present on the valve epidermis, which are not present on pre-anthesis gynoecial valves ([Supplemental Figure 1B](#)) (Smyth *et al.*, 1990). The presence of stomata on floral organs was previously reported (Smyth *et al.*, 1990; Geisler *et al.*, 1998; Nadeau and Sack, 2002a) ([Supplemental Figure 1C-E](#)), however, the progression of stomatal development on these organs has not been fully described. We addressed this knowledge gap by examining the developmental progression of stomata on the siliques of *A. thaliana*.

We started by surveying the mRNA levels of master regulators of stomatal development as reported by publicly available transcriptomics datasets of flower and gynoecium development ([Figure 1A-B](#)). We focused on the expression of three bHLH transcription factor-coding genes, *SPCH*, *MUTE*, and *FAMA*, which are necessary and sufficient to promote different stages of the stomatal lineage (Ohashi-Ito and Bergmann, 2006; MacAlister *et al.*, 2007; Pillitteri *et al.*, 2007). A distinct peak of *SPCH* and *MUTE* mRNA was detected at stage 6-7 and 8 of flower development, respectively, in genome-wide expression profiling using a synchronous flowering system ([Figure 1A](#)) (Ryan *et al.*, 2015). These peaks probably corresponded to the initiation and progression of the stomatal lineage on sepals, which began to mature at approximately stage 10 ([Supplemental Figure 1C](#)). Levels of both *SPCH* and *MUTE* mRNA then decreased until anthesis, although *MUTE* levels plateaued between stages 11 and 13. In contrast, *FAMA* levels continued to increase as flower development progressed ([Figure 1A](#)). A more recent transcriptomics datasets derived from laser-microdissected gynoecia at different stages of flower development provided improved spatial resolution (Kivivirta *et al.*, 2020). These data are not complicated by the presence of other floral organs that bear stomata, such as the sepals and stamens, although the latest stage analyzed was stage 12 of flower development. *SPCH* mRNA levels were relatively high from approximately stage 5 and rose slightly until stage 11 and then decreased ([Figure 1B](#)). *MUTE* mRNA was not detectable before stage 11 when it was weakly expressed and then increased ~3-fold by stage 12 ([Figure 1B](#)). *FAMA* mRNA was also not detected at stage 5 but increased steadily by ~8-fold between stages 9 and 12 ([Figure 1B](#)). Based on these data, we

concluded that *MUTE* and *FAMA* transcription in the gynoecium was initiated at approximately stage 11-12, whereas *SPCH* transcription was initiated earlier.

To cross-examine these transcriptomic data and to provide a better resolution of the expression of these genes in the gynoecium, we obtained transgenic lines that harbor fluorescent translational or transcriptional reporters for *SPCH* (Davies and Bergmann, 2014), *MUTE* (Pillitteri *et al.*, 2007), and *FAMA* (Adrian *et al.*, 2015) whose transcription is driven by their endogenous regulatory elements (Figure 1C-E). Using laser-scanning confocal microscopy, we could not detect SPCH-YFP in stage 11 gynoecial valves, but the SPCH-YFP protein was abundant in the epidermis of late stage 12 and stage 13 gynoecial valves (Figure 1C). The contrast between mRNA and protein accumulation may indicate the presence of a post-translational mechanism to control SPCH protein levels (Figure 1B, C). A pattern of protein accumulation was found for MUTE-GFP, which was largely in agreement with the transcriptomics data in gynoecia (Figures 1B and 1D). The accumulation of YFP from the *FAMAPro:NLS-2xYFP* transgene was delayed relative to either of these reporters, with few fluorescent foci present at stage 12, which became slightly more abundant at stage 13 (Figure 1E). At stage 14, accumulation of YFP was observed as fluorescent foci throughout the gynoecial valve (Figure 1E). Therefore, the SPCH and MUTE proteins accumulate after stage 11 but before late stage 12, whereas *FAMA* transcription is initiated after late stage 12.

We then examined the formation of stomatal cell lineage types on gynoecial valves using scanning electron microscopy (SEM). We categorized the stomatal lineage into ‘early’ (including meristemoids and young GMCs), ‘mid’ (including GMCs before signs of asymmetric cell division were observed), and ‘late’ stages (including GMCs where asymmetric cell division had started through to mature stomata). We observed early-stage stomatal lineage cells on gynoecial valves during stage 12 of flower development whereas more mature stomatal cell lineages were largely absent (Figure 2A, Supplemental Table 7). At stage 13, early-stage lineage cells remained most common, however, mid-stage cells became more prevalent and late-stage cells were occasionally observed (Figure 2B, Supplemental Table 7). Post-fertilization, at stage 15-16, the frequency of early-stage cells reduced dramatically with a commensurate increase in mid-stage and late-stage cells (Figure 2C, Supplemental Table 7), until most stomatal lineage cells were of late-stage on older (>stage 17) siliques (Figure 2D, Supplemental Table 7). Stomata also formed on the style of the gynoecium prior to the formation of stomata on the valves, with mature stomata present on the style of stage 12 gynoecia (Supplemental Figure 1E). However, the style comprises only a small fraction of the total area of the

mature silique ([Supplemental Figure 1F](#)), suggesting that stomata on the style contribute only a small fraction of the total carbon fixation and transpiration capacity of the silique.

Given the close association between fertilization and progression of stomatal development, we tested whether a dependent relationship existed ([Supplemental Figure 2](#)). To this end, we emasculated flowers before anthesis and examined gynoecial valves 5 d after anthesis by SEM ([Supplemental Figure 2A](#)). We found mature stomatal complexes on these unfertilized gynoecial valves ([Supplemental Figure 2B](#)), which indicated that maturation of stomatal complexes on the gynoecium does not depend on fertilization.

### **Expression of *AG* and *SEP3* during the formation of stomatal complexes**

The C class protein *AG* controls the specification and development of stamens and carpels by forming dimers or tetramers with the E class protein *SEP3* (Bowman *et al.*, 1989; Pelaz *et al.*, 2000a; Honma and Goto, 2001; Hugouvieux *et al.*, 2018). Notably, it has been shown that *AG* suppresses the formation of trichomes on carpel valves by controlling the expression of other developmental regulators (Ó'Maoiléidigh *et al.*, 2013, 2018). In addition, trichomes form on the gynoecia of plants with reduced *SEP* activity (Ditta *et al.*, 2004). Because stomata, like trichomes, are formed by the epidermis, we hypothesized that an *AG-SEP3* complex might also repress the formation of stomata on carpel valves. To explore this, we first assessed published expression data for *AG* and *SEP3* in gynoecial tissues over the course of stomatal initiation and maturation ([Figure 2E-F](#), [Supplemental Figure 3](#)). *In situ* hybridizations showed *AG* mRNA accumulation throughout the carpels during stage 8 before its expression becomes restricted and absent from stage 12 gynoecial valves (Bowman *et al.*, 1991a). A similar expression pattern was observed of *SEP3* mRNA in the gynoecium, although its expression was restricted from gynoecial valves from approximately stage 10 (Mandel and Yanofsky, 1998). This is consistent with the results of transcriptomics experiments, where *AG* and *SEP3* mRNA levels begin to decrease from approximately stage 11 of flower development ([Figure 2E-F](#)) (Ryan *et al.*, 2015; Kivivirta *et al.*, 2020).

*AG-GFP* and *SEP3-GFP* protein accumulation was analyzed in transgenic plants that harbored *AGpro:AG-GFP* and *SEP3pro:SEP3-GFP* transgenes that drive transcription from the corresponding endogenous regulatory elements (Urbanus *et al.*, 2009). *AG-GFP* fluorescence was observed throughout stage 12 gynoecial valves in these plants even though *AG* mRNA is not present at this stage (Bowman *et al.*, 1991a), possibly due to the low turnover of *AG* at these stages (Urbanus *et al.*, 2009). We verified

these results for AG-GFP using a similar but independently generated *AGpro:AG-GFP* transgenic line using laser-scanning confocal microscopy (Supplemental Figure 3A-B) (Ó'Maoiléidigh *et al.*, 2013). We also imaged stage 13 gynoecia and found that fluorescent signal stemming from AG-GFP was more intense in the replum and style relative to the valves (Supplemental Figure 3C). Additionally, SEP3-GFP protein accumulated to much higher levels in the replum and valve margins at stage 12 when compared to the gynoecial valves (Urbanus *et al.*, 2009). Taken together with the transcriptomics data, it appears that the abundance of AG and SEP3 proteins in the gynoecial valves reduces significantly between stages 11 and 13 of flower development.

