

1 **Control of patterning, growth and differentiation by floral organ identity**
2 **genes**

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Short statement

MADS domain proteins determine floral organ identity; recent work has given insight into how these proteins modify organ development by interacting with genes involved in organ patterning, growth and differentiation.

Abstract

In spite of the different morphologies of sepal, petals, stamen and carpels, all these floral organs are believed to be modified versions of a ground-state organ similar to the leaf. Modifications of the ground-state developmental program are orchestrated by different combinations of MADS-domain transcription factors encoded by floral organ identity genes. In recent years, much has been revealed about the gene regulatory networks controlled by the floral organ identity genes and about the genetic pathways that control leaf development. Here, I review how floral organ identity is connected with the control of morphogenesis and differentiation of shoot organs, focusing on the model species *Arabidopsis thaliana*. Direct links have emerged between floral organ identity genes and genes involved in abaxial-adaxial patterning, organ boundary formation, tissue growth and cell differentiation. In parallel, predictive models have been developed to explain how the activity of regulatory genes can be coordinated by intercellular signaling and constrained by tissue mechanics. Combined, these advances provide a unique opportunity to reveal how exactly leaf-like organs have been “metamorphosed” into floral organs during evolution and to reveal crucial regulatory points in the generation of plant form.

1 **Introduction**

2

3 Over two decades ago, molecular genetics of floral development was in its
4 heyday. Work in *Arabidopsis* and snapdragon had converged on the well-
5 known ABC model, which explained how each type of floral organ is
6 specified by a different combination of floral organ identity genes, which are
7 expressed in overlapping regions of the flower (Bowman *et al.*, 1991; Coen
8 and Meyerowitz, 1991; Schwarz-Sommer *et al.*, 1990). Within a few years, all
9 floral organ identity genes had been cloned, and all but one turned out to
10 encode transcription factors containing the MADS DNA binding domain
11 (named after yeast MCM1, *Arabidopsis* AGAMOUS, snapdragon DEFICIENS
12 and mammalian Serum Response Factor)(Sommer *et al.*, 1990; Yanofsky *et*
13 *al.*, 1990). Similar combinations of homologous genes encoding MADS-
14 domain proteins were found to determine floral organ identity across distant
15 species, including monocotyledons (Bowman *et al.*, 2012; Irish and Litt,
16 2005; Ito, 2011; Wellmer *et al.*, 2014).

17

18 Subsequently, the genetic interactions between *MADS* organ identity genes
19 were neatly mirrored by protein-protein interactions in what became known
20 as the quartet model (Theißen and Saedler, 2001). The MADS-domain
21 proteins required for each type of organ directly interact with each other to
22 form different multimeric complexes (Melzer and Theissen, 2009), which are
23 sufficient to convert any type of shoot organ into a specific floral organ
24 (Honma and Goto, 2001; Pelaz *et al.*, 2000; Pelaz *et al.*, 2001). In
25 *Arabidopsis*, the following combinations of MADS-domain proteins specify
26 each floral organ type: APETALA1 (AP1) and SEPALLATA (SEP) proteins
27 (SEP1, 2, 3 and 4) direct sepal development; petals are specified by AP1
28 and SEP1-3 together with APETALA3 (AP3) and PISTILLATA (PI); AP3 and
29 PI combined with SEP1-3 and AGAMOUS (AG) direct stamen development,
30 and AG combined with SEP1-3 specifies carpels (Figure 1; reviewed by
31 (Wellmer *et al.*, 2014)).

32

33 The findings that any shoot organ can be converted to a floral organ, and that
34 in the absence of organ identity genes, floral organs become leaf-like

1 (Bowman *et al.*, 1991; Ditta *et al.*, 2004) matched the idea proposed by
2 Goethe in the 18th century, that floral organs are modified versions of a leaf-
3 like archetypal organ (Goethe, 1790; Pelaz *et al.*, 2001). Therefore, the
4 diverse morphology of each type of floral organ would be expected to arise
5 from modifications of a basic, leaf-like developmental program. It would also
6 be expected that genes targeted by the floral homeotic genes would reveal
7 key control points where morphological diversity can be generated. Until
8 recent years, however, not enough was known about the genes that control
9 leaf and floral organ growth to suggest what aspects of the basic, leaf-like
10 program would be modified by organ identity genes.

