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# Molecular cartography of leaf development — role of transcription factors



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Organ elaboration in plants occurs almost exclusively by an increase in cell number and size. Leaves, the planar lateral appendages of plants, are no exception. Forward and reverse genetic approaches have identified several genes whose role in leaf morphogenesis has been inferred from their primary effect on cell number and size, thereby distinguishing them as either promoters or inhibitors of cell proliferation and expansion. While such classification is useful in studying size control, a similar link between genes and shape generation is poorly understood. Computational modelling can provide a conceptual framework to re-evaluate the known genetic information and assign specific morphogenetic roles to the transcription factor-encoding genes. Here we discuss recent advances in our understanding of the roles of transcription factors in the planar growth of leaf lamina in two orthogonal dimensions.

#### Addresses

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#### Introduction

Leaves evolved from ancestral branching systems multiple times among the vascular plant lineages [1]. Angiosperm leaves share a common phylogenetic origin, which is reflected in the conserved sequence of developmental events leading up to the initiation of a rod-shaped bulge called primordium on the flanks of the shoot apical meristem (SAM), having distinct upper and lower sides [2]. Yet, angiosperm leaves are distinguished by their tremendous architectural diversity in the final size, shape, and complexity [2,3]. This is attributed to the variation in

the post-initiation growth patterns along the 'base-to-tip' and the 'middle-to-margin' axes [2,4]. The simple leaves of the winter annual Arabidopsis, the lobed leaves of lettuce, the compound leaves of tomato and even the intricately-designed insect-trapping leaves of the bladderwort arise by the modification of the growth parameters, such as growth duration and growth direction, along these two perpendicular axes.

Growth in a leaf primordium occurs by an increase in cell number and cell size, which is regulated in space and time primarily through the activities of growth-promoting and growth-repressing transcription factors, sometimes themselves expressed in gradients, as a result of their transcriptional regulation and/or post-transcriptional control by the upstream regulatory microRNAs. Transcriptional output is often a modulation of cellular properties and response to hormones, which act as intercellular messengers. These molecular players lay down the blueprint for growth patterns in space and time. The challenge is to decipher the connection between proximal effects of gene activities and its ultimate manifestation on organ growth. The current review highlights the attempts made over the past decade in joining the dots by making use of the insights gleaned from computational models.

#### How do leaves grow?

A leaf is characterized by a flat, bifacial lamina with a stalk-like base. Laminar growth proceeds by division and expansion of cells thought to arise from a short-lived meristematic activity localized at the base of the primordium [5]. Initially, active proliferation occurs throughout the primordium but is quickly limited to the base by the sudden appearance of an arrested zone at the distal end [6–9]. Cells distal to the proliferation-differentiation boundary, also called arrest front, contribute to growth by undergoing expansion and maturation, whereas cells proximal to the arrest front continue to increase in number till the complete cessation of proliferating activity. Thereafter, growth is propelled by post-mitotic cell expansion alone till the mature size is achieved. This linear gradient of cell division and expansion along the proximo-distal axis, though common to most monocot and dicot model species is, however, not universal and several other types of growth gradient exist, at least among the eudicot species (Box 1, Figure 3) [10,11\*\*]. Nevertheless, whereas the link between the base-to-tip growth gradient and final leaf size has been well-studied [12,13], how the

#### Box 1 Divergent growth polarity patterns and their evolution

Leaves of all model plants studied so far display a common 'basipetal' pattern of progression of the cell-proliferation arrest from the distal tip towards the proximal base [6,7,15,76]. Even though such a growth pattern was initially assumed to be universal in plants, a broader study using a large number of dicot species revealed three other types of proximo-distal gradient: firstly, acropetal pattern where the arrest front progresses in the base-to-tip direction, secondly, bi-directional pattern where two simultaneous arrest fronts progress from base-to-tip and tip-to-base and thirdly, diffuse growth where there is no progression of the growth arrest and cells throughout the lamina proliferate and differentiate simultaneously (Figure 3) [11\*\*]. These divergent patterns are also closely associated with the expression of growth promoting transcription factors (such as the GRFs) and growth repressing genes (such as miR396) (Figure 3) [11\*\*]. A phylogenetic analysis showed that some of these growth patterns evolved independently in several plant lineages [32°]. It has been speculated that the direction of the growth pattern is under the control of a conserved gene regulatory module, which was co-opted to novel expression domains during evolution to create diverse growth gradients, either by mutation in the proximalregulatory genes (e.g. CIN-TCPs that regulate miR396 and hence GRF expression) or in the regulatory sequences of the individual genes (e.g. GRFs) [11\*\*].

patterns of cell division and expansion generate speciesspecific shapes is still an enigma.

Classical studies tracking laminar growth by observing the movement of artificial landmarks during development or by clonal analysis in fig, tobacco and cotton leaves, had already established that the growth of a primordium does not occur by uniform enlargement of the primordial bulge [14–18]. Rather, different regions of the primordium grow at different rates and orientations that can be quantified using computational tools [19,20°]. Growth orientations diverge at the leaf base and converge towards the tip [21°]. Not only do the growth rates progressively decrease from base to tip as reflected in the patterns of cell division and cell expansion, they differ along the medio-lateral axis as well, with more growth in the lateral than in the medial domain [21\*\*]. Recently, tracking lamina growth in fluorescently-labelled Arabidopsis leaf cells has revealed that the growth pattern is established early on during the primordium development [21<sup>\*\*</sup>].

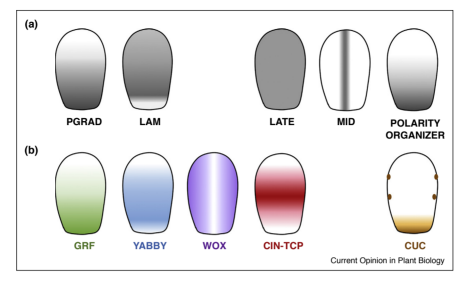
Although such studies provide a roadmap for describing growth patterns, the major challenge is to understand how they are specified at the genetic level [4,22]. Gene expression patterns in space and time, when combined with mutant phenotypes, inform us how a gene specifies growth rate and/or orientation locally. However, cells in a tissue are mechanically constrained by being connected to their neighbours and a specified growth pattern of cells in a region may conflict with that in the neighbouring regions leading to buckling of the tissue out of plane generating curvature or bending [4]. Thus, a 'resultant' growth pattern is an emergent property of any anisotropically-growing organ and may not be intuited from studying gene mutant phenotypes and/or expression domains in isolation [4].

