The genetic control of leaf and petal allometric variations in Arabidopsis thaliana

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Research article

Keywords: Arabidopsis thaliana, Leaf and petal, Allometric variation, QTL mapping, Multiparent Advanced Generation Intercross Lines

DOI: https://doi.org/10.21203/rs.2.20553/v3

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Abstract

Background

Organ shape and size covariation (allometry) are essential concepts for the study of evolution and development. Although ample research has been conducted on organ shape and size, little research has considered the correlated variation of these two traits and quantitatively measured the variation in a common framework. The genetic basis of allometry variation in a single organ or among different organs is also relatively unknown.

Results

A principal component analysis (PCA) of organ landmarks and outlines was conducted and used to quantitatively capture shape and size variation in leaves and petals of multiparent advanced generation intercross (MAGIC) populations of *Arabidopsis thaliana*. The PCA indicated that size variation was a major component of allometry variation and revealed negatively correlated changes in leaf and petal size. After quantitative trait loci (QTL) mapping, five QTLs for the fourth leaf, 11 QTLs for the seventh leaf, and 12 QTLs for petal size and shape were identified. These QTLs were not identical to those previously identified, with the exception of the ER locus. The allometry model was also used to measure the leaf and petal allometry covariation to investigate the evolution and genetic coordination between homologous organs. In total, 12 QTLs were identified in association with the fourth leaf and petal allometry covariation, and eight QTLs were identified to be associated with the seventh leaf and petal allometry covariation. In these QTL confidence regions, there were important genes associated with cell proliferation and expansion with alleles unique to the maximal effects accession. In addition, the QTLs associated with life-history traits, such as days to bolting, stem length, and rosette leaf number, which were highly coordinated with climate change and local adaption, were QTL mapped and showed an overlap with leaf and petal allometry, which explained the genetic basis for their correlation.

Conclusions

This study explored the genetic basis for leaf and petal allometry and their interaction, which may provide important information for investigating the correlated variation and evolution of organ shape and size in *Arabidopsis*.

Background

Organ morphology is determined by organ shape and size, and coordinated variation in shape and size is a major component of natural diversity. Allometry refers to the size-related changes in morphological traits and can be used to describe the correlated variation in shape and size that can occur within one type of organ or can involve the relative proportions of different organs (Langlade et al., 2005; Feng et al., 2009; Klingenberg, 2016). Even closely related species can still show very different allometries, possibly due to the correlations resulting from selection (Galen, 2006; McDonald et al., 2003) and developmental
constraints (Smith et al., 2014). Leaves and petals, for example, are homologous organs that share mechanisms of developmental control (Anastasiou and Lenhard, 2007; Powell and Lenhard, 2012) so that genes that act pleiotropically on both organ types might give rise to coordinating changes in shape or size. The genetic and evolutionary basis for allometric variation is integral to our understanding of plant development. However, these are poorly understood.

Addressing this question requires a quantitative genetic framework that incorporates allometric relationships to allow intraspecies differences to be evaluated in relation to gene action. Size differences can be expressed in terms of rather simple one-dimensional measurements (e.g., length, height). Shape, on the other hand, is a much more complex feature, and it is far more difficult to measure and compare. In the 1980s, advances in the development of statistical analysis tools and their combination with outline and landmark data revolutionized the field of geometric morphometrics (Prpic and Posnien, 2016). A point and outline approach was first amplified to quantify allometric variation within the leaves of the snapdragon (Antirrhinum) species (Langlade et al., 2005). Covariation in the positions of multiple points around leaf outlines was described in terms of principal components (PCs) that captured variation in both shape and size. The principal component analysis (PCA) on the parameters then allowed the major sources of variation to be identified. This method captures allometric variation directly without resulting in shape and size separation and achieves the incorporation of different types of organs within the same framework. It was applied to quantify the allometric variation between leaves and flowers in Antirrhinum, and the resulting measures of allometry allowed underlying genes to be mapped as QTLs in a genetically segregating population (Feng et al., 2009; Costa et al., 2012).