### **Transcriptional response of master regulators of stomatal development to AG perturbation**

To test whether a perturbation of AG activity would result in the differential expression of master regulators of stomatal development, we harvested stage ~10-13 gynoecia from *L-er* wild-type plants harboring a fully functional copy of AG and those that were homozygous for the weak *ag-10* mutation (Liu *et al.*, 2011). Plants that are homozygous for *ag-10* form gynoecia that are relatively normal in appearance, although they tend to bulge (Liu *et al.*, 2011). We extracted total RNA from these plants and using reverse-transcription combined with quantitative PCR (RT-qPCR), we found that the mean mRNA levels of *MUTE* and *FAMA* were increased in *ag-10* carpels to ~3.4 ( $p = 0.0002$ , two-tailed paired t-test) and ~1.9-fold ( $p = 0.0004$ , two-tailed paired t-test), respectively, relative to wild-type counterparts (Figure 3A, Supplemental Table 3). In contrast, the mean levels of *SPCH* mRNA were mildly decreased to 0.8-fold ( $p = 0.09$ , two-tailed paired t-test) while the mean mRNA levels of *SCRM* and *SCRM2* appeared unchanged (Figure 3A, Supplemental Table 3). Next, we used a transgenic line that harbors a dexamethasone (DEX) inducible artificial microRNA that is designed to target the AG mRNA (*AG-amiRNA<sup>i</sup>*) (Ó'Maoiléidigh *et al.*, 2013). We treated the inflorescences of these plants with a DEX-containing solution and a mock solution, and harvested stage ~10-13 gynoecia 24 h later. By RT-qPCR, we found that mean *MUTE* mRNA levels were increased to ~1.4-fold ( $p = 0.08$ , two-tailed paired t-test) in DEX-treated samples relative to the mock control (Figure 3B, Supplemental Table 3), consistent with the increased expression detected in *ag-10*. None of the other bHLH-coding genes were consistently differentially expressed (Figure 3B, Supplemental Table 3).

We then used the *AG-amiRNA<sup>i</sup>* line in a time-course experiment where we harvested stage 13 gynoecia of untreated plants (0 d) and then treated the inflorescences

of plants with a DEX-containing solution. We harvested only gynoecia that had reached anthesis every day for seven days following that treatment. Therefore, gynoecia that were harvested 1 d following DEX-treatment would have been ~late-stage 12 gynoecia at the time of DEX treatment. Gynoecia harvested at 2 d following DEX-treatment would have been ~early stage 12 gynoecia, and so on (Figure 3C). Following harvesting, we processed the samples and determined mRNA levels of *SPCH*, *MUTE*, *FAMA*, and *AG* using RT-qPCR. One day after treatment, there was very little change in the levels of *SPCH*, *FAMA*, or *MUTE* mRNAs whereas *AG* mRNA was reduced to 0.2-fold when compared with 0 d ( $p.adj < 10^{-9}$ , pairwise two-tailed t-tests), as has been previously described (Figure 3C, Supplemental Table 3) (Ó'Maoiléidigh *et al.*, 2013). The lack of response from the bHLH-encoding genes may have been because the *AG-miRNA* was activated in late-stage 12 gynoecia when *AG* mRNA levels were already very reduced in the gynoecial valve, although *AG* mRNA continued to accumulate in other tissues (Bowman *et al.*, 1991a). At 2 d post-induction, which roughly corresponded to activation of the *AG-miRNA* in early stage 12 gynoecia, the mRNA levels of both *MUTE* and *FAMA* increased to ~1.8 ( $p.adj = 0.006$ , pairwise two-tailed t-tests) and ~2-fold ( $p.adj = 0.02$ , pairwise two-tailed t-tests) relative to 0 d, respectively, as *SPCH* mRNA levels decreased to 0.6-fold ( $p.adj = 0.006$ , pairwise two-tailed t-tests) (Figure 3C, Supplemental Table 3). *MUTE* and *FAMA* expression then continued to increase until 4 d post-induction where they peaked at ~2.2 ( $p.adj = 0.0004$ , pairwise two-tailed t-tests) and ~3.9-fold ( $p.adj < 10^{-5}$ , pairwise two-tailed t-tests) relative to 0 d, respectively, and then began to decrease. At 5 d post-induction, *AG* mRNA levels began to recover, which correlated with a reduction in *MUTE* mRNA levels, which *FAMA* partially mimics, until the end of the experiment at 7 d post induction (Figure 3C, Supplemental Table 3). Taken together, we concluded that *AG* represses the expression of *MUTE*, and *FAMA* during gynoecium development.

To improve our understanding of the spatial resolution with which *AG* controls the expression of these stomatal genes, we introduced the *FAMAPro:2xYFP* reporter into the *ag-10* background and imaged gynoecia at late stage 12 and stage 13 using confocal laser scanning microscopy (Lee *et al.*, 2019). We observed very few fluorescent foci on the valves of stage 12 wild-type gynoecia and stage 13 valves (Figure 3D-E, Supplemental Table 8), as previously described for the *FAMAPro:NLS-2xYFP* reporter (Figure 1E). In contrast, over 20 times the number of foci were observed on the valves of stage 12 ( $p = 0.06$ , two-tailed paired t-test) and stage 13 *ag-10* gynoecia ( $p = 0.01$ , two-tailed paired t-test) (Figure 3D-E). We then introduced the *FAMAPro:2xYFP* reporter into

the *AG-amiRNA<sup>i</sup>* transgenic line. We treated inflorescences with a DEX-containing or mock solution and imaged stage 12 and 13 gynoecia every day for 3 days (Figure 3F-G, Supplemental Figure 4, Supplemental Table 8). Similar to the *ag-10* observations (Figure 3D-E) and mirroring the results of the RT-qPCR experiments (Figure 3C), fluorescent foci were prevalent on valves of the DEX-treated gynoecia at 48 and 72 h (Figure 3F-G). Far fewer fluorescent foci were observed on the valves of untreated samples (0 h, Figure 3F-G), mock-treated counterparts (Supplemental Figure 4B, D), or DEX-treated samples after 24 h (Supplemental Figure 4A, C). Statistical summaries of these *AG-amiRNA<sup>i</sup>* data are presented in Supplemental Table 8. These confocal data support the observations of the RT-qPCR experiments and clarify that expression of *FAMA* is not elevated throughout the gynoecial epidermis but only in a pattern that is consistent with stomatal formation on the gynoecial valves.

### **Interaction between AG, SHP, SEP3, and the first intron of MUTE**

*AG* activity is negatively correlated with stomatal development on gynoecial valves and perturbation of *AG* results in elevated mRNA levels of key stomatal regulators, such as *MUTE* and *FAMA*. To understand whether the regulation of *MUTE* and *FAMA* transcription was directly controlled by *AG*, we interrogated chromatin immunoprecipitation followed by next-generation sequencing (ChIP-Seq) data available for *AG* (Ó'Maoiléidigh *et al.*, 2013). We found that *AG* bound to the first intron of *MUTE* (Figure 4A), which was previously identified as a direct target (Ó'Maoiléidigh *et al.*, 2013). *MUTE* was also identified as a putative direct target of *SEP3* (Pajoro *et al.*, 2014) (Figure 4A), which forms heterodimers and quaternary complexes with *AG* to control floral organ development (Honma and Goto, 2001; Theißen *et al.*, 2016; Hugouvieux *et al.*, 2018). Within the first intron of *MUTE* are two motifs with similarity to CArG motifs, to which these MADS domain transcription factors bind (Wuest *et al.*, 2012; Ó'Maoiléidigh *et al.*, 2013; Pajoro *et al.*, 2014). Analysis of the genomic sequences bound by *AG* revealed an enrichment of CArG motifs, which were very similar to the motifs identified in the first intron of *MUTE* (Figure 4B, Supplemental Table 4). *MUTE* was also identified as a direct target of the *AG-SEP3* complex through sequential immunoprecipitation followed by DNA affinity purification and sequencing experiments (seq-DAP-Seq) (Supplemental Figure 5) (Lai *et al.*, 2020).