# Modelling leaf growth predicts a molecular toolkit for shape specification

Time-lapse growth analyses in combination with computational modelling can reveal coherent links between gene activity and the resultant growth patterns [8,21°,23,24°,25°]. While elegant reports described modelling growth in the petals or leaf margin [24°,25°], Kuchen et al built a model to simulate the observed leaf shape and growth patterns in Arabidopsis by incorporating two key systems — a network of factors specifying the *rate* of growth and a factor for determining the orientation of growth along the two orthogonal laminar axes [21\*\*] (Figures 1a and 2 a). Two growth-promoting factors, PGRAD that is expressed in a decreasing gradient from base to tip, and LAM that is expressed uniformly, promote growth along the proximo-distal and the medio-lateral axes, respectively. In addition, two growth-inhibitory factors were introduced into the model to account for the cessation of leaf growth—a uniformly-distributed late-acting factor LATE and a midline-restricted factor MID that repress growth along the proximo-distal and the medio-lateral axes, respectively. Finally, a tissue polarity organizer, expressed in a decreasing base-to-tip gradient, was included that determined the direction of growth (Figure 1a). The regulation of growth rate was thus uncoupled from that of growth orientation. Further, the distribution of these factors changed in space and time with growth, allowing a feedback from tissue deformation to specified growth pattern. This 'deforming growth-orientation organizer' model could accurately reproduce shape changes and growth patterns observed experimentally in a wild-type Arabidopsis leaf. Varying the parameters of the model generated a variety of shapes that are observed among the simple-leaved species with smooth margins, indicating that the specification of growth patterns through interactions of growth-modulating and polarity-determining factors along the orthogonal axes underlies diverse simple leaf forms. An independent study incorporated the directional growth of veins as the major determinant of the orientation of specified growth and the resulting model accounted for the generation of diversity in complex leaf shapes, including serrations, lobes and leaflets [26°].

The value of computational modelling lies in its testable predictions about gene action in morphogenesis, leading to new insights. For example, modelling the development of a petal primordium that shows divergent growth pattern at its tip relative to its base, in contrast to the leaf, led to the prediction of the existence of a distally-expressed polarity organizer in addition to a proximally-expressed organizer [24\*\*]. Based on its expression domain and mutant phenotype which matched the requirement of the model, the

Figure 1



Spatial expression domains of the growth-regulating factors along the longitudinal axis of the early leaf primordia represented as rod-shaped structures. Both the hypothetical factors of the growing polarized tissue framework model in Kuchen *et al.* (a) and the proposed counterparts discussed in the text (b) are shown. The expression domains depicted in (b) are approximated from published studies on GRF, [30]; YABBY, [45\*\*]; WOX, [63\*]; CIN-TCP, [7]; CUC, [25\*]. The primordium depicted in (b) is assumed to be at an early growth stage (up to ~2.0 mm in length), reflecting the dynamic expression of the growth-regulatory genes described here.

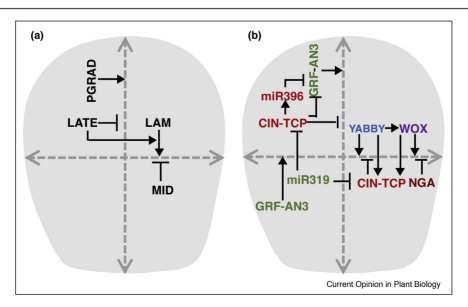
JAGGED gene encoding a Zn-finger transcription factor was identified and validated as the distal polarity organizer in addition to its previously-known function in petal cell division [24\*\*]. Likewise, the model proposed by Kuchen et al. suggests candidate transcription factor-encoding genes that could function as the 'leaf architects', based on their known expression patterns and developmental

roles [21\*\*] (Figures 1b and 2 b). Following is a discussion on such factors and their modes of action.

#### Growth-promoting factors along the proximodistal axis

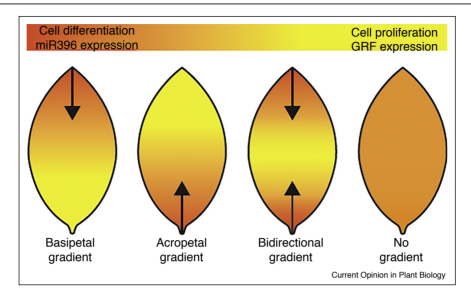
*GROWTH-REGULATING FACTORS (GRFs)* — These are a conserved group of plant-specific transcription factors that

Figure 2



Growth-regulatory network consisting of positive and negative regulatory factors in the early leaf primordium as hypothesized by Kuchen et al. (a) and as discussed in the text (b). The proximo-distal and the medio-lateral axes are represented by the orthogonally intersecting broken grey lines, along which growth is regulated by transcription factors as indicated.

Figure 3



Multiple growth polarities in eudicot leaves. The young leaf primordia show cell proliferation throughout the organ at inception. A pattern of cell differentiation is later superposed on this proliferating sheet of cells. The direction of the progression of the arrest front (black arrow) determines leaf growth polarity. The arrest front progresses from tip-to-base (basipetal gradient), base-to-tip (acropetal gradient) or from both ends to the middle (bidirectional gradient). In some cases, cells throughout the leaf exit proliferation simultaneously, thereby showing no growth polarity. The differentiating regions of the leaves show a strong correlation with the expression of miR396 while the proliferating region shows a gradient of GRF expression.

promote lateral organ growth [27]. Arabidopsis grf mutants have smaller leaves with reduced size and cell number, pointing towards a role in enhancing cell proliferation, although some family members also enhance cell size [28,29]. GRF expression is strongly associated with the base-anchored proliferation zone in the lamina [30] and is down-regulated in differentiated cells at the tip due, at least in part, to the activation of miR396 that targets seven of the nine GRFs in Arabidopsis [30,31]. The expression gradient of the miR396-GRF module along the proximo-distal axis is conserved in other species with basipetal growth pattern, suggesting that this module is a primary factor in promoting growth in this axis, similar to the PGRAD factor in the Kuchen et al. model [21\*\*] (Figures 1b and 2 b). However, grf mutants have narrower leaves, indicating that growth along the medio-lateral axis (perpendicular to midvein in the Kuchen et al. model) is also affected, leading to a change in shape. Remarkably a recent study has identified a correlation between the polarity of miR396-GRF expression gradient and that of the leaf growth gradient (Box 1) (Figure 3) [11°,32°].