*Arabidopsis thaliana* is an ideal organism for the study of natural variation in leaf and petal shape and size because there are extensive variations among worldwide accessions for both of these traits and for many life-history traits (Alonso-Blanco and Koornneef, 2000; Kover et al., 2009; Weigel, 2012). Leaves and petals have an advantage, as both their shapes and sizes can be readily captured for an initial approximation by a two-dimensional (2D) outline. Previous studies specifically featured a QTL analysis of leaf and petal shape and size in *Arabidopsis thaliana*. Recombinant inbred lines (RILs) from a *Ler-0 × Col-4* cross identified a total of 16 and 13 QTL-harboring, naturally occurring alleles that contributed to natural variations in the architecture of juvenile and adult leaves, respectively (Pérez-Pérez et al., 2002). In the *Ler × Cvi* RIL population, eight QTLs for petal traits and three QTLs for leaf traits were identified (Juenger et al., 2005). Abraham et al. (2013) found 23 QTLs for variation in petal length, width, area, and shape in two RIL populations (*Col-0 × Est-1* and *Ler-0 × Col-4*). In addition, many factors controlling leaf and petal shape and size have been identified and have been shown to be regulated by hormonal signals, transcription factors and miRNAs during leaf and petal development, and recent findings have highlighted the contribution of mechanical signals to leaf and petal growth (Czesnick and Lenhard, 2015; Moyroud and Glover, 2017; Maugarny-Calès and Laufs, 2018). However, neither these QTLs nor the factors identified could capture the allometry variation of leaves and petals due to the limitation of the common measures in capturing the shape variation fully and in integrating the analysis of shape and size (Klingenberg, 2003).
In this study, we investigated the genetic basis of natural allometry variation in leaves and petals using a set of RILs of *Arabidopsis thaliana* that were derived from multiparent advanced generation intercross (MAGIC) lines, which were constructed by 19 founder accessions (Kover et al., 2009). Multiparent lines are better for addressing genetic correlations due to the larger number of alleles and recombination events, which allows for mapping to smaller intervals (Kover et al., 2009). In addition, the larger number of alleles improves the ability to determine whether the distributions of allelic effects are compatible with pleiotropy. Moreover, we used a quantitative approach based on a PCA of landmark positions to define allometric spaces that captured variation in shape and size (Langlade et al., 2005; Bensmihen et al., 2008; Feng et al., 2009; Costa et al., 2012), which was treated collectively to allow allometric relationships to be defined.

**Results**

**Allometry models of leaves and petals in MAGIC lines**

To examine the allometric variation of leaves and petals within *Arabidopsis thaliana*, an allometric method based on a PCA of organ landmarks and outlines was used to quantify this trait. Leaf4, Leaf7 and petals from MAGIC lines were modelled and generated a separate data set (Supplementary Figure 1). Then, PCA was applied to detect the variation in the positions of points and to identify trends in shape and size variations among MAGIC lines. The resulting principal components (PCs) were ranked according to the proportions of the total variance that each of them described (Figure 1).

In Leaf4, the PCA revealed that 90.92% of the variance in organ shape and size was attributed to two PCs (Figure 1A). Leaf4.PC1 of this model accounted for 76.84% of the total variance and affected the leaf size. Higher PC1 values corresponded to larger leaves, whereas lower values yielded smaller leaves. PC2 accounted for 14.08% of the variance and affected mainly shape. Lower values of PC2 yielded rounded leaves with a short petiole, whereas high values yielded elongated leaves with a long petiole. PC3 accounted for 3.30% of the variance and reflected the way the petiole twisted when the leaves were flattened. Its values were not significantly different between genotypes, and it was therefore excluded from further analysis.

In Leaf7, the PCA revealed that 95.25% of the variance in organ shape and size could be attributed to three PCs (Figure 1B). Leaf7.PC1 of this model accounted for 80.27% of the total variance and mostly affected size; however, there was also a minor effect on shape. Higher PC1 values corresponded to larger, more elongated leaves, whereas lower values yielded smaller and more rounded leaves. PC2 accounted for 11.42% of the variance and mostly affected the steepness of the transition from petiole to blade, with low values yielding a very gradual transition and high values yielding a long petiole with a steep transition. PC3 accounted for 3.56% of the variance and affected mainly the shape. Lower values of PC3 yielded more elongated and narrower leaves, whereas higher values of PC3 yielded more rounded and wider leaves.
In petals, the PCA revealed that 92.02% of the variance in organ shape and size could be attributed to two PCs (Figure 1C). The PC1 of this model accounted for 85.43% of the total variance and affected petal size. Higher PC1 values corresponded to larger petals, whereas lower values yielded smaller petals. PC2 accounted for 6.59% of the variance and affected mainly the shape. Low values of PC2 yielded elongated petals with a narrower shape, and high values of PC2 yielded rounded petals with a wider shape. PC3 accounted for 3.63% of the variance and was reflected in petal twisting when the petals were flattened. Its values were not significantly different between genotypes, and it was therefore excluded from further analysis.