11

12 In the last few years, much has been learned about the gene regulatory
13 network controlled by floral homeotic genes and about the mechanisms that
14 control growth and morphogenesis of shoot organs. As reviewed below, links
15 have emerged between floral homeotic genes and general regulators of
16 lateral organ growth, including molecular links to the cellular activities that
17 support tissue growth and shape organs (cell division, cell wall functions).
18 More recently, molecular work and computer models have started to
19 converge to explain how the control of organ growth unfolds from the
20 molecular to cellular to organ scale. This creates new opportunities to reveal
21 key regulatory points in the generation of morphological diversity between
22 organs in the same plant and potentially between the same organs across
23 plant species.

24

25 **MADS-domain organ identity proteins orchestrate gene expression** 26 **throughout floral organ development**

27

28 The finding that all floral homeotic genes encode transcription factors
29 prompted numerous studies of the changes in gene expression downstream
30 of the floral organ identity genes (reviewed in (Wellmer *et al.*, 2014)). Initial
31 comparisons of organ identity mutants and wild-type revealed large numbers
32 of changes in gene expression, most of which are likely to be indirectly
33 caused by the organ identity genes. Subsequent studies using inducible
34 versions of the MADS-domain proteins revealed immediate target genes and

1 were extended more recently by Chromatin Immunoprecipitation – High
2 throughput Sequencing (ChIP-Seq) to identify genome-wide binding sites of
3 the MADS-domain protein complexes. Comprehensive stage-specific
4 expression and ChIP-Seq data are now available for all classes of organ
5 identity genes in Arabidopsis: AP1 (Kaufmann *et al.*, 2010; Pajoro *et al.*,
6 2014), AP3/PI (Wuest *et al.*, 2012), AG (Ó'Maoiléidigh *et al.*, 2013) and
7 SEP3 (Kaufmann *et al.*, 2009; Pajoro *et al.*, 2014).

8

9 These studies revealed that MADS-domain organ identity proteins directly
10 interact with thousands of loci (between 1500 high-confidence target sites for
11 AP3/PI and more than 4000 for SEP3). In part, the large number of
12 downstream targets reflects the fact that, as indicated by early work using
13 temperature-sensitive alleles, organ identity genes are required at all stages
14 of organ development (Bowman *et al.*, 1989), and as shown by time-course
15 transcriptome analysis, control distinct sets of genes at different stages of
16 floral development (Gómez-Mena *et al.*, 2005; Wellmer *et al.*, 2006). In
17 contrast to the detailed analysis of temporal changes in gene expression,
18 much less is known about cell type-specific target genes. It is possible that
19 even in a single developmental stage, organ identity genes will control
20 different genes in specific tissues and regions of the organ. Thus the gene
21 expression programs directed by organ identity genes in individual cells
22 might be less complex than our current picture based on whole developing
23 buds.

24

25 Context-specific interactions with target genes are also suggested by
26 comparing ChIP-Seq and expression data. In the case of SEP3, 72% of the
27 bound genes were differentially expressed at some point in flower
28 development or in at least one of the floral homeotic mutants, suggesting that
29 the majority of SEP3 binding sites are functionally relevant (Kaufmann *et al.*,
30 2009). However, this does not imply that every binding event causes a
31 transcriptional response. This has been shown clearly for AP1 and AG, for
32 which only about 10% of genes bound during early floral development also
33 showed differential expression in the same experimental conditions
34 (Kaufmann *et al.*, 2009; Ó'Maoiléidigh *et al.*, 2013). The discrepancy between

1 binding and expression differences suggests that many of the binding sites
2 identified by ChIP-Seq may be functionally irrelevant. Alternatively, MADS-
3 domain proteins could “tag” genes that are due to be regulated at some point
4 or in some cell type during floral organ development, but only be able to
5 change their activity when co-factors become available. This has been
6 confirmed in the case of AG: a significant number of loci bound by AG early
7 in development only showed AG-dependent transcriptional changes at later
8 developmental stages (O’Maoileidigh 2013).