How GRFs stimulate proliferation directly is at present unclear. They seem to enhance the duration of cell proliferation and the number of cells undergoing proliferation [33°]. GRFs are proposed to regulate their target genes through a conserved interaction with GRF-INTERACTING FACTORs (GIFs) (Figure 2b). Loss of GIFs also results in narrower leaves with reduced cell number [34-36]. They function as transcriptional coactivators that lack a DNA-binding domain [34,35], though GIF1/ANGUSTIFOLIA3 (AN3) interacts and co-regulates its target genes with several components of the SWI/SNF chromatin remodelling complex containing BRAHMA or SPLAYED ATPases [37°]. Furthermore, AN3 transcript is specifically enriched in the mesophyll cells and its protein product moves to the epidermis, coordinating cell proliferation across the clonally distinct cell types [38°]. Transcriptome profiling of the 35S:AN3-GR transgenic line revealed up and down-regulation of transcripts that are inversely regulated during the transition from proliferation-driven to expansion-driven growth phases, supporting a role for AN3-GRF in balancing proliferation versus differentiation in developing leaves [12,37°]. In addition, AN3 promotes its own transcription and that of GRF3/5/6, thereby providing a molecular basis for synergistic effect of simultaneous overexpression of AN3 and GRF in leaf development [29,37°]. Interestingly, in maize, ZmAN3 is associated preferentially with ZmGRF1 in proximal division zone and with ZmGRF10 in the distal expansion zone; this preference reflects the mRNA and protein abundance of the respective ZmGRF partners [39]. Because ZmGRF1 stimulates and ZmGRF10 limits cell proliferation, it is hypothesized that the competition for ZmAN3 binding by different GRFs along the proximo-distal axis determines the position of the transition zone between that of division and expansion in growing leaves [39,40].

GRFs may promote laminar outgrowth by other indirect mechanisms. A study suggests that GRFs repress class I KNOX genes in both monocots and dicots [41]. Overexpression of GRF5 in Arabidopsis and GRF2 in Brassica napus leads to an increased chloroplast division and chlorophyll accumulation with strong induction of the PORA gene encoding an enzyme in the tetrapyrrole pathway for chlorophyll biosynthesis [33°,42]. Enhanced chlorophyll content is linked to the promotion of cell-cycle progression as the intermediates of the chlorophyll biosynthesis activate cyclin-dependent kinases through chloroplast-tonucleus signalling, suggesting that GRFs link chloroplast division with cell division [33°,43]. Further, AtGRF7 represses several stress-response genes under normal conditions which would otherwise be detrimental to growth [44].

# Growth-promoting factors along the mediolateral axis

The YABBY and WUSCHEL RELATED HOMEOBOX (WOX) transcription factors—These proteins promote lamina outgrowth downstream to the adaxial-abaxial polarity establishment [45\*\*,46]. YABBYs encode small proteins with zinc-finger and helix-loop-helix domains, conserved among all seed plants [47]. Arabidopsis plants mutated for four abaxially-expressed, vegetative YABBY genes form leaves with severe to moderate loss of lamina and marginal tissues with polarity defects [45°]. A subgroup of WOX genes encoding transcriptional repressors, comprising of the PRESSED FLOWERS (PRS) and the MAEWEST/WOX1 subclades, regulate laminar expansion specifically along the medio-lateral axis [46,48,49\*\*]. These factors are expressed in the so-called middle domain between the adaxial and the abaxial domains [49°°,50°]. The prs wox1 double mutant in Arabidopsis produces narrow leaves with perturbed polarity but unaltered length [49\*\*]; similar phenotype is associated with the loss of WOX1 homologues in other dicots and that of PRS homologues in maize [46,48,51].

Given their lamina-wide expression pattern and role as 'lamina identifiers', the YABBY and the WOX genes qualify as the LAM factor that determines growth along the medio-lateral axis in the Kuchen et al model (Figure 1). YABBY activity may be upstream to that of WOX, as the YABBY member FILAMENTOUS FLOWER (FIL) was shown to up-regulate WOX1 expression [49°]. Vegetative YABBYs regulate a broad lamina-specific genetic program involving the repression of SAM identity and maintenance genes (WUSCHEL and KNOXI), promoting expression of the polarity and lamina maturation markers [45°,52]. The YABBY factors may act in concert with AINTEGUMENTA (encoding an AP2/ERF family transcription factor), as the fil ant and the yab3 ant double mutants have reduced lamina growth compared to the single mutants [53]. Whereas, the WOX genes mainly control cell proliferation along the medio-lateral axis,

possibly by recruiting transcriptional repressors like TOPLESS to its targets, resulting in the indirect activation of growth-promoting transcription factors (SCARE-CROW-like), enzymes (KLU/CYP78A5), cell-cycle factors (D-type cyclins) and metabolic pathways leading to auxin biosynthesis and cytokinin signalling [46,49°,54]. Both YABBY and WOX factors regulate lamina outgrowth non-cell autonomously, affecting differentiation of tissues lacking their expression [45°,49°,52,55,56]. It is possible that they generate a mobile signal, such as the phytohormones. YABBYs have been postulated to control auxin response and flux along the leaf margin; the quadruple *yabby* mutant in Arabidopsis shows perturbed venation and lack of marginal structures [45°]. The WOX1 homologue in Nicotiana sylvestris, LAM1, promotes auxin biosynthesis and co-application of auxin and cytokinin can rescue lamina growth defect in the lam1 mutant [46].