We performed gel shift assays to cross-validate these experiments and to determine whether both CArG motifs in the *MUTE* first intron are required for *AG* and *SEP3* binding. We synthesized probes with the same sequence as a region of the *MUTE*



first intron that contains the two CArG motifs (*AtMUTE\_i1*) and incubated them with the AG and SEP3 proteins (Supplemental Table 6, Figure 4C). We also used a truncated version of SEP3 (SEP3 $\Delta$ C), which has been shown to retain activity (Melzer *et al.*, 2009), to visualize shifts between homodimers of SEP3 and AG-SEP3 heterodimers. We found that incubation of SEP3, SEP3 $\Delta$ C, or AG individually resulted in the appearance of a single retarded fraction of the DNA probe, suggesting that homodimers of each of these proteins can bind the *MUTE* first intron (Figure 4C). In accordance to the similar size of the full-length proteins, the retarded fractions produced by SEP3 and AG migrated with approximately the same speed, whereas the retarded fraction produced by SEP3 $\Delta$ C migrated faster, likely due to the reduced protein size. Co-incubation of AG with SEP3 $\Delta$ C resulted in the appearance of an additional retarded fraction of intermediate electrophoretic mobility, probably corresponding to AG-SEP3 $\Delta$ C heterodimers. We also incubated the SHP1 and SHP2 paralogs of AG with SEP3 $\Delta$ C in separate reactions and observed a shifted band when SHP1 was incubated with SEP3 $\Delta$ C at a height similar to that of the putative AG-SEP3 $\Delta$ C heterodimers. Only a very weak band was present to suggest that a SHP2-SEP3 $\Delta$ C heterodimer could bind the *AtMUTE\_i1* probe and there was very weak support for SHP1 or SHP2 homodimer binding. Furthermore, incubation of all 4 proteins together did not result in a shift, suggesting that a multimeric complex does not form at the *MUTE* intron (Figure 4C).

Next, we tested whether the CArG motifs present in this region were required for binding. We incubated combinations of the above proteins with probes harboring modifications of the CArG motifs individually (*mCArG\_1*, *mCArG\_2*) or simultaneously (*mCArG\_1+2*), which were intended to disrupt protein binding (Supplemental Table 6). Similar results to the *AtMUTE\_i1* probe were obtained when *mCArG\_1* was incubated with the same combinations of proteins as above (Figure 4D). However, incubation with the *mCArG\_2* probe abolished the formation of bands corresponding to heterodimer formation. Furthermore, the formation of bands corresponding to homodimers were reduced significantly (Figure 4E). Disruption of both CArG motifs (*mCArG\_1+2*) eliminated all shifted bands (Figure 4F). This suggests that CArG<sub>2</sub> interacts with heterodimers and that both CArG motifs can interact with homodimers, although the tested proteins have the highest affinity for CArG<sub>2</sub>.

We compared the sequence of the first intron of *A. thaliana MUTE* to the sequences of the first intron from orthologs of *MUTE* from other members of the Brassicaceae family (Supplemental Table 5). We found that a region of approximately 60

bp was well-conserved, which contained both CArG motifs (Figure 4G). For CArG\_2, the AT track in the center of the CArG motif was well maintained in four species. However, a single T to C transition was observed in the AT track of CArG\_2 in *Capsella rubella*, *Capsella grandiflora* and *Eutrema salsugineum* (Figure 4J). Therefore, we tested whether AG, SEP3, and/or the SHP proteins could bind to these diverged CArG\_2 motifs. We chose the sequences from *C. rubella*, as the CArG\_2 motif was identical to *C. grandiflora* (Figure 4H), and *E. salsugineum* (Figure 4I), and generated probes that encompassed both CArG motifs. We observed near identical results when these probes were incubated with the above proteins when compared to the *AtMUTE\_i1* probe (Figure 4H-I). This suggests that the interaction between *MUTE*, AG, SEP3, and to a lesser extent SHP1, is conserved in the Brassicaceae.

### Stomatal development on gynoecial valves upon AG perturbation

We next asked whether the elevated transcription of *MUTE* and *FAMA* observed in the plants lacking complete AG activity was sufficient to modify stomatal formation on the gynoecial valves. Using SEM, we imaged the gynoecia and siliques of wild-type *L-er* and *ag-10* plants (Figure 5A-B, D-E, I-J). At stage 12, as described above, mid and late-stage stomatal lineage cells are largely absent from wild-type gynoecial valves. In contrast, stage 12 *ag-10* gynoecial valves have eight times the number of mid-stage stomatal lineage cells ( $p.adj = 0.0003$ , pairwise unpaired two-tailed t-tests) (Figure 5A-B, I, Supplemental Table 7). At stage 13, mid ( $p.adj = 0.02$ , pairwise unpaired two-tailed t-tests) and late-stage ( $p.adj < 10^{-7}$ , pairwise unpaired two-tailed t-tests) stomatal lineage cells were much more abundant on *ag-10* gynoecial valves in comparison to wild-type (Figure 5D-E, J, Supplemental Table 7). Concomitantly, we observed a severe reduction in the number of early-stage stomatal lineage cells ( $p.adj = 10^{-6}$ , pairwise unpaired two-tailed t-tests) (Figure 5D-E, J, Supplemental Table 7). We also observed a substantial increase in the number of late-stage stomatal lineage cells after perturbation of AG activity through the *AG-amiRNA* relative to uninduced plants ( $p.adj < 10^{-5}$ , pairwise unpaired two-tailed t-tests) or similarly treated *L-er* wild-type plants ( $p.adj < 10^{-5}$ , pairwise unpaired two-tailed t-tests) (Supplemental Figure 6A-D, Supplemental Table 9).

It has been previously observed that the mutant of another MADS-domain transcription factor-coding gene, *FRUITFULL* (*FUL*), does not bear stomata on the silique (Gu *et al.*, 1998). Notably, introgression of the *shp1-1 shp2-1* mutations into the strong *ful-1* mutant results in the recovery of stomata on siliques (Ferrández *et al.*, 2000b). We asked whether introduction of the *ag-10* mutant into the *ful-1* background would also

be sufficient to rescue the formation of stomata on siliques. We found stomatal formation was partially restored on *ag-10 ful-1* siliques although the stomata are not completely normal in appearance and are not distributed throughout the silique (Figure 5G-H). Mean expression of *SHP1* was not reduced ( $p = 0.87$ , two-tailed paired t-test) and *SHP2* was only mildly reduced 0.87-fold ( $p = 0.12$ , two-tailed paired t-test) in the *ag-10* background compared to *L-er* (Supplemental Figure 6E, Supplemental Table 3). Therefore, recovery of stomata on *ag-10 ful-1* siliques is probably independent of *SHP* activity. Together these data demonstrate that AG is required to suppress stomatal development during gynoecium development.

### Comparative activity of AG and SHPs during stomatal development

To understand more about the redundancy between AG and the *SHP* genes during stomatal development, we generated an *ag-10 shp1 shp2* triple mutant and examined the stomatal indices (i.e. the ratio cell belonging to the stomatal lineage to the total number of stomata and epidermal cells) on gynoecial and silique valves. Late-stage stomatal lineage cells were more common on the valves of triple mutant gynoecia at stage 12 relative to *ag-10* ( $p_{adj} = 0.03$ , pairwise two-tailed t-test). However, slightly fewer late-stage stomata formed on the valves of triple mutant gynoecia relative to *ag-10* at stage 13 (Figure 5B, C, E, F, I-J, Supplemental Table 7). We also tested the mRNA levels of *SPCH*, *MUTE*, and *FAMA* in this *ag-10 shp1 shp2* triple mutant background in comparison to parental genotypes (Figure 6A, Supplemental Figure 7E, Supplemental Table 3). Mean *SPCH* levels were reduced in the *ag-10 shp1 shp2* triple mutant relative to *ag-10* to ~0.84 fold, although the variation in expression was quite high ( $p = 0.55$ , pairwise two-tailed t-test) (Supplemental Figure 7E, Supplemental Table 3). Mean *MUTE* mRNA levels were elevated in the *ag-10 shp1 shp2* triple mutant relative to *ag-10* to ~1.33 fold ( $p_{adj} = 0.012$ , pairwise two-tailed t-test). *FAMA* mRNA levels were also elevated to ~1.4 fold in the *ag-10 shp1 shp2* triple mutant relative to *ag-10* ( $p_{adj} = 0.022$ , pairwise two-tailed t-test, Figure 6A, Supplemental Table 3). In contrast, levels of *SPCH*, *MUTE*, and *FAMA* mRNAs were not as strongly affected in the *shp1-1 shp2-1* double mutant background (Figure 6A, Supplemental Figure 7E, Supplemental Table 3). We concluded that the SHP proteins probably suppress the expression of *MUTE* in parallel with AG, which results in its elevated expression and the elevated expression of *FAMA* in the triple mutant background relative to *ag-10*.