9

10 One mechanism by which MADS-domain proteins could prime target genes
11 for subsequent regulation by other factors could be by inducing changes in
12 chromatin accessibility. This idea has been supported by careful comparison
13 between binding of AP1 and SEP3 and genome-wide changes in DNase I
14 sensitive sites (Pajoro *et al.*, 2014), and by the direct interaction between
15 MADS-domain proteins and chromatin-modifying enzymes (Smaczniak *et al.*,
16 2012; Sridhar *et al.*, 2006). In addition to chromatin regulators, MADS-
17 domain proteins interact with several other transcription factors, such as
18 BELL-like homeodomain and AUXIN RESPONSE FACTOR (ARF) proteins
19 (Smaczniak *et al.*, 2012). These transcription factors control patterning and
20 growth of both leaves and floral organs (see below), so direct interaction with
21 general regulators of organ development appears to be one of the
22 mechanisms by which MADS-domain proteins modify the basal leaf-like
23 developmental program. The ubiquitous, but context-specific function of
24 organ identity proteins, combined with their direct interaction with core
25 regulators of shoot organ development, support the idea that MADS domain
26 proteins function as organ identity co-factors that modify the function of a
27 variety of transcription factors with more specialized functions (Sablowski,
28 2010), as proposed for Hox proteins in *Drosophila* (Akam, 1998).

29

30 **Interaction with genetic pathways for organ patterning**

31

32 The sets of target genes in early organ development are especially enriched
33 in genes that encode additional transcription factors (Gómez-Mena *et al.*,
34 2005; Kaufmann *et al.*, 2010; Ó’Maoiléidigh *et al.*, 2013; Wuest *et al.*, 2012).

1 Relevant to morphogenesis, these transcription factors provide multiple links
2 between the organ identity genes and regulatory networks that control
3 adaxial-abaxial patterning, formation of organ boundaries and development
4 of the organ margins.
5
6 One of the earliest acting patterning networks establishes the differences
7 between the adaxial (facing the meristem) and the abaxial (facing away from
8 the meristem) sides of the organ. The initial clue that distinguishes the
9 adaxial and abaxial sides of the primordium is likely derived from the radial
10 axis of the shoot apex (meristem in the center, initiating organs in the
11 periphery) (Emery *et al.*, 2003; McConnell *et al.*, 2001). The different
12 identities of the two sides are consolidated and maintained by the
13 antagonistic activity of adaxial (*AS1*, *AS2*, *HD-ZIPIII* genes) and abaxial
14 identity genes (*YABBY*, *KANADI* and *ETT/ARF4* genes)(reviewed in (Khan *et al.*,
15 2014; Rodriguez *et al.*, 2014)). These genes perform comparable
16 functions during leaf and floral organ development, but there is some
17 specialization of family members. For example, mutation of the *YABBY* gene
18 *FILAMENTOUS FLOWER (FIL)* is sufficient to cause severe abaxial-adaxial
19 polarity defects in floral organs (Sawa *et al.*, 1999), whereas in leaves there
20 is a higher level of redundancy between *YABBY* genes, and comparable
21 defects are only seen in the triple mutant *fil yab3 yab5* (Stahle *et al.*, 2009).
22 Other *YABBY* genes function in abaxial-adaxial patterning specifically in
23 flowers: *CRABS CLAW (CRC)* in carpels (Bowman and Smyth, 1999) and
24 *INNER NO OUTER (INO)* in ovules (Villanueva *et al.*, 1999). Organ identity
25 genes interact directly with abaxial-adaxial polarity genes, for example,
26 *AP1/SEP3* bind to *FIL*, *AS1* and *AS2* (Pajoro *et al.*, 2014) and *CRC* is directly
27 activated by *AG* (Gómez-Mena *et al.*, 2005). These interactions may have a
28 role in floral-specific variations in adaxial-abaxial patterning, but it is not clear
29 yet what role this may play in morphological differences between leaves and
30 floral organs.
31
32 The boundary between adaxial and abaxial regions of the organ primordium
33 is important for establishing domains at the organ margins, which promote
34 lateral growth (Eshed *et al.*, 2004; Waites and Hudson, 1995) to produce the