The allometric variation captured by PCs reflected both genetic differences and environmental variations within the plant growth chamber in which plants were grown. Extensive phenotypic variation was observed for all traits measured among the MAGIC lines (Table 1). An estimate of the relative genetic contribution was made by comparing the variance of each PC among MAGIC lines (which was largely due to genetic differences) to that within each line (Supplementary Figure 2). Estimates from an average of approximately eight plants from each of the collected lines suggested that most of the variance (>60% for the PCs) had an underlying genetic basis (Table 1).

A correlation analysis between shape and size was also performed, and a number of significant pairwise correlations were observed (Figure 2). Leaf4.PC1 was significantly positively correlated with Leaf7.PC1, which represented the leaf size. Leaf4.PC2 was significantly correlated with Leaf7.PC2 and leaf7.PC3, which represented the leaf shape. Moreover, leaf shape and size showed significant correlations with petals. Petal.PC1 was significantly correlated with Leaf4.PC1 and Leaf7.PC1, which showed a negative size correlation between leaves and petals. Furthermore, both Leaf4.PC2 and Leaf7.PC2 were significantly positively correlated with Petal.PC1 and negatively correlated with Petal.PC2. The correlation between the leaf and petal allometry model indicated the genetic dependency and evolution correlation controlling leaf and petal allometry. Besides, a pairwise correlation analysis was performed between the life history traits and the leaf and petal allometry model (Figure 2). Leaf4.PC1 was correlated with rosette leaf number and stem height; additionally, Leaf4.PC2 was highly correlated with branch number and pod number; Leaf7.PC1 was correlated with days to bolting, days to flower and stem height; Leaf7.PC2 was highly positively correlated with days to bolting, days to flower, rosette leaf number, and branch number; Petal.PC1 was correlated with rosette leaf number and branch number; and Petal.PC2 was correlated with days to bolting and days to flower.

**QTLs accounted for leaf and petal allometry**

To examine the genetic basis for shape and size variation of leaves and petals along the PCs in the MAGIC lines, we treated each PC as a quantitative trait, whose variation frequency showed a normal distribution (Figure 2, Supplementary Figure 3) for QTL mapping. In QTL mapping of the MAGIC lines, the PCs for the leaf and petal allometry model and 1260 SNP markers among the 19 founder ecotypes were used. We then calculated a series of QTLs associated with the variance of leaf and petal shape and size (Figure 5, Supplementary Table 2, Supplementary Figure 4, 5 and 6). In the leaf model, the QTL analysis
for Leaf4.PC1 identified four QTLs located on chromosomes 1 and 3 and one QTL located on chromosome 2 for Leaf4.PC2. For Leaf7.PC1, five QTLs were observed on chromosome 3, one QTL was located on chromosome 2 for Leaf7.PC2, and four QTLs were located on chromosomes 1 and 2 for Leaf7.PC3. In the petal model, three QTLs were identified on chromosomes 1 and 4 in Petal.PC1, and nine QTLs were identified on chromosomes 1, 2, 3, and 5 in Petal.PC2.

After comparing the positions for all the QTLs identified, there was some QTL overlapping in the leaf and petal allometry model. The QTLs for PC2 of the leaf (Leaf4.PC2: LF4_2.1, Leaf7.PC2: LF7_2.1) and petal (Petal.PC2: PE_2.5) on chromosome 2 (~11 Mb) overlapped, and the alleles from the Ler-0 accession formed the most rounded leaves and petals with the widest shape (Supplementary Table 3). This QTL likely stemmed from the mutation of *ERECTA*, which is known to affect fruit length and is due to the allele from the *Ler-0* accession (Abraham et al., 2013). With the exception of the ER locus for leaf and petal shape, the QTLs LF7_1.1, LF7_1.2, LF7_1.3, LF7_1.4, and LF7_1.5 for Leaf7.PC1 on chromosome 3 overlapped with QTL PE_2.6 for Petal.PC2. Moreover, the QTLs LF7_1.3, LF7_1.4, and LF7_1.5 also overlapped with QTL PE_2.7 for Petal.PC2, whereas these QTLs all showed an uncorrelated allele effect distribution (Supplementary Table 3). For the fourth and seventh leaves, except for the overlapping ER locus (Leaf4.PC2: LF4_2.1, Leaf7.PC2: LF7_2.1) for PC2 described above, the QTLs LF4_1.3 and LF4_1.4 for Leaf4.PC1 overlapped with QTL LF7_1.6 for Leaf7.PC1 on chromosome 3 and showed the same allele effect distribution with a maximum value in the *Mt-0* accession and a minimum value in the *Can-0* accession (Supplementary Table 3). The overlapping QTLs might have explained the phenotypic correlation and indicated the correlated genetic modules for leaf and petal allometry in evolution.