Two other genes that are associated with AG binding are also involved in stomatal development, *ERL1* and *ERL2* (Ó'Maoiléidigh *et al.*, 2013). These genes encode

receptor-like kinases that control stomatal patterning in concert with TMM (Shpak *et al.*, 2005). Ligands, such as STOMAGEN (STOM) and EPF1, bind to complexes of TMM, ER, ERL1, and ERL2, to maintain stomatal spacing and density (Torii, 2021). Therefore, we tested the mRNA levels of these genes in response to *AG* and *SHP* perturbation. *ERL1* and *ERL2* expression was mildly decreased in *ag-10*, however, their mRNA levels were reduced to ~0.4 ( $p_{adj} = 0.038$ , pairwise two-tailed t-test) and ~0.5-fold ( $p_{adj} = 0.009$ , pairwise two-tailed t-test) in the triple *ag-10 shp1 shp2* mutant relative to *L-er*, respectively (Figure 6A, Supplemental Table 3). *EPF1* mRNA levels were elevated in both *ag-10* and the triple mutant to ~3-fold ( $p_{adj} = 0.009$ , pairwise two-tailed t-test) and ~4.3-fold ( $p_{adj} = 0.009$ , pairwise two-tailed t-test), respectively (Figure 6A, Supplemental Table 3). *EPF1* levels were elevated by ~1.4-fold in the triple mutant relative to *ag-10* ( $p_{adj} = 0.03$ , pairwise two-tailed t-test) (Figure 6A, Supplemental Table 3). Similarly, mean *STOM* expression was elevated in *ag-10* and the triple mutant by ~1.7 ( $p_{adj} = 0.06$ , pairwise two-tailed t-test) and ~1.9-fold ( $p_{adj} = 0.14$ , pairwise two-tailed t-test) (Figure 6A, Supplemental Table 3). In contrast to these observations, mean *TMM* mRNA levels were not changed in *ag-10* or the *ag-10 shp1 shp2* triple mutant backgrounds ( $p = 0.39$ , two-way ANOVA) (Supplemental Figure 7E, Supplemental Table 3). Furthermore, there was little change in the expression of these genes in the *shp1 shp2* mutant background (Figure 6A, Supplemental Figure 7E-F, Supplemental Table 3).

Although the accumulation of *ERL1*, *ERL2*, *STOM*, and *EPF1* mRNAs were changed in the *ag-10* and/or the *ag-10 shp1 shp2* mutant backgrounds, we did not observe stomatal clustering in the samples in which *AG* or *SHP* were perturbed (Figure 6B-D). This suggests that either the observed changes are not substantial enough to modify patterning or that an initial change in gene expression mediated by *AG* and/or *SHP* was buffered by the regulatory network to maintain patterning. Notably, combinations of *er*, *erl1*, and *erl2* mutations results in a reduction of silique length (Shpak *et al.*, 2004), which is similar to the reduction of silique length observed for *ag-10 shp1 shp2* plants relative to parental genotypes and wild-type (Figure 6E-F, Supplemental Table 10). Therefore, it is likely that *AG*-mediated control of *ERL1* and *ERL2* expression is related only to gynoecial growth rather than stomatal patterning.

## Discussion

The progression of stomatal development on the gynoecium/silique has not been previously described. Here, we show that stomatal development initiates on the gynoecial valves between late stage 11 and early stage 12 (Figure 2A-D, Supplemental Table 7).

Stomatal initiation in the gynoecium correlates with the increased expression of stomatal bHLH transcription factors during this period (Figure 1). Most stomates do not reach maturity until after fertilization, although their maturation does not depend on fertilization (Supplemental Figure 2). This fertilization-independent differentiation is reminiscent of other differentiation processes, such as the continued increase in chlorophyll concentration in the gynoecia of emasculated pistils (i.e. unfertilized) (Carbonell-Bejerano *et al.*, 2010), which also temporally correlate with but are not dependent on fertilization (Wagstaff *et al.*, 2009). Given that fertilization does not control the onset or progression of stomatal development on gynoecial valves, we searched for an alternative mechanism.

The floral organ identity gene *AG* has been implicated previously in the differentiation of the gynoecial valve and in the suppression of leaf-like traits (Ó'Maoiléidigh *et al.*, 2013, 2018). Here, we present evidence that *AG* forms a complex with *SEP3* to suppress the formation of stomata (Figure 5), which are also typical of leaves, until the point of anthesis when their expression dissipates from gynoecial valves (Figure 2E-F, Supplemental Figure 3). *AG* and *SEP3* accomplish this by directly repressing the transcription of *MUTE*, which encodes a key bHLH transcription factor required for stomatal development (Figure 3, 4). *MUTE* mRNA levels are elevated in the gynoecia of plants with reduced *AG* activity. Furthermore, the expression of *FAMA*, which is directly promoted by *MUTE* (Han *et al.*, 2018), is elevated when *AG* activity is perturbed (Figure 3). By visualizing *FAMA* expression using a transgenic reporter line (Lee *et al.*, 2019), we showed that perturbation of *AG* activity does not release *FAMA* expression throughout the epidermis of the gynoecium but only in what are likely stomatal lineage cells (Figure 3D-G). This suggests that SPCH drives *MUTE* expression in a similar manner as during leaf development and that *AG* counteracts this SPCH activity in stomatal lineage cells. *MUTE* expression gradually increases in stomatal lineage cells once *AG* protein levels are reduced after late stage 12, probably enabling SPCH to fully activate *MUTE* transcription (Figure 1). Therefore, stomatal lineage progression, but not entry, is modified by *AG*.

*AG* and *SEP3* bind to the first intron of *MUTE*, which was previously demonstrated by ChIP-Seq and seq-DAP-seq experiments. The bound regions contain two CARG motifs and we showed that each CARG motif influences binding in a different way (Figure 4). We confirmed that both homodimers and heterodimers of *AG* and *SEP3* can bind to the first intron of *MUTE*. We also showed that heterodimers of *SH1-SEP3* bind to the same region. Heterodimer binding is dependent on the presence of only one

of these CArG motifs (CArG<sub>2</sub>), whereas homodimers do not fully depend on the presence of CArG<sub>2</sub> although homodimer affinity is reduced in its absence. Mutagenesis of both CArG motifs, however, abolishes binding of both homodimers and heterodimers (Figure 4C-F). It has been suggested that AG-SEP3 quartets are required only to maintain floral meristem determinacy (Hugouvieux *et al.*, 2018). Here, we did not observe the formation of higher order complexes with the generated probes suggesting that homodimers or heterodimers are sufficient to suppress *MUTE* expression. This is consistent with the idea that quartets of AG-SEP3 are not functional in the control of differentiation (Hugouvieux *et al.*, 2018).

The observation that heterodimers of SHP1-SEP3 and, to a lesser extent, SHP2-SEP3 are also capable of binding to the *MUTE* first intron (Figure 4) is in keeping with observations of redundant activities between AG, SHP1, and SHP2 (Figure 5, Figure 6). At stage 12, stomatal lineage cells had progressed further in the triple mutant than *ag-10* while *shp1-1 shp2-1* mutants appeared similar to *L-er* (Figure 5). The progression of stomatal development on *ag-10* and the *ag-10 shp1 shp2* triple mutant was, however, similar at stage 13 although there was a large amount of variation associated with these measurements. Therefore, there is only weak phenotypic evidence to support the redundant control of stomatal development between AG and SHP. On a molecular level, there is stronger support for redundancy. Key regulatory genes, such as *MUTE* and *FAMA*, were differentially expressed to a greater degree in the triple mutant when compared to *ag-10* and were generally not differentially expressed in *shp1-1 shp2-1* double mutants relative to *L-er* (Figure 6). Patterning genes such as *ERL1*, *ERL2*, and *EPF1* show a similar trend although there is no change in stomatal clustering. As such, the effect of AG and SHP on *ERL1* and *ERL2* expression may be more related to gynoecium/silique growth than stomatal development. Considering both the molecular and phenotypic data, we conclude that functional redundancy between AG and SHP does play a role in the control of stomatal development and gynoecial/silique growth.