1 planar structures of leaves, petals, sepals and carpel walls. In leaf
2 development, these marginal regions can retain the activity of a subset of
3 genes that control meristem function: the homeodomain-encoding *KNOX*
4 family, which includes the meristem maintenance genes *SHOOT*
5 *MERISTEMLESS (STM)* and *BREVIPEDICELLUS (BP)* (Hay and Tsiantis,
6 2010), and *CUP-SHAPED COTYLEDON (CUC) 1* and *2*, which are initially
7 required for the establishment of *KNOX* expression but subsequently repress
8 *KNOX* genes to establish the lateral boundaries of shoot organs (Aida and
9 Tasaka, 2006). This meristematic “module” also functions in the leaf margins
10 to control the formation of leaf lobes and leaflets, and has been repeatedly
11 involved in the independent evolution of compound leaves in different clades
12 (Blein *et al.*, 2008; Townsley and Sinha, 2012). In extreme cases, such as in
13 *Kalanchoe*, *KNOX* gene expression in the sinuses of serrations maintain
14 meristematic regions that generate new plants (Garcês *et al.*, 2007).

15
16 In the gynoecium, organ margins also have an organogenic role. The
17 gynoecium is likely derived from leaf-like organs that fused at their margins
18 (Hawkins and Liu, 2014). The similarity between each of the fused units
19 (carpels) and leaves is readily apparent in homeotic mutations such as *ap2-*
20 *2*, which replace sepals by single carpels with ovules on their margins
21 (Bowman *et al.*, 1989). The region of the gynoecium corresponding to the
22 fused carpel margins is called the carpel margin meristem (CMM), which
23 produces the inner structures of the gynoecium, including the placenta,
24 ovules, septum and transmitting tract (Hawkins and Liu, 2014). Numerous
25 mutations affect carpel fusion and development of the CMM, many of which
26 affect flower-specific regulatory genes such as *CRC*, *SPATULA (SPL)*,
27 *ALCATRAZ (ALC)* and *INDEHISCENT (IND)* (Reyes-Olalde *et al.*, 2013).
28 Thus CMM development appears to be a particularly specialized aspect of
29 the gene expression program downstream of the organ identity genes.
30 However, there are also aspects shared with leaf margin development, in
31 particular the central role of the meristematic module including *CUC* and
32 *KNOX* genes (Hasson *et al.*, 2011) (Kamiuchi *et al.*, 2014). Organ identity
33 proteins directly interact with *CUC* genes (Kaufmann *et al.*, 2009;
34 Ó'Maoiléidigh *et al.*, 2013; Wuest *et al.*, 2012), but it is not known whether

1 this interaction is involved in elaborating the function of *CUC* genes in
2 marginal tissues, such as the carpel CMM.
3
4 *CUC* genes are not the only organ boundary genes that modify organ shape.
5 Development of the basal region of shoot organs is controlled by a different
6 set of organ boundary genes, notably *BLADE-ON-PETIOLE (BOP) 1* and *2*
7 (Ha *et al.*, 2003; Hepworth *et al.*, 2005; Khan *et al.*, 2014; Norberg *et al.*,
8 2005), and *ARABIDOPSIS THALIANA HOMEODOMAIN 1 (ATH1)* (Gómez-Mena
9 and Sablowski, 2008). *BOP1/2* are required for proper development of the
10 leaf petiole, preventing outgrowth of the leaf lamina at least in part by
11 regulating adaxial-abaxial polarity genes and antagonizing *KNOX* genes (Ha
12 *et al.*, 2007; Jun *et al.*, 2010). Both *BOP1/2* and *ATH1* are also required for
13 the development of basal organ structures, such as the abscission zone. The
14 direct interaction of organ identity proteins with *BOP1*, *BOP2* and *ATH1*
15 (Ó'Maoiléidigh *et al.*, 2013; Wuest *et al.*, 2012) may play a role in generating
16 the diverse basal structures of floral organs: sepals lack recognizable
17 petioles, carpels normally have very short petiole-like structures (the
18 gynophores), petals have petioles comparable to those of leaves, and the
19 petiole-like structures of stamens (the filaments) are very enlarged compared
20 to the other floral organs. It must be noted, however, that *BOP* genes also
21 interact genetically with *AP1* at the transition from inflorescence meristem to
22 floral meristem (Xu *et al.*, 2010), so the interaction with *AP1/SEP3* may
23 reflect functions that precede floral organ development.
24
25 The development of distinct tissues along the apical-basal axis patterning
26 has also been linked to auxin function, particularly in carpel development. It
27 was initially proposed that an auxin gradient patterns the gynoecium, but
28 more recent evidence supports a model in which the apical-basal defects
29 seen in auxin-related mutants result from growth defects very early in
30 primordium development (Hawkins and Liu, 2014). There is evidence that
31 input from organ identity genes is important for this role of auxin in carpel
32 development: *SEP3* binds to genes involved in auxin transport and auxin
33 responses (*PIN-LIKE 4*, *PINOID*, *ARF3*, *ARF8*, *IAA4*), and the *sep1 sep2*
34 *sep3* triple mutant has elongated gynophores similar to those seen in the *pid*