**Candidate genes for leaf and petal allometry**

The genes that explain natural variations in leaf and petal allometry have remained largely unknown. To identify possible candidate genes, we searched for genes containing nonsynonymous SNPs unique to accession according to PC distribution among these accession alleles (Supplementary Table 3). Based on the resequencing and reannotation of the 19 parental accessions (Gan et al., 2011), we identified candidate genes with unique alleles referring to the maximal effects accession in the 95% confidence region (Supplementary Table 4). In the Leaf4 allometry model, the auxin receptor *TIR1*, brassinolide signalling regulator *BSL3*, and *TIR1*, contributing to flowering time repression, had allelic variation in the coding sequence unique to the accession. In the Leaf7 allometry model, hormonal-related genes, such as *SUA* (a suppressor of *abi3-5*), *ARGOS*, serine/threonine-protein kinase *PID2*, *BRI1 suppressor 1 (BSU1)-like 3*, and *ABI4* genes, had allelic variation in the coding sequence. Moreover, the flower time regulators *ELF3* and *ELF4*, the receptor kinase *ERECTA*, cell wall modification-related genes and some transcription factors conferred allelic variations unique to the maximal effects accession.

In the petal allometry model, 23 genes were identified with variations unique to the accession. Among these genes, *PTL* in Petal.PC2 encodes a trihelix transcription factor whose expression is limited to the margins of floral and vegetative organs. It is involved in limiting lateral growth of organs, and recessive
mutations have been found to be defective in organ initiation and orientation in the second whorl (Kaplan-Levy et al., 2014). The OFP13 in Petal.PC2 encodes a member of the plant-specific OVATE family of proteins. Members of this family have been shown to bind to KNOX and BELL-like TALE class homeodomain proteins and function as transcriptional repressors that suppress cell elongation (Wang et al., 2011). The SEU in Petal.PC1 encodes a transcriptional coregulator of AGAMOUS that coordinates with LEUNIG to repress AG in the outer floral whorls. Other genes, including the cell cyclin-related protein Cyclin A1;1, the protein kinase, the CYP family protein, the photoperiod-associated ELF6, and the transcription-related genes with nonsynonymous SNPs also contribute to petal PCs. The identified QTLs and candidate genes provided us with a valuable reference for insight into leaf and petal allometry.

The genetic basis for leaf and petal covariation in allometry models

Leaves and petals are homologous organs sharing mechanisms of developmental control, such that genes that act pleiotropically on both organ types might give rise to coordinating changes in shape or size. To examine the genetic basis for shape and size covariation between leaves and petals, the allometry model was also used. The petal and leaf modelled data sets obtained above were combined, which allowed overall trends to be identified. To ensure equal weighting of the data from different organs, the organ size for all plants was multiplied by a constant factor so that the variance in the Leaf4, Leaf7, and petal data sets was equal. The Leaf4, Leaf7, and petal data sets were then combined to create Leaf4-Petal and Leaf7-Petal data sets containing each plant from the MAGIC line groups. Additionally, a PCA was applied to the Leaf4-Petal and Leaf7-Petal data sets to detect correlated variation in the positions of points and to identify trends in shape and size variation between the two organs.

In the Leaf4-Petal model, PC1 accounted for 53.58% of the total variance, representing the negative size covariation between Leaf4 and petals. The higher the PC1 value, the larger the petal size, and the smaller the fourth leaf size. PC2 accounted for 30.26% of the total variance, representing the positive size covariation between the fourth leaf and petal. The higher the PC2 value, the larger the petal and leaf size. PC3 accounted for 5.92% of the total variance representing the positive shape (mainly in width) covariation between the fourth leaf and the petal. The higher the PC3 value, the more rounded the leaves and petal, and the shorter the petiole. PC4 accounted for 3.23% of the total variance representing the negative shape (mainly in width) covariation between the fourth leaf and the petal. The higher the value, the narrower the leaves, the longer the petiole, and the more rounded the petals were. The other PCs represented only one organ shape or size variance, so they were not considered for further analysis (Figure 3).