### Growth-repressing factors — common to proximo-distal and medio-lateral axes?

CINCINNATA-LIKE **TEOSINTE** BRANCHED1/ CYCLOIDEA/PROLIFERATING CELL**FACTORS** (CIN-TCP) transcription factors—These belong to the plant-specific, non-canonical, bHLH domain-containing TCP family that redundantly repress growth [57]. Five Arabidopsis CIN-TCPs are post-transcriptionally co-regulated by microRNA319 (miR319) [58]. Perturbation of the conserved miR319-TCP module either by mutation of CIN-TCPs or by ectopic expression of miR319 results in larger leaves with altered shape and loss of flatness due to prolonged cell proliferation phase more towards the margin [7,58,59]. On the other hand, premature or increased activation of these factors leads to precocious cellular, organ and organism maturation, suggesting that they are heterochronic regulators of morphogenesis [60-62].

CIN-TCP expression occurs in a dynamic spatio-temporal gradient during the primordium development, as an output of their transcriptional and the miR319-mediated post-transcriptional control [63°] (Figure 1b). Initially, the CIN expression in snapdragon and TCP4 expression in Arabidopsis starts at the tip and later gets restricted to the leaf base, overlapping with the base-anchored proliferation zone, before disappearing with the cessation of mitotic activity [7,63°]. Given the relatively late onset of CIN-TCP growth-repressor activity, compared to the growth-promoters like YABBYs, CIN-TCPs could serve as the late-acting growth-inhibitor LATE (Figure 2b). However, LATE is proposed to be uniformly distributed and retards growth along proximo-distal axis while promoting growth along medio-lateral axis. This is inconsistent with the expression pattern and activity of CIN-TCPs which restrict growth along both the axes, as evident from phenotypes of their gain-of-function and loss-of-function mutants [61].

CIN-TCPs are postulated to mainly promote the onset of cell maturation program, thereby indirectly inhibiting cell proliferation [61]; although some studies suggest a direct link with cell-cycle suppression [64,65]. CIN-TCPs also activate miR396 expression leading to temporal and spatial decline in its cognate GRF target levels [30,65] (see Box 1). They also down-regulate GRF5/6 and AN3 expression independent of miR396 [30], thus efficiently repressing the overall growth-promoting activity of the GRF-AN3 complex along the proximo-distal axis, similar to the antagonistic activities proposed for the PGRAD and LATE factors (Figure 2). In addition, CIN-TCPs regulate the level and/or response to several growthregulating phytohormones such as auxin, cytokinin, gibberellic acid and jasmonic acid among others [66,67].

In snapdragon, CIN expression shows a strong mediolateral gradient; more expression at the margin than at the centre of lamina. This is consistent with more de-repression of growth and mitotic marker expression at the margin than at the centre in the cin mutant, suggesting that CIN-TCPs also function as a component of the proposed MID that inhibits growth along the mediolateral axis, though the observed CIN-TCP expression pattern is in contrast to that hypothesized for MID [7,21°]. Two recent studies have shed light on the relevance of CIN-TCP activity in the medio-lateral axis to the regulation of leaf morphogenesis [68°,69°]. Transcriptome profiling of young wild-type snapdragon and cin mutant leaves allowed the identification of AmHISTI-DINE KINASE4 (encoding a homolog of Arabidopsis cytokinin receptor) and AmIAA3/SHY2 (encoding a homolog of the Arabidopsis AUX/IAA repressors of auxin signalling), as the direct downstream targets of CIN [68°]. Interestingly, these two genes are expressed more strongly at the margins than at the medial region, similar to CIN, raising the possibility that CIN regulates growth suppression at the margins by modulating the balance between auxin and cytokinin signals [70]. On the other hand, direct identification of the margin and centreenriched genes in Arabidopsis enabled an analysis of their differential regulation in the tcp2/3/4/10 quadruple mutant [69\*\*]. The margin-enriched genes (genes expressed more in the margins than in the centre) were more down-regulated in the cin-tcp mutant than the centre-enriched genes and included several transcription factors known to control margin development, such as the NGATHA, STYLISH and even the miR319-resistant CIN-TCP family members. Other differentiation markers such as photosynthesis-related genes, shown to be expressed during the proliferation-to-expansion transition [9], were more down-regulated at the margins than at the centre of cin-tcp mutant; in contrast, the mitosis markers and the growth-promoting transcription factors such as ANT and WOX1 were up-regulated [69\*\*]. These studies have established CIN-TCPs as the major growth repressors not only along the proximo-distal axis but also along the medio-lateral axis: thus CIN-TCPs could serve as the LATE and MID factors that inhibit growth (Figure 2b).

Another study has revealed a novel role of CIN-TCPs and NGA genes in regulating determinate leaf growth [63°]. Simultaneous down-regulation of miR319-targeted CIN-TCPs and four NGA genes, either throughout the lamina or only at the margin, results in a dramatic indeterminate growth with sustained de novo organogenesis at the margin, whose molecular signature resembles that of undifferentiated initiating leaf primordia. Strikingly, there was no ectopic expression of SAM-specific genes at the margin; the authors proposed that the phenotype rather results from the de-repression of a short-lived bonafide 'leaf meristem', which is normally active only transiently at the leaf base and is kept suppressed in the distal margins by the coordinated CIN-TCP and NGA activity [63°]. CIN-TCPs have been shown to directly activate NGA genes; however, the phenotype of their combined down-regulation, absent from individual cin-tcp and nga mutants, suggests a synergistic relationship between these two family members [63°,71]. Possibly, CIN-TCPs interact with NGA to co-regulate margin-enriched genes for determinate growth [69\*\*] (Figure 2b).

# Growth orientation-determining factors

CUP-SHAPED COTYLEDON transcription factors— These are the NAC (NAM, ATAF1/2, CUPULIFOR-MIS) domain-containing transcription factors, characterized by their roles in organ boundary formation and serration development [72,73]. CUC2 and CUC3 are expressed at the leaf base and margin, demarcating the boundaries of incipient serrations [73]. Anisotropic growth requires a mechanism for the cells to determine the orientation of growth in response to growth regulatory factors. In the Kuchen et al. model, this information is derived from a polarity organizer expressed at the leaf base and growth occurs along the axes parallel and perpendicular to its proximo-distal gradient (Figure 1a). The CUC genes could serve as the candidate organizer based on their expression pattern and function [25°] (Figure 1b).