1 mutant (Kaufmann *et al.*, 2009). Furthermore, plants in which SEP3 was
2 converted from a transcriptional activator to a repressor by fusion to the EAR
3 (ERF-associated Amphiphilic Repression) domain showed severe defects in
4 floral organ development, including defects in apical-basal development of
5 carpels, similar to those of the auxin-related mutants *pin1* and *arf3* or of
6 plants treated with the auxin transport inhibitor NPA (Kaufmann *et al.*, 2009).
7 Thus organ identity complexes are likely to influence apical-basal patterning
8 through direct regulation of genes involved in auxin transport and signaling.

9
10 Given the extensive use of hormone signaling in all aspects of plant
11 development, it is not surprising that in addition to the auxin-related genes
12 mentioned above, organ identity genes have many direct links to hormone
13 synthesis and signaling. These include regulation by AG of jasmonic acid
14 synthesis (Ito *et al.*, 2007), which is essential for anther development (Ito *et al.*,
15 2007) and direct regulation of genes involved in GA biosynthesis
16 (GA2ox1) and response (RGL2) by multiple organ identity genes (Gómez-
17 Mena *et al.*, 2005; Kaufmann *et al.*, 2010), although in the later case the
18 specific consequences for floral organ development are not clear.

19

20 **Interaction with growth regulatory genes**

21

22 Ultimately, organ identity genes alter organ shape by controlling rates and
23 directions of tissue growth (Coen *et al.*, 2004). This role likely involves
24 interactions with intermediate regulatory genes that control growth of both
25 vegetative and floral organs.

26

27 One of the best-characterized genetic pathways that control organ growth is
28 centered on the *GRF* (*GROWTH REGULATING FACTOR*) genes (Rodriguez
29 *et al.*, 2014). GRFs are a family of transcription factors that promote cell
30 proliferation during lateral organ development (Kim *et al.*, 2003; Rodriguez *et al.*,
31 2010), in association with the co-activator GRF-INTERACTING FACTOR
32 (GIF), also called *ANGUSTIFOLIA 3* (*AN3*) (Horiguchi *et al.*, 2005; Kim and
33 Kende, 2004). In *Arabidopsis*, *GRFs* are antagonized by the micro RNA
34 miR396, which targets seven of the nine family members for degradation,

1 and is in turn activated by transcription factors of the TCP family
2 (TEOSINTE-BRANCHED 1, CYCLOIDEA and PROLIFERATING CELL
3 FACTORS 1 and 2) (Rodriguez *et al.*, 2010). Both *TCPs* and *GRFs* are
4 overrepresented among the targets of organ identity genes (Kaufmann *et al.*,
5 2009), and SEP3 binds to all 9 *GRF* genes (Pajoro *et al.*, 2014). DNA
6 sequences bound by TCP proteins are enriched in the vicinity of genomic
7 binding sites for SEP3, suggesting that organ identity proteins and TCPs
8 could influence the regulation of at least a subset of TCP target genes
9 (Kaufmann *et al.*, 2009). The combined data make the TCP/GRF pathway an
10 obvious candidate to mediate the effects of organ identity genes on organ
11 growth.