As previously suggested, the divergent expression patterns of AG, SHP1, and SHP2 underlie many of their functional differences. The SHPs are broadly expressed in the gynoecium, including the valves, from stage 7 but become restricted to the valve margins during stage 12 of gynoecium development. This expression pattern is largely imposed by the FUL transcription factor, which represses SHP expression in the valves. Notably, *ful* mutants lack stomata on the valves although their formation can be restored when SHP activity is simultaneously perturbed. We observed a similar scenario as stomata were also recovered on *ag-10 ful-1* mutant valves (Figure 5G-H). This supports

the idea that AG and the SHP proteins are suppressing stomatal development by repressing *MUTE* expression.

We propose that AG acts as a timer to control the emergence of stomata on the developing gynoecium/silique valve in *A. thaliana*. The conserved CArG motifs in the first intron of *MUTE* and the ability of AG, SEP3, and the SHPs to bind even slightly diverged sequences in *C. rubella* and *E. salsugineum* support the idea that this function is conserved in the Brassicaceae (Figure 4H-I). The correlation between stomatal development and fertilization suggests that atmospheric carbon assimilation by the silique stomata is used during silique photosynthesis to support seed development, particularly in terms of seed oil biosynthesis. Oils are stored in the seeds and mobilized during germination and early seedling development before autotrophic growth is established (Graham, 2008). Energy derived from silique photosynthesis supplements the accumulation of storage oils in the seeds in a variety of species including *A. thaliana* and *Brassica napus* (Sheoran *et al.*, 1991; Gammelvind *et al.*, 1996; Wang *et al.*, 2016; Zhu *et al.*, 2018).

The timing of stomatal formation on the silique valves may have been selected for a variety of reasons. Pre-anthesis formation of stomata may be disadvantageous due to selective pressure by pathogens (Wu and Liu, 2022). Pathogens can penetrate the plant through stomata and photosynthetic activity would probably be low in pre-anthesis gynoecia due to the shading effect of sepals. Furthermore, transpiration within a closed flower bud is unlikely to be particularly effective. These factors may have resulted in the selection for stomatal formation to coincide with flower bud opening. It is also possible that the timing of stomatal progression on gynoecial valves differs between species as stomatal development had not been completed even once siliques were exposed to light (e.g. stage 15-16 Supplemental Table 1). Therefore, there appears to be opportunity to initiate stomatal development at an earlier stage so that a higher number of active stomatal complexes are developed once the silique is exposed to light. If so, this may result from the modulation of AG function caused by divergence of the *cis* regulatory elements AG binds to. Assessing the correlation between stomatal development and fertilization in other species will partially address these questions.