Cellular anisotropy may result from unequal or polarized distribution of molecules within the cytoplasm (e.g. microtubule cytoskeleton) or at the plasma membrane (e.g. receptors) [4]. CUC activity is associated with the generation of membrane anisotropy. CUC2 promotes reorientation of the plasma-membrane-localized auxin efflux carrier protein PINFORMED1 (PIN1) in the epidermis to form PIN1 convergence points that result in auxin maxima formation at the margins and the subsequent serration outgrowth [25°]. Whether a similar CUC activity regulates growth polarity during laminar outgrowth requires further experimental validation.

Interestingly, CUC activity is repressed by miR319-regulated CIN-TCPs. CIN-TCPs activate the transcription of miR164 that targets CUC2 transcript for degradation [74]. In addition, TCP4 interacts with and inhibits CUC2-CUC3 dimerization and dampens their transactivation potential. This effect is ameliorated by the sequestration of TCP4 by the miR156-targeted SPL transcription factors in the adult vegetative-phase leaves, causing agedependent changes in the leaf margin shape [75°]. Thus, the polarity organizer activity may be subject to regulation by growth-regulatory factors.

#### Concluding remarks

Computational models provide a useful prism through which a complex biological phenomenon can be viewed in order to reveal its underlying component network(s) of interacting molecules. On the other hand, mutant analyses provide valuable mechanistic details that may not be predicted by modelling alone. For example, the yabby and the wox1 mutants (in Arabidopsis and Medicago truncatula, respectively) show down-regulation of CIN-TCP expression, suggesting that the induction of growth-repressors by the growth-promoters is required for balanced determinate growth [45°,46]; although Kuchen et al. hypothesized that the LATE factor enhances the extent to which LAM promotes medio-lateral growth, there is no evidence that CIN-TCPs directly regulate YABBY and WOX genes. Likewise, inhibition of CUCs by CIN-TCPs indicates that the determination of growth orientation may not be independent of the growth regulatory network. It would be interesting to model these interactions and observe the impact on the simulated leaf shape.

Leaf form is a complex trait that requires several genetic regulators [77], of which only a few have been discussed in this review. Transcription factors such as ANT, STRUWWELPETER (a component of RNA Polymer-II-associated Mediator complex), RESPONSE FACTOR2 and SPATULA control the duration of proliferation or the number of cells undergoing proliferation; most of these mutants show alteration in size but not shape, suggesting that they are general regulators of growth [78-81]. The PEAPOD transcription factors, on the other hand, control the onset of a secondary proliferation arrest of the dispersed meristematic cells, more at the centre than the margins [82]. Thus, they can form a component of the putative LATE or MID factors. Another candidate polarity organizer is JAGGED, which serves a similar function in petal growth [24°]. Detailed spatio-temporal analysis of these genes coupled with time-lapse growth analysis of their mutants should clarify their morphogenetic role.

Though modelling and experimental approaches together explain how diverse and complex leaf shapes can be generated, many questions still remain. Firstly, what is the evolutionary significance of this diversity. Indeed, the existence of any strong selection pressure on leaf shape in a specific environment is still debated [2].

Secondly, what is the advantage of evolving diverse growth gradients, as many species grow leaves without any gradient [11°]. It has been suggested that specific growth patterns confer adaptive advantage depending on the ecological niche of the plant species [32°]. A detailed 'eco-evo-devo' approach will be required to gain deeper insights.

#### Conflict of interest statement

Nothing declared.

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#### References and recommended reading

Papers of particular interest, published within the period of review, have been highlighted as

- of special interest
- of outstanding interest
- Harrison CJ, Morris JL: The origin and early evolution of vascular plant shoots and leaves. Philos Trans R Soc B Biol Sci 2018. 373:20160496
- Nicotra AB, Leigh A, Boyce CK, Jones CS, Niklas KJ, Royer DL, Tsukaya H: The evolution and functional significance of leaf shape in the angiosperms. Funct Plant Biol 2011, 38:535-552.
- Chitwood DH, Sinha NR: Evolutionary and environmental forces sculpting leaf development. Curr Biol 2016, 26:R297-R306.
- Whitewoods CD, Coen E: Growth and development of threedimensional plant form, Curr Biol 2017, 27:R910-R918.
- Ichihashi Y, Kawade K, Usami T, Horiguchi G, Takahashi T, Tsukaya H: Key proliferative activity in the junction between the leaf blade and leaf petiole of Arabidopsis. Plant Physiol 2011, **157**:1151-1162
- Donnelly PM, Bonetta D, Tsukaya H, Dengler RE, Dengler NG: Cell cycling and cell enlargement in developing leaves of Arabidopsis. Dev Biol 1999, 215:407-419
- Nath U, Crawford BC, Carpenter R, Coen E: Genetic control of surface curvature. Science (80-) 2003, 299:1404-1407
- Kazama T, Ichihashi Y, Murata S, Tsukaya H: The mechanism of cell cycle arrest front progression explained by a KLUH/ CYP78A5-dependent mobile growth factor in developing leaves of Arabidopsis thaliana. Plant Cell Physiol 2010, 51:1046-1054
- Andriankaja M, Dhondt S, DeBodt S, Vanhaeren H, Coppens F, DeMilde L, Mühlenbock P, Skirycz A, Gonzalez N, Beemster GTS et al.: Exit from proliferation during leaf development in Arabidopsis thaliana: a not-so-gradual process. Dev Cell 2012, 22:64-78
- 10. Nelissen H, Gonzalez N, Inzé D: Leaf growth in dicots and monocots: So different yet so alike. Curr Opin Plant Biol 2016, **33**:72-76.
- 11. Das Gupta M, Nath U: Divergence in patterns of leaf growth polarity is associated with the expression divergence of miR396. Plant Cell 2015, 27:2785-2799.

This study demonstrated that the proximo-distal leaf growth polarity is divergent among angiosperm species and is associated with the miR396 expression pattern.

12. Gonzalez N, Vanhaeren H, Inzé D: Leaf size control: complex coordination of cell division and expansion. Trends Plant Sci 2012, **17**:332-340.