12
13 Another transcription factor with well-studied roles in organ growth is
14 JAGGED (*JAG*). Mutations in *JAG* and in its paralog, *NUBBIN* (*NUB*)
15 enhance leaf serration and impair growth of the apical region of floral organs
16 (Dinneny *et al.*, 2006; Dinneny *et al.*, 2004; Ohno *et al.*, 2004). The
17 preferential role of *JAG* in the distal region of floral organs led to the
18 suggestion that it functions as a mediator between organ patterning and
19 growth processes (Breuninger and Lenhard, 2010). This idea has been
20 corroborated by the finding that *JAG* directly binds to genes involved in
21 boundary formation (e.g. *BOP1*, *BOP2*) and organ growth (*TCP4*, *GRF5*,
22 *AN3* and *miRNA396*), in addition to directly regulating genes involved in the
23 cellular activities required for tissue growth, such as cell cycle control and cell
24 wall functions (Schiessl *et al.*, 2014). Quantitative analysis of the effects of
25 *JAG* at the cellular level revealed roles in both the rate of cell growth and
26 proliferation, and in promoting oriented cell expansion (Schiessl *et al.*, 2012),
27 and genetic analysis confirmed that *JAG* stimulates organ growth to a large
28 extent by repressing the expression of cell cycle inhibitors (Schiessl *et al.*,
29 2014). *JAG* and *NUB* are direct target genes of AG, SEP3 and AP3/PI
30 (Gómez-Mena *et al.*, 2005; Kaufmann *et al.*, 2009; Ó'Maoiléidigh *et al.*, 2013;
31 Wuest *et al.*, 2012), so *JAG/NUB* are also good candidates to mediate
32 between organ identity and growth.

33

1 As with most of the examples discussed above, direct targets of organ
2 identity proteins reveal molecular links to processes such as organ growth.
3 However, exactly how the temporal or spatial expression pattern of these
4 genes is altered by organ identity genes, and how these changes are
5 translated into the patterns of cell proliferation and expansion that shape
6 organs, remains virtually unknown. Some insight into how *JAG* may shape
7 different organ types came from recent computational models of organ
8 growth (Sauret-Güeto *et al.*, 2013). The model had three main components:
9 i) a polarity factor whose distribution and orientation in the tissues depends
10 on the location of proximal and distal organizers; ii) growth factors that
11 determine rates of growth perpendicular and parallel to local polarity; iii)
12 mechanical connectedness leads to accumulation of stresses during growth,
13 which are resolved in part by tissue deformation. Using this modeling
14 framework, the different shapes of petals and leaves were simulated by
15 assuming different patterns of tissue polarity (convergent at the distal end in
16 leaves, divergent in petals). Considering its expression pattern and mutant
17 phenotype, *JAG* was proposed as a candidate for the distal growth factor,
18 which would preferentially promote growth perpendicular to local polarity.
19 Based on the effect of *jag* mutation on reporters for auxin transport and
20 response (which are connected to local tissue polarity), *JAG* was also
21 proposed to be required for establishing a continuous distal organizer along
22 the edge of the petal. Changes in the corresponding parameters resulted in
23 models that correctly captured the main features of *jag* petals (narrow organs
24 with serrated edges, Figure 2). Therefore this type of modeling approach has
25 the potential to reveal how organ identity genes generate different organ
26 morphologies through changes in tissue polarity and the localized activity of
27 growth regulators.