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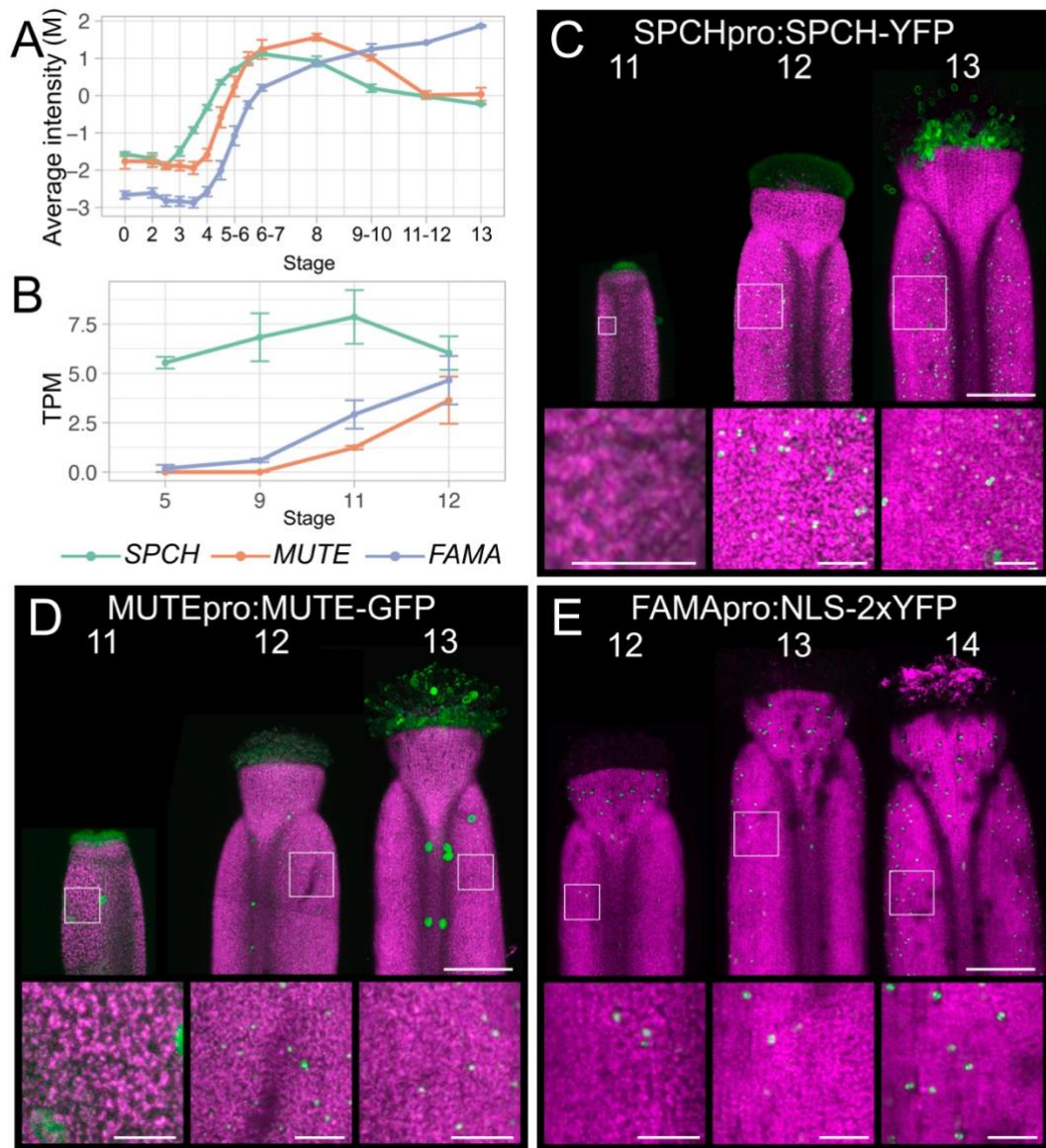
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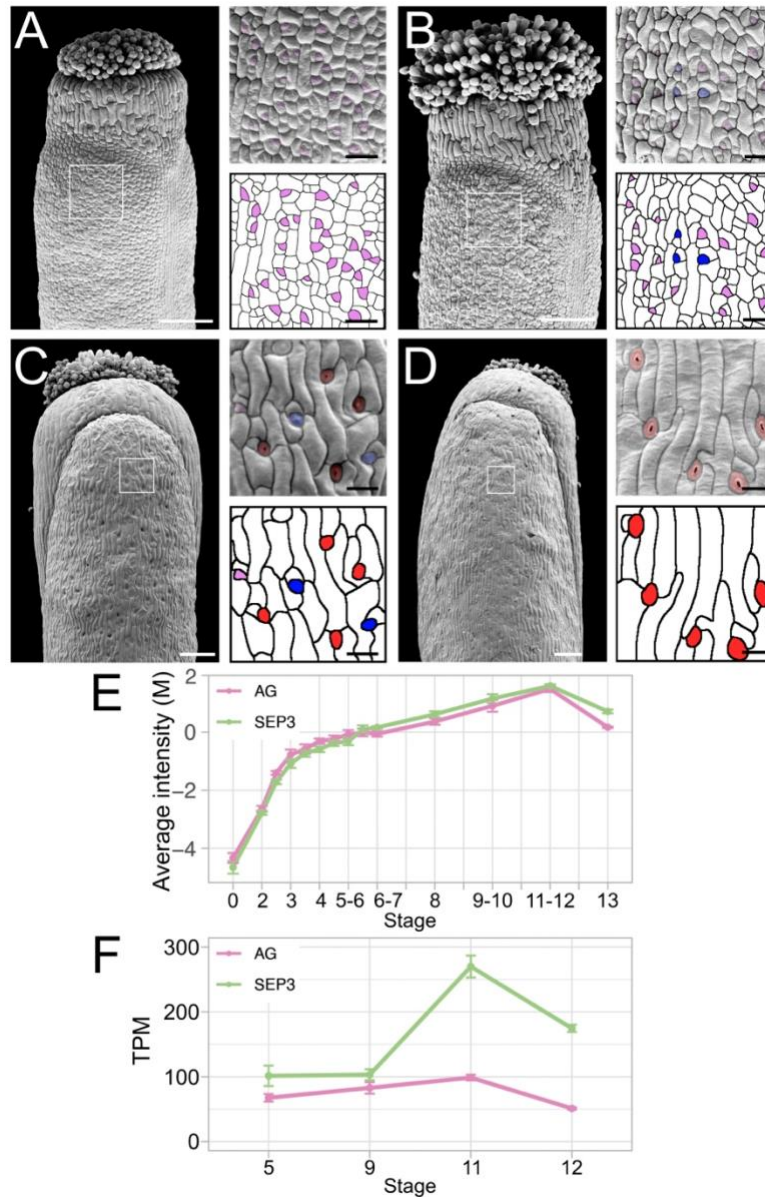
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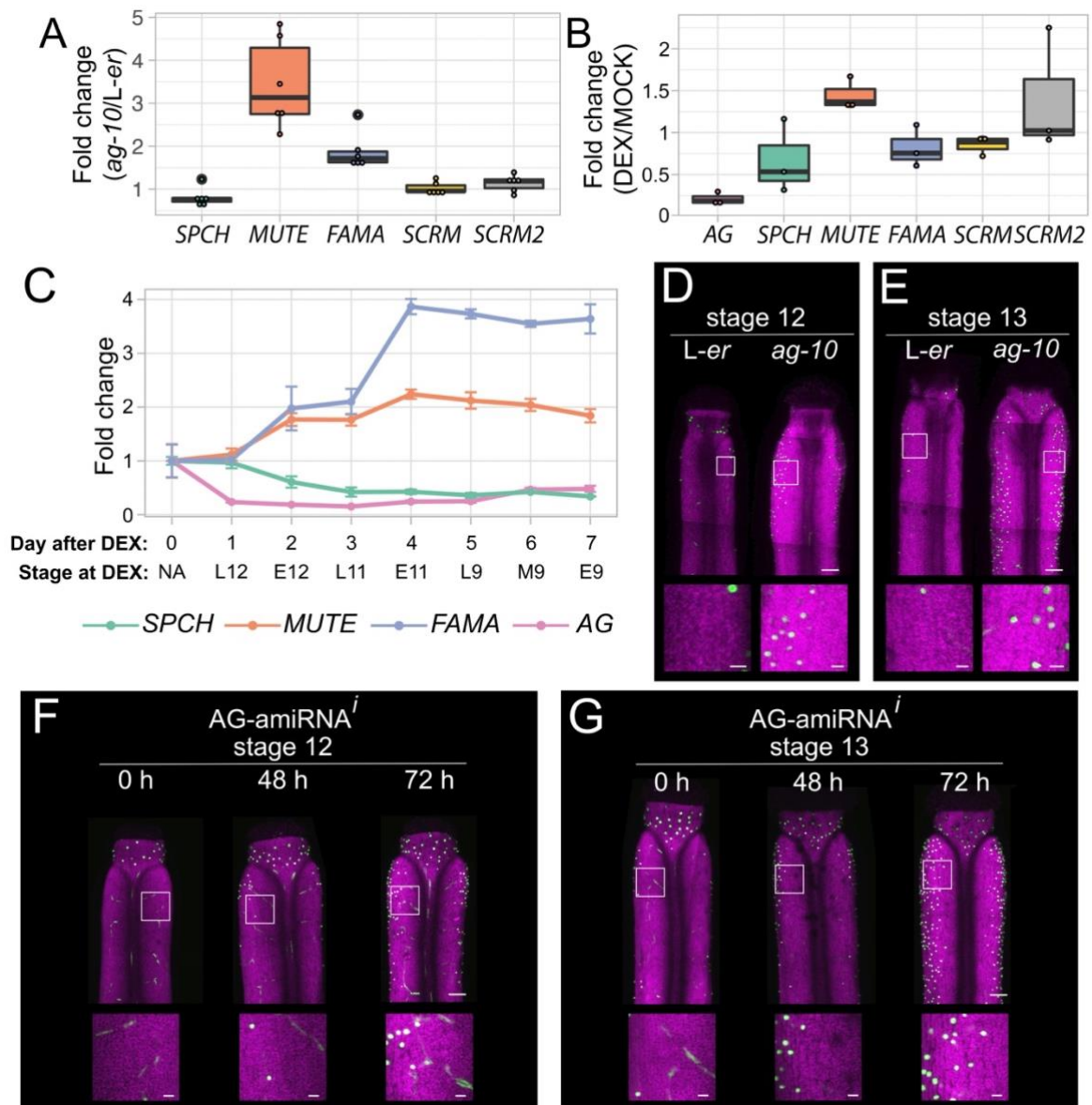


**Figure 1. Expression of stomatal bHLH transcription factors during gynoecium development.** (A-B) Levels of *SPCH*, *MUTE*, and *FAMA* mRNAs at described stages of flower development in (A) whole flower buds (Ryan *et al.* 2015) and (B) laser-microdissected gynoecia (Kivivirta *et al.* 2020). Error bars in (A-B) are s.e.m of three and four biological replicates, respectively. M, log-ratio of time-point to a common reference; TPM, transcripts per million. (C-E) Maximum intensity projections of stitched confocal laser scanning z-stacked micrographs of (C) *SPCHpro:SPCH-YFP*, (D) *MUTEpro:MUTE-GFP*, and (E) *FAMApr:NLS-2xYFP*, transgenes at different stages of gynoecium development as indicated. White boxes indicate areas that were magnified to produce the insets. Fluorescent protein is coloured green and chlorophyll fluorescence is coloured magenta. Scale bars for images of whole gynoecia are 100  $\mu$ m. Scale bars for insets are 20  $\mu$ m.