- 13. Czesnick HR, Lenhard M: Size control in plants lessons from leaves and flowers. Cold Spring Harb Perspect Biol 2015:7
- 14. Avery GS: Structure and development of the tobacco leaf. Am J Bot 1933. 20:565-592.
- 15. Poethig RS, Sussex IM: The cellular-parameters of leaf development in tobacco - a clonal analysis. Planta 1985,
- 16. Dolan L, Poethig RS: Clonal analysis of leaf development in cotton. Am J Bot 1998, 85:315-321.
- 17. Wolf SD, Silk WK, Plant RE: Quantitative patterns of leaf expansion: comparison of normal and malformed leaf growth in Vitis vinifera cv. Ruby Red. Am J Bot 1986, 73:832-846.
- Saurer W, Possingham JV: Studies on the growth of spinach leaves (Spinacea oleracea). J Exp Bot 1970, 21:151-158.
- Remmler L, Rolland-Lagan A-G: Computational method for quantifying growth patterns at the adaxial leaf surface in three dimensions. Plant Physiol 2012, 159:27-39.
- Rolland-Lagan A-G, Remmler L, Girard-Bock C: Quantifying 20. shape changes and tissue deformation in leaf development. Plant Physiol 2014, **165**:496-505.

The authors used computational tools to determine how spatial heterogeneity in growth patterns largely preserve the shape of the Arabidopsis leaves during growth.

- 21. Kuchen EE, Fox S, De Reuille PB, Kennaway R, Bensmihen S,
- Avondo J, Calder GM, Southam P, Bangham A, Coen E: Generation of leaf shape through early patterns of growth and tissue polarity. Science 2012, 335:1092-1096.

This study showed that a simple network of growth-regulatory and growth polarity-determining components underlies diverse simple leaf forms.

- Coen E, Kennaway R, Whitewoods C: On genes and form. Development 2017, 144:4203-4213.
- Prusinkiewicz P. Runions A: Computational models of plant development and form. New Phytol 2012, 193:549-569
- 24. Sauret-Güeto S, Schiessl K, Bangham A, Sablowski R, Coen E:
- JAGGED controls arabidopsis petal growth and shape by interacting with a divergent polarity field. PLoS Biol 2013, 11:

An example of a computational model predicting a novel morphogenetic role for a transcription factor based on its expression pattern and mutant

Bilsborough GD, Runions A, Barkoulas M, Jenkins HW, Hasson A, Galinha C, Laufs P, Hay A, Prusinkiewicz P, Tsiantis M: **Model for** the regulation of Arabidopsis thaliana leaf margin development. *Proc Natl Acad Sci U S A* 2011, **108**:3424-3429.

This study presented a molecular framework describing the interactions between CUC2, PIN1 and auxin flux to explain the initiation and placement of serrations along the leaf margin.

- Runions A, Tsiantis M, Prusinkiewicz P: A common 26.
- developmental program can produce diverse leaf shapes. New Phytol 2017, 216:401-418.

The authors propose a computational model to explain the generation of diversity in complex leaf shapes, based on the assumption that three inter-dependent processes of marginal growth, directional vein growth and passive growth of lamina determine the final leaf shape.

- Omidbakhshfard MA, Proost S, Fujikura U, Mueller-Roeber B: Growth-regulating factors (GRFs): a small transcription factor family with important functions in plant biology. Mol Plant 2015, 8:998-1010.
- Kim JH, Choi D, Kende H: The AtGRF family of putative transcription factors is involved in leaf and cotyledon growth in Arabidopsis. Plant J 2003, 36:94-104.
- Debernardi JM, Mecchia MA, Vercruyssen L, Smaczniak C, Kaufmann K, Inzé D, Rodriguez RE, Palatnik JF: Posttranscriptional control of GRF transcription factors by microRNA miR396 and GIF co-activator affects leaf size and longevity. Plant J 2014, 79:413-426.
- Rodriguez RE, Mecchia MA, Debernardi JM, Schommer C, Weigel D, Palatnik JF: Control of cell proliferation in

- Arabidopsis thaliana by microRNA miR396. Development 2010,
- 31. Liu D, Song Y, Chen Z, Yu D: Ectopic expression of miR396 suppresses GRF target gene expression and alters leaf growth in Arabidopsis. *Physiol Plant* 2009, **136**:223-236.
- 32. Das Gupta M, Nath U: On the evolution of developmental mechanisms: Divergent polarities in leaf growth as a case study. Plant Signal Behav 2016, 11:e1126030.

An evolutionary basis of divergent growth polarity in angiosperm leaves was discussed.

Vercruyssen L, Tognetti VB, Gonzalez N, Van Dingenen J, De Milde L, Bielach A, De Rycke R, Van Breusegem F, Inzé D: **GROWTH REGULATING FACTOR5 stimulates arabidopsis** chloroplast division, photosynthesis, and leaf longevity. Plant Physiol 2015, 167:817-832.

A link between the proliferation status of cells and the number of chloroplast, regulated by GRF5, was elucidated.

- 34. Kim JH, Kende H: A transcriptional coactivator, AtGIF1, is involved in regulating leaf growth and morphology in Arabidopsis. Proc Natl Acad Sci USA 2004, 101:13374-13379.
- Lee BH, Ko J-H, Lee S, Lee Y, Pak J-H, Kim JH: **The Arabidopsis GRF-INTERACTING FACTOR gene family performs an** overlapping function in determining organ size as well as multiple developmental properties. Plant Physiol 2009, 151:655-
- 36. Horiquchi G, Kim GT, Tsukaya H: The transcription factor AtGRF5 and the transcription coactivator AN3 regulate cell proliferation in leaf primordia of Arabidopsis thaliana. Plant J
- 37. Vercruyssen L, Verkest A, Gonzalez N, Heyndrickx KS,

  Eeckhout D, Han S-K, Jegu T, Archacki R, Van Leene J, Andriankaja M et al.: ANGUSTIFOLIA3 binds to SWI/SNF chromatin remodeling complexes to regulate transcription during Arabidopsis leaf development. Plant Cell 2014, 26:210-229

Molecular basis for the gene-regulatory function of AN3 in leaf development was revealed.