28

29 **Interaction with cellular differentiation pathways**

30

31 As the organ grows and takes shape, cell differentiation is initiated. Floral
32 organs differ from leaves not only in morphology, but also in their repertoire
33 of cell types. Accordingly, organ identity genes directly interact with genes

1 that control cell identity, both to repress leaf-specific cell types and to
2 promote floral-specific differentiation.
3
4 Photosynthetic capacity is a prominent feature of leaves that is lost in petals
5 and stamens. Presumably B-function genes suppress the differentiation of
6 photosynthetic tissues, but the molecular basis for this is only partially
7 understood. One of the few direct targets of organ identity genes specifically
8 regulated in petals is the *BANQUO3* (*BNQ3*) gene, which encodes an
9 atypical bHLH protein that does not have a DNA binding domain but is
10 believed to interact with other bHLH transcription factors to modify their
11 function (Mara *et al.*, 2010). *BNQ3* is widely expressed in the shoot but is
12 directly repressed by AP3/PI in developing petals. Loss of *BNQ3* function
13 caused reduced chlorophyll levels in cauline leaves, stems, sepals and
14 carpels, while *BNQ3* overexpression interfered with light-induced hypocotyl
15 elongation. Thus one way in which organ identity genes turn green leaves
16 into pale petals is by interfering with light signaling and chloroplast
17 development through repression of *BNQ3*.

18
19 An example of a leaf cell type whose development is suppressed during floral
20 development are the branched trichomes, whose development is promoted
21 by *GLABROUS1* (*GL1*) (Larkin *et al.*, 1994) and inhibited by *CAPRICE*
22 (*CPC*) (Schellmann *et al.*, 2002). AP1, AP3, PI and AG all directly bind to the
23 *GL1* and *CPC* loci, and consistent with the absence of trichomes on stamens
24 and carpels, AG repressed *GL1* and activated *CPC* (Ó'Maoiléidigh *et al.*,
25 2013). Further support for the role of AG in repressing this aspect of the leaf
26 development program came from experiments in which loss of AG function
27 was caused during development by artificial miRNAs. Loss of AG function
28 during mid-stages of organ development, when cell differentiation is
29 underway, caused ectopic trichome development on carpels (Ó'Maoiléidigh
30 *et al.*, 2013).

31
32 There are also examples of differentiation pathways that are unique to floral
33 organs and are directly activated by the organ identity genes.
34 *SPOROCTELESS* (*SPL*), also known as *NOZZLE* (*NZZ*), is required for

1 development of sporogenic tissues, which produce the male and female
2 gametophytes (Schiefthaler *et al.*, 1999; Yang *et al.*, 1999). AG directly binds
3 to and activates *SPL* (Ito *et al.*, 2004; Ó'Maoiléidigh *et al.*, 2013) and ectopic
4 activation of *SPL* is sufficient to activate pollen development in petals,
5 revealing that male sporogenesis is a developmental module invoked by AG
6 through *SPL* (Ito *et al.*, 2004). However, competence to respond to ectopic
7 *SPL* was only seen in the inner organs of the *ag-1* mutant and was limited to
8 the distal petal margins, showing that the exact timing and location of *SPL*
9 function depend on additional, unidentified floral factors.

10

11 Another differentiation pathway that is specific to flowers leads to the
12 formation of conical cells on the petal epidermis, which have characteristic
13 cuticular wax ridges. These conical cells are a conserved feature of petals
14 that has been implicated in the interaction with pollinators (Glover and Martin,
15 1998). *SHINE1*, which is directly bound by AP1 during petal development
16 (Pajoro *et al.*, 2014), coordinates the expression of biosynthetic genes
17 required for the production of the cuticular ridges (Shi *et al.*, 2011). The latter
18 example illustrates how organ identity genes direct the gene expression
19 program up to the finishing touches in floral organ development.

20

21 **Conclusions and perspectives**

22

23 We now have a much better understanding of the genetic networks that
24 control leaf development, and numerous direct molecular links between floral
25 organ identity genes and key nodes in these networks. Rather than
26 functioning as master genes at the top of a regulatory hierarchy that
27 overrides the leaf developmental program, MADS-domain proteins directly
28 modify every step of organ development, from early patterning to growth to
29 final differentiation (Figure 3). The current picture suggests that the
30 interaction between organ identity genes and general regulators of organ
31 development may produce the overall structure of floral organs, on which
32 organ-specific cell types and structures are added or suppressed by
33 interaction with more specialized gene expression programs.