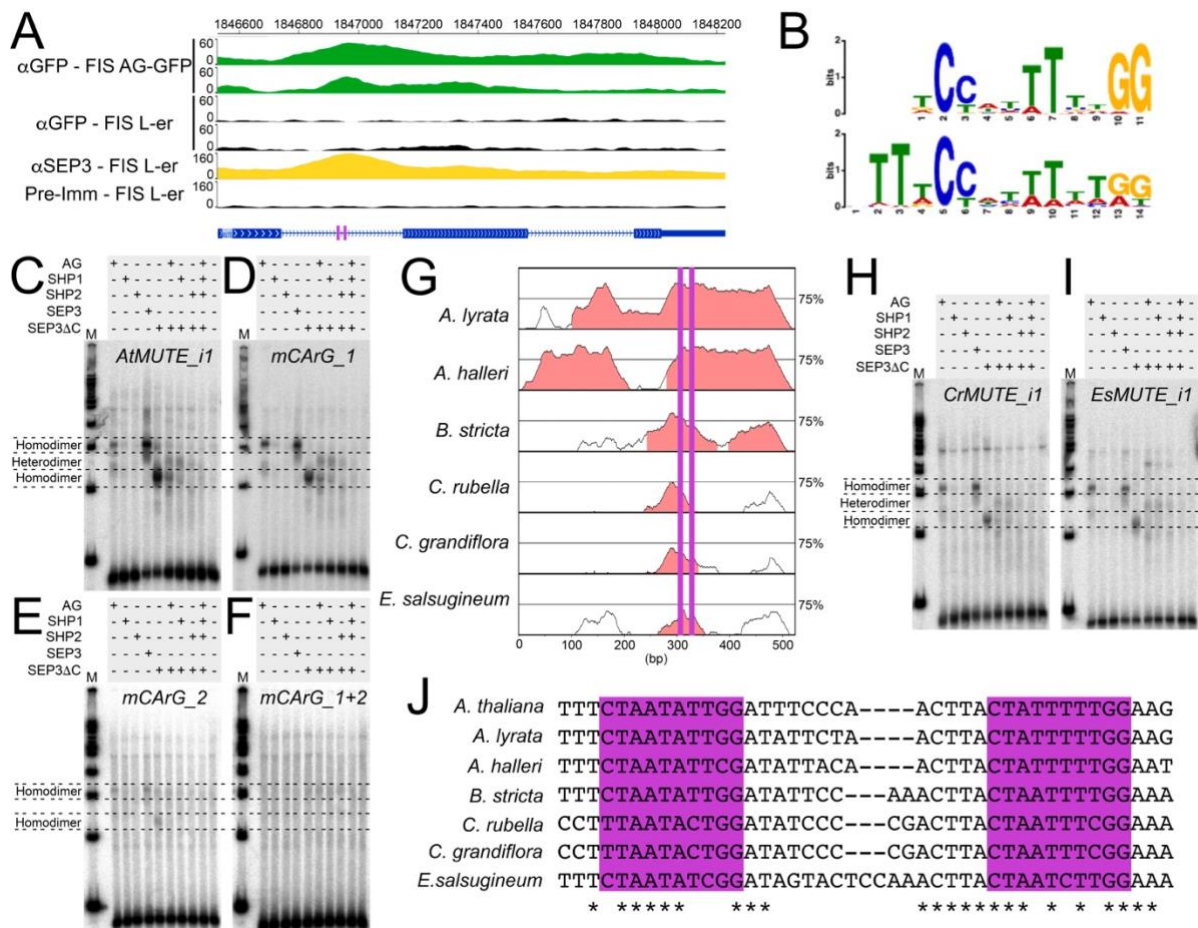


**Figure 2. Progression of stomatal development on gynoecial and silique valves.** (A-D) Scanning electron micrographs of gynoecia at floral stage (A) 12, (B) 13, (C) 15-16 gynoecia and (D) a stage >17 silique with magnifications of the cell surface (from white box). Scale bars for images of whole gynoecia are 100  $\mu$ m. Scale bars for magnifications are 20  $\mu$ m. Purple, blue, and red highlights indicate early, mid, and late stage stomatal lineage morphology, respectively. (E-F) Levels of *AG* and *SEP3* mRNAs over the course of (E) flower development (Ryan *et al.* 2015) and (F) gynoecium development (Kivivirta *et al.* 2020). Error bars in (E-F) are s.e.m of three and four biological replicates, respectively. M, log-ratio of time-point to a reference; TPM, transcripts per million.



**Figure 3. Transcriptional response of master regulators of stomatal development to perturbation of AG.** (A-B) Levels of mRNAs encoding stomatal bHLH transcription factor regulators, as determined by RT-qPCR, in (A) *ag-10* stage 10-13 gynoecia relative to *L-er* stage 10-13 gynoecia, (B) dexamethasone-treated *AG-amiRNA<sup>i</sup>* (*OPpro:AG-amiRNA/35Spro:GR-LhG4*) in stage 10-13 gynoecia relative to mock-treated *AG-amiRNA<sup>i</sup>* in stage 10-13 gynoecia 24 h after treatments. Each dot in (A-B) represents the mean of an individual biological replicate. (C) Levels of *SPCH*, *MUTE*, *FAMA*, and *AG* mRNAs, as determined by RT-qPCR, in stage 13 gynoecia after treatment with dexamethasone relative to untreated (0 d) in stage 13 gynoecia. “Day after DEX” indicates the number of days that gynoecia were treated with DEX before being harvested at anthesis (stage 13), with “0 d” representing the untreated sample. “Stage at DEX” indicates the approximate stage of the flower/gynoecium when DEX treatment was applied. Error bars are s.e.m of three biological replicates. (D-G) Maximum intensity

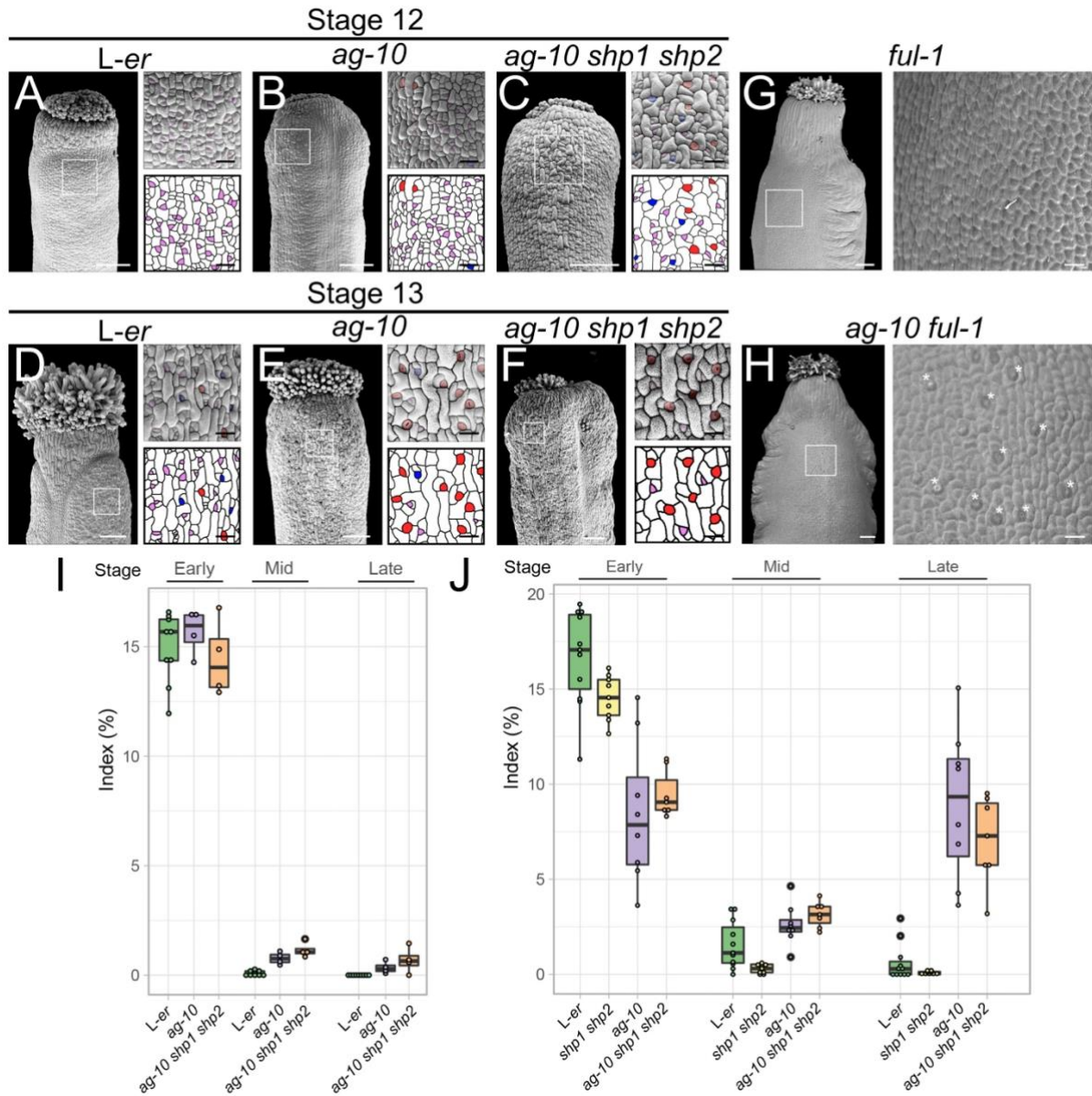
projections of stitched confocal laser scanning z-stack micrographs of (D, F) stage 12 and (E, G) stage 13 gynoecia from plants harbouring a *FAMAPro:2xYFP* transgene in (D-E) *L-er* and *ag-10* backgrounds, and (F-G) the *AG-amiRNA<sup>i</sup>* (*OPpro:AG-amiRNA/35Spro:GR-LhG4*) background before treatment (0 h) and after dexamethasone treatment (48 h and 72 h). YFP is coloured green and chlorophyll fluorescence is coloured magenta. Scale bars for images of whole gynoecia are 100  $\mu\text{m}$ . Scale bars for insets are 20  $\mu\text{m}$ .