- 38. Kawade K, Horiguchi G, Usami T, Hirai MY, Tsukaya H:
- ANGUSTIFOLIA3 signaling coordinates proliferation between clonally distinct cells in leaves. Curr Biol 2013, 23:788-792. This study demonstrated that the AN3 protein moves from mesophyll to epidermis and coordinates cell proliferation between the two clonallydistinct cell layers.
- Nelissen H, Eeckhout D, Demuynck K, Persiau G, Walton A, van Bel M, Vervoort M, Candaele J, De Block J, Aesaert S et al. Dynamic changes in ANGUSTIFOLIA3 complex composition reveal a growth regulatory mechanism in the maize leaf. Plant Cell 2015, 27:1605-1619.
- Wu L, Zhang D, Xue M, Qian J, He Y, Wang S: Overexpression of the maize GRF10, an endogenous truncated growthregulating factor protein, leads to reduction in leaf size and plant height. *J Integr Plant Biol* 2014, **56**:1053-1063.
- 41. Kuijt SJH, Greco R, Agalou A, Shao J, 't Hoen CCJ, Overnas E, Osnato M, Curiale S, Meynard D, van Gulik R et al.: Interaction between the GROWTH-REGULATING FACTOR and **KNOTTED1-LIKE HOMEOBOX Families of Transcription** Factors. Plant Physiol 2014, 164:1952-1966.
- 42. Liu J, Hua W, Yang HL, Zhan GM, Li RJ, Deng L, Bin, Wang XF, Liu GH, Wang HZ: The BnGRF2 gene (GRF2-like gene from Brassica napus) enhances seed oil production through regulating cell number and plant photosynthesis. J Exp Bot
- 43. Kobayashi Y, Kanesaki Y, Tanaka A, Kuroiwa H, Kuroiwa T, Tanaka K: Tetrapyrrole signal as a cell-cycle coordinator from organelle to nuclear DNA replication in plant cells. Proc Natl Acad Sci USA 2009, 106:803-807.
- 44. Kim J-S, Mizoi J, Kidokoro S, Maruyama K, Nakajima J, Nakashima K, Mitsuda N, Takiguchi Y, Ohme-Takagi M, Kondou Y et al.: Arabidopsis GROWTH-REGULATING FACTOR7 functions as a transcriptional repressor of abscisic acid- and osmotic stress-responsive genes, including DREB2A. Plant Cell 2012, 24:3393-3405.

- 45. Sarojam R, Sappl PG, Goldshmidt A, Efroni I, Floyd SK, Eshed Y, Bowman JL: Differentiating Arabidopsis shoots from leaves by combined YABBY activities. Plant Cell 2010, 22:2113-2130. Through in-depth expression and mutant analyses, the authors clarify the role of YABBY genes as lamina identifiers during leaf development.
- Tadege M, Lin H, Bedair M, Berbel A, Wen J, Rojas CM, Niu L, Tang Y, Sumner L, Ratet P et al.: STENOFOLIA regulates blade outgrowth and leaf vascular patterning in Medicago truncatula and Nicotiana sylvestris. Plant Cell 2011, 23:2125-
- 47. Bowman JL, Smyth DR: CRABS CLAW, a gene that regulates carpel and nectary development in Arabidopsis, encodes a novel protein with zinc finger and helix-loop-helix domains. Development 1999, **126**:2387-2396.
- 48. Vandenbussche M, Horstman A, Zethof J, Koes R, Rijpkema AS, Gerats T: Differential recruitment of WOX transcription factors for lateral development and organ fusion in petunia and Arabidopsis. Plant Cell 2009, 21:2269-2283.
- 49. Nakata M, Matsumoto N, Tsugeki R, Rikirsch E, Laux T, Okada K:
- Roles of the middle domain-specific WUSCHEL-RELATED HOMEOBOX genes in early development of leaves in Arabidopsis. *Plant Cell* 2012, **24**:519-535.

49\*\*] and [50\*] together elucidate the molecular basis for the requirement of juxtaposition of adaxial and abaxial tissues for laminar outgrowth by revealing the role of WOX genes in the 'middle domain

- 50. Nakata M, Okada K: A new model for the early development of leaves in Arabidopsis thaliana. The three-domain model. Plant Signal Behav 2012, 7:1-5.
- 49\*\*] and [50\*] together elucidate the molecular basis for the requirement of juxtaposition of adaxial and abaxial tissues for laminar outgrowth by revealing the role of WOX genes in the 'middle domain
- 51. Scanlon MJ, Schneeberger RG, Freeling M: The maize mutant narrow sheath fails to establish leaf margin identity in a meristematic domain. Development 1996, 122:1683-1691.
- Kumaran MK, Bowman JL, Sundaresan V: YABBY polarity genes mediate the repression of KNOX homeobox genes in Arabidopsis. Plant Cell 2002, 14:2761-2770.
- 53. Nole-Wilson S, Krizek BA: AINTEGUMENTA contributes to organ polarity and regulates growth of lateral organs in combination with YABBY genes. Plant Physiol 2006, 141:977-
- Zhang F, Wang Y, Li G, Tang Y, Kramer EM, Tadege M: STENOFOLIA recruits TOPLESS to repress ASYMMETRIC LEAVES2 at the leaf margin and promote leaf blade outgrowth in Medicago truncatula. Plant Cell 2014, 26:650-664.
- 55. Siegfried KR, Eshed Y, Baum SF, Otsuga D, Drews GN, Bowman JL: Members of the YABBY gene family specify abaxial cell fate in Arabidopsis. Development 1999, 126:4117-
- Goldshmidt A, Alvarez JP, Bowman JL, Eshed Y: Signals derived from YABBY gene activities in organ primordia regulate growth and partitioning of Arabidopsis shoot apical meristems. Plant Cell 2008, 20:1217-1230.
- 57. Martín-Trillo M, Cubas P: TCP genes: a family snapshot ten years later. Trends Plant Sci 2010, 15:31-39.
- Palatnik JF, Allen E, Wu X, Schommer C, Schwab R, Carrington JC, Weigel D: Control of leaf morphogenesis by microRNAs. Nature 2003, **425**:257-263.
- 59. Koyama T, Furutani M, Tasaka M, Ohme-Takagi M: TCP transcription factors control the morphology of shoot lateral organs via negative regulation of the expression of boundaryspecific genes in Arabidopsis. Plant Cell 2007, 19:473-484.
- 60. Ori N, Cohen AR, Etzioni A, Brand A, Yanai O, Shleizer S, Menda N, Amsellem Z, Efroni I, Pekker I et al.: Regulation of LANCEOLATE by miR319 is required for compound-leaf development in tomato. Nat Genet 2007, 39:787-791.
- 61. Efroni I, Blum E, Goldshmidt A, Eshed Y: A protracted and dynamic maturation schedule underlies Arabidopsis leaf development. Plant Cell 2008, 20:2293-2306.