34

1 One important next step will be to test how floral organ identity genes modify
2 organ morphology through changes in the temporal or spatial expression
3 patterns of general regulators of shoot development. To achieve this, at least
4 three challenges lie ahead. First, we will need higher resolution, quantitative
5 measurements of gene expression during organ development. An example of
6 how this type of data can be integrated into three-dimensional models of
7 floral buds has been produced for early sepal development (La Rota *et al.*,
8 2011). Second, it will be necessary to reveal the links between the relevant
9 regulatory genes and the cellular activities that constrain tissue growth (e.g.
10 oriented cell expansion, cell cycle progression) (Schiessl *et al.*, 2014). Third,
11 spatial modeling will be required to simulate and predict the feedbacks
12 between gene expression, growth and tissue mechanics. Progress has been
13 made in establishing predictive models of leaf and floral organ growth
14 (Robinson *et al.*, 2011; Sauret-Güeto *et al.*, 2013), and in understanding the
15 feedbacks between tissue mechanics and growth (Kierzkowski *et al.*, 2012;
16 Routier-Kierzkowska and Smith, 2013). In years to come, these approaches
17 may finally give a full understanding of how exactly shoot organs can be
18 “metamorphosed” as described by Goethe.

19

20 Another interesting point is the question of how variation on developmental
21 programs between organs in the same organisms relates to variation across
22 organisms. In particular, it would be of interest to what extent the regulatory
23 pathways that produced evolutionary variation in leaf development (Townsend
24 and Sinha, 2012; Tsukaya, 2014) also played a role in establishing the
25 differences between leaves and floral organs. Parallels between
26 morphological diversity between segments of the same organisms and
27 across species may give insight not only into organ development in individual
28 species, but also into the evolutionary diversity of plant form.

29

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33

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Figure legends

Figure 1:

MADS-domain proteins function combinatorially to modify organ identity. Coloured circles represent the organ identity MADS-domain proteins from *Arabidopsis*; for simplicity, SEP_n represents multiple, partially redundant SEP proteins. A) The ground state of floral organs is similar to leaves (here, an *Arabidopsis* cauline leaf is shown); in different floral whorls, different combinations of organ identity modify the ground state organ to sepals (B), petals (C), stamens (D) or carpels (E). Scale bar: 1 mm.

Figure 2:

Example of computational modeling of the effect of *JAG* (one of the targets of organ identity proteins) on organ growth (based on Sauret-Güeto et al, 2013).

A) Schematic wild-type organ primordium with key model assumptions represented: proximal and distal organizers (orange and green lines, respectively) orient local tissue polarity (blue arrows); a growth factor expressed more highly in the distal region of the organ (red gradient) preferentially promotes growth perpendicular to local polarity. B) *JAG* function is assumed to correspond to the growth factor (red gradient) in A, and in addition is required to establish a continuous distal organizer (green line); the dotted black arrow represents growth perpendicular to local polarity (blue arrow). C) Running the model to a state corresponding to a mature petal results in a morphology similar to that of a wild-type petal (D). E, F) Initial state and assumptions of the model corresponding to the *jag* mutant: growth perpendicular to local polarity is reduced, and the distal organizer (green line) is discontinuous. G,H) Running the simulation to a state corresponding to a mature petal results in a narrow organ with serrated edges, which are features of *jag* petals (G). Scale bar: 1 mm.

Figure 3:

Complexes of organ identity MADS-domain proteins (represented by the coloured circles, see Figure 1) directly regulate processes required at all

stages of organ development, including early organ patterning, subsequent organ growth and final cellular differentiation. The lower panels show: on the left, expression of *CUC1* (as an example of patterning gene) revealed by *in situ* RNA hybridization on a section through an early floral bud; middle: outlines of a growing petal (based on Sauret et al. 2013); right: scanning electron micrograph of conical cells of the petal epidermis (as an example of differentiated cell in a mature organ). Scale bar: 50 μm .

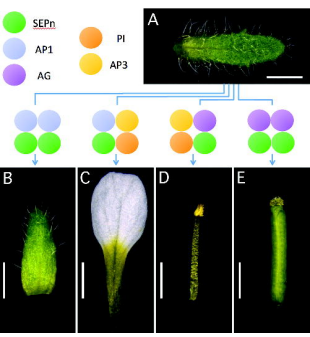


Figure 2