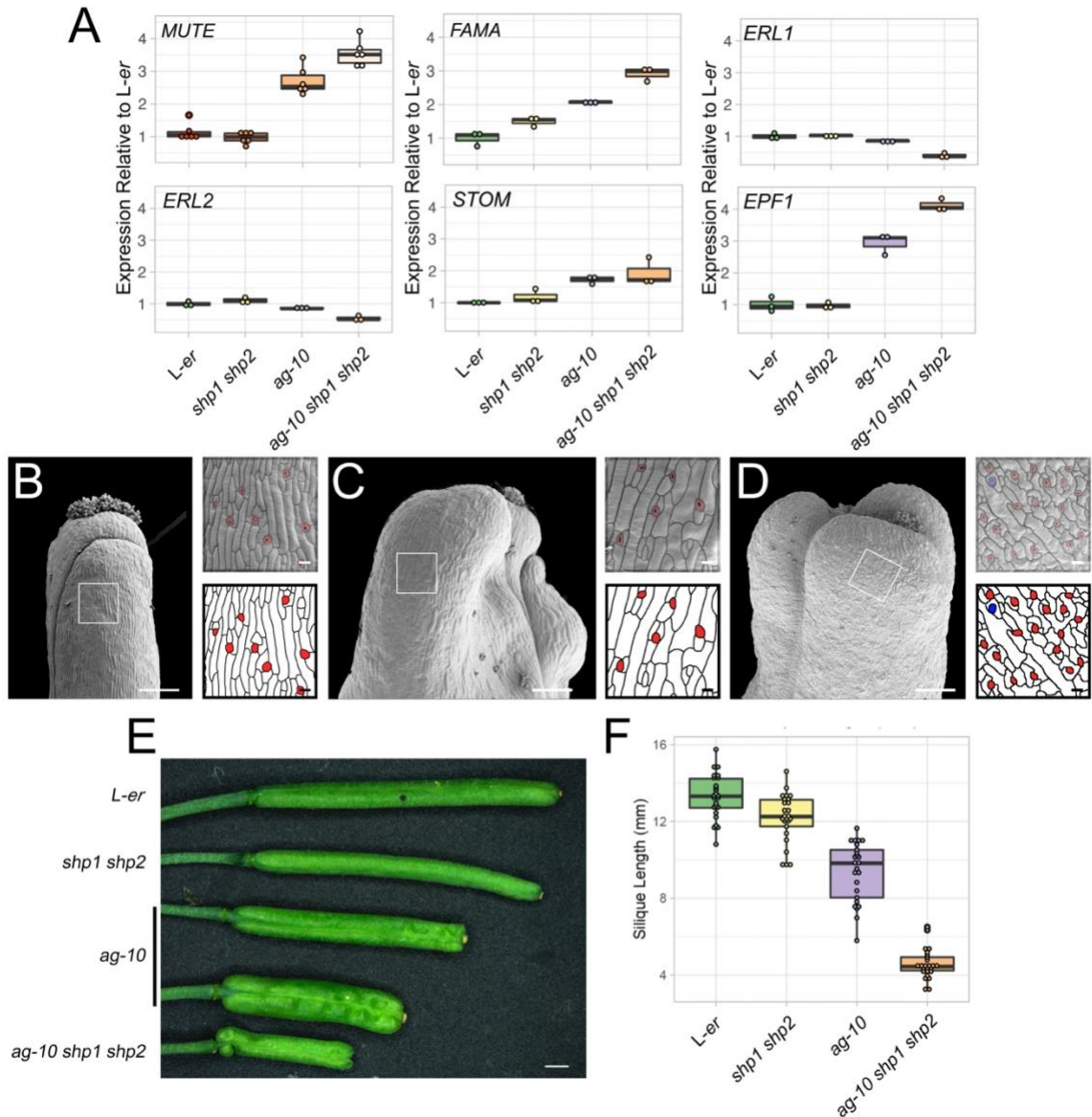
**Figure 4. Interaction between AG, SHP1, SHP2, SEP3, and first intron of MUTE.**

(A) Tracks indicating enrichment of sequences in the first intron of *MUTE* from two replicates of ChIP-Seq of AG-GFP (green tracks) and SEP3 (yellow tracks, combined results of two replicates), proteins in the floral induction system (FIS) (Ó'Maoiléidigh *et al.*, 2013; Pajoro *et al.*, 2014). Corresponding control ChIP-Seq experiments are coloured in black. A schematic of the gene structure of *MUTE* blue is below (exon, rectangles; introns, lines). Two CARG motifs were identified within the first intron of *MUTE* (purple rectangles). (B) Binding logos from MEME and STREME analyses of 1421 binding sites identified in a ChIP-Seq of AG (Ó'Maoiléidigh *et al.*, 2013). (C-F) Protein-DNA gel shift assays using combinations of AG, SEP3, SEP3ΔC, SHP1, and SHP2 recombinant protein and (C) a wild-type *AtMUTE* probe (*AtMUTE\_i1*), (D) a probe where CARG\_1 is mutated (*mCArG\_1*), (E) a probe where CARG\_2 is mutated (*mCArG\_2*), and (F) a probe where both CARG motifs are mutated (*mCArG\_1+2*). M, molecular weight marker. (G) An mVISTA alignment of the first intron of *MUTE* from various members of the Brassicaceae relative to *A. thaliana*. Regions highlighted in salmon have been designated as “Conserved Non-Coding Sequences” by mVISTA. Purple lines indicate the position of each CARG motif. (H-I) Protein-DNA gel shift assays using combinations of AG, SEP3, SEP3ΔC, SHP1, and SHP2 protein and (H) a wild-type *CrMUTE* probe

(*CrMUTE\_il*) and (I) a wild-type *EsMUTE* probe (*EsMUTE\_il*). M, molecular weight marker. (J) Sequence alignments using some of the conserved region identified in (B) containing both CArG motifs. Asterisks indicate conserved nucleotide and the two CArG motifs are highlighted by purple rectangles. Unadjusted images for both replicates of the protein-DNA gel shift assays can be found in [Supplemental Figure 8](#).



**Figure 5. Stomatal development on gynoecial valves in response to AG and SHP perturbation.** (A-F) Scanning electron micrographs of (A-C) stage 12 and (D-F) stage 13 *L-er*, *ag-10*, and *ag-10 shp1 shp2* gynoecia, as indicated. Purple, blue, and red highlights indicate early, mid, and late-stage stomatal lineage morphology, respectively. (G-H) Scanning electron micrographs of a (G) *ful-1* and (H) *ag-10 ful-1* silique. Asterisks indicate presence of stomata. Scale bars for images of whole gynoecia are 100  $\mu$ m. Scale bars for magnifications are 20  $\mu$ m. (I-J) Index of early, mid, and late stomatal lineages based on morphology from scanning electron micrographs from (I) stage 12 and (J) stage 13 *L-er*, *shp1 shp2*, *ag-10*, *ag-10 shp1 shp2* gynoecia. Each dot represents an individual sample.



**Figure 6. Redundancy between AG and SHP1/2 during gynoecium and silique development.** (A) Levels of *MUTE*, *FAMA*, *ERL1*, *ERL2*, *STOM*, and *EPF1* mRNAs as determined by RT-qPCR, in *L-er*, *shp1 shp2*, *ag-10*, *ag-10 shp1 shp2* stage 10-13 gynoecia relative to average of *L-er* samples. Each dot represents an individual independent biological replicate. (B-D) Scanning electron micrographs of mature siliques of (B) *L-er*, (C) *ag-10*, and (D) *ag-10 shp1 shp2*. Purple, blue, and red highlights indicate early, mid, and late-stage stomatal lineage morphology. Scale bars for images of whole gynoecia are 200  $\mu$ m. Scale bars for magnifications are 20  $\mu$ m. (E) Mature siliques of *L-er*, *shp1 shp2*, *ag-10*, *ag-10 shp1 shp2*. Scale is 1 mm. (F) Length of *L-er*, *shp1 shp2*, *ag-10*, *ag-10 shp1 shp2* mature siliques.