- 62. Sarvepalli K, Nath U: Hyper-activation of the TCP4 transcription factor in Arabidopsis thaliana accelerates multiple aspects of plant maturation. Plant J 2011, 67:595-607.
- 63. Alvarez JP, Furumizu C, Efroni I, Eshed Y, Bowman JL: Active suppression of a leaf meristem orchestrates determinate leaf growth. Elife 2016, 5:e15023

Simultaneous knockdown of CIN-TCPs and NGA genes led to a dramatic indeterminate growth throughout the leaf margin, suggesting a novel role for these transcription factors in suppressing leaf meristem.

- 64. Masuda HP, Cabral LM, De Veylder L, Tanurdzic M, De Almeida Engler J, Geelen D, Inzé D, Martienssen RA, Ferreira PCG, Hemerly AS: ABAP1 is a novel plant Armadillo BTB protein involved in DNA replication and transcription. EMBO J 2008, 27:2746-2756.
- 65. Schommer C, Debernardi JM, Bresso EG, Rodriguez RE, Palatnik JF: Repression of cell proliferation by miR319regulated TCP4. Mol Plant 2014. 7:1533-1544.
- 66. Nicolas M, Cubas P: TCP factors: new kids on the signaling block. Curr Opin Plant Biol 2016, 33:33-41.
- 67. Challa KR, Aggarwal P, Nath U: Activation of YUCCA5 by the transcription factor TCP4 integrates developmental and environmental signals to promote hypocotyl elongation in Arabidopsis. *Plant Cell* 2016, **28**:2117-2130.
- 68. Das Gupta M, Aggarwal P, Nath U: CINCINNATA in Antirrhinum
- majus directly modulates genes involved in cytokinin and auxin signaling. New Phytol 2014, 204:901-912.
   This study linksCINCINNATA with the phytohormone response in the

regulation of leaf growth along the proximo-distal and the medio-lateral

- 69. Bresso EG, Chorostecki U, Rodriguez RE, Palatnik JF,
- Schommer C: Spatial control of gene expression by miR319regulated TCP transcription factors in leaf development. Plant Physiol 2017, 176:1694-1708.

First study in Arabidopsis to identify the margin-enriched versus centerenriched genes during lamina development in the wild-type and cin-tcp mutant leaves.

- 70. Sarvepalli K, Nath U: CIN-TCP transcription factors: Transiting cell proliferation in plants. IUBMB Life 2018 http://dx.doi.org/ 10.1002/jub.1874.
- 71. Ballester P, Navarrete-Gomez M, Carbonero P, Onate-Sanchez L, Ferrandiz C: Leaf expansion in Arabidopsis is controlled by a TCP-NGA regulatory module likely conserved in distantly related species. Physiol Plant 2015, 155:21-32.
- 72. Aida M. Ishida T. Tasaka M: Shoot apical meristem and cotyledon formation during Arabidopsis embryogenesis: interaction among the CUP-SHAPED COTYLEDON and SHOOT MERISTEMLESS genes. Development 1999, 126:1563-
- 73. Hasson A, Plessis A, Blein T, Adroher B, Grigg S, Tsiantis M, Boudaoud A, Damerval C, Laufs P: Evolution and diverse roles of the CUP-SHAPED COTYLEDON genes in Arabidopsis leaf development. Plant Cell 2011, 23:54-68.
- 74. Koyama T, Mitsuda N, Seki M, Shinozaki K, Ohme-Takaqi M: TCP transcription factors regulate the activities of ASYMMETRIC LEAVES1 and miR164, as well as the auxin response, during differentiation of leaves in Arabidopsis. Plant Cell 2010, 22:3574-3588.
- Rubio-Somoza I, Zhou CM, Confraria A, Martinho C, Von Born P,
   Baena-Gonzalez E, Wang JW, Weigel D: Temporal control of leaf complexity by miRNA-regulated licensing of protein complexes. Curr Biol 2014, 24:2714-2719.

This study revealed the mutual, age-dependent interactions of CUC, CIN-TCP and SPL transcription factors and how they shape leaf development.

- Poethig RS, Szymkowiak EJ: Clonal analysis of leaf development in maize. Maydica 1995, 40:67-76.
- 77. Powell AE, Lenhard M: Control of organ size in plants. Curr Biol 2012. 22:R360-R367.
- 78. Mizukami Y, Fischer RL: Plant organ size control: AINTEGUMENTA regulates growth and cell numbers during organogenesis. Proc Natl Acad Sci U S A 2000, 97:942-947

- 79. Autran D, Jonak C, Belcram K, Beemster GTS, Kronenberger J, Grandjean O, Inzé D, Traas J: Cell numbers and leaf development in Arabidopsis: a functional analysis of the **STRUWWELPETER gene**. *EMBO J* 2002, **21**:6036 LP-6049.
- 80. Schruff MC, Spielman M, Tiwari S, Adams S, Fenby N, Scott RJ: The AUXIN RESPONSE FACTOR 2 gene of Arabidopsis links auxin signalling, cell division, and the size of seeds and other organs. Development 2006, 133:251-261.
- 81. Ichihashi Y, Horiguchi G, Gleissberg S, Tsukaya H: **The bHLH** transcription factor **SPATULA** controls final leaf size in *Arabidopsis thaliana*. *Plant Cell Physiol* 2010, **51**:252-261.
- 82. White DWR: **PEAPOD regulates lamina size and curvature in Arabidopsis**. *Proc Natl Acad Sci USA* 2006, **103**:13238-13243.