After QTL mapping in the MAGIC lines for the Leaf4-Petal model, three significant QTLs for PC1, one significant QTL for PC2, two significant QTLs for PC3, and six significant QTLs for PC4 were identified (Figure 5, Supplementary Table 5, Supplementary Figure 7). In each QTL, the candidate genes containing nonsynonymous SNPs unique to the maximal effects accession in the 95% confidence region were identified (Supplementary Tables 6 and 7). In PC1, there were five genes with the unique maximal effects accession allele, including the cell-proliferation-related genes, such as ARGOS, LOM2, and EXPB5. In PC3,
which represented the shape (mainly in width) covariation, four genes were identified: \textit{ARGOS}, \textit{FRS3}, \textit{BSL3}, and \textit{extensin proline-rich1}. In PC4, representing the negative shape (mainly in width) covariation, there were also four genes containing the unique accession allele. Among these genes, the \textit{CYCD2;1} gene acting on the G1 phase of the cell cycle to control the cell division rate in both the shoot and root meristems had an allele unique to the \textit{Hi-0} accession, and the \textit{PRX53} gene influencing cell elongation had an allele unique to the \textit{Po-0} accession.

Similar to the Leaf4-Petal model, in the Leaf7-Petal model, PC1 accounted for 68.58\% of the total variance, representing the negative size covariation between the seventh leaf and petal, whereas PC2 accounted for 22.51\% of the total variance, representing the positive size covariation between the seventh leaf and the petal and the seventh leaf shape variance. PC3 accounted for 2.84\% of the total variance, representing the positive shape (mainly in width) covariation, and PC4 accounted for 1.99\% of the total variance, representing the negative shape (mainly in width) covariation. The other PCs represented only one organ shape or size variance, so they were not considered for further analysis (Figure 4).

After QTL mapping in the MAGIC lines for the Leaf7-Petal model, two significant QTLs for PC3 and six significant QTLs for PC4 were identified, whereas no significant QTL was identified in PC1 and PC2 (Figure 5, Supplementary Table 5, Supplementary Figure 8). Moreover, candidate genes were also identified (Supplementary Tables 6 and 7). The QTL LF7PE_3.2 in PC3, which represented the positive shape (mainly in width) covariation between the seventh leaf and petal, had the most rounded leaf and petal in \textit{Ler-0} and the narrowest leaf and petal in the \textit{No-0} accession. In the 95\% confidence region, there were 34 genes conferring alleles unique to the \textit{Ler-0} or \textit{No-0} accession. Among these genes, the GRF gene \textit{AT2G22840}, pentatricopeptide repeat protein \textit{SLOW GROWTH1} (\textit{SLO1}), \textit{ORGAN BOUNDARY1} (\textit{OBO1}) and \textit{OVATE} family of protein OFP16 have been reported to affect organ shape or size (Kim et al., 2003; Sung et al., 2010; Cho et al., 2011; Wang et al., 2011). Furthermore, the cyclin-dependent kinase inhibitor \textit{KRP4} (Schiessl et al., 2014) and the serine/threonine-protein kinase \textit{PINOID} (\textit{PID}) are involved in the regulation of auxin signalling (Saini et al., 2017). \textit{Growth-regulating factor 3} (\textit{GRF3}), which regulates cell expansion in leaf and cotyledon tissues (Kim et al., 2003), as well as other genes associated with cell differentiation, cell expansion, cell wall modification, and flower time control genes, were also identified. The QTL LF7PE_4.4 in PC4, which represented the negative shape (mainly in width) covariation, had the narrowest leaves with the longest petiole and the most rounded petals in the \textit{Po-0} accession. There were three genes with an allele unique to the \textit{Po-0} accession, including \textit{DME}, a transcriptional activator involved in gene imprinting; \textit{peroxidase 2}, which influences cell elongation (Jin et al., 2011); and \textit{CYP712A2}, a member of \textit{CYP712A}.

**Discussion**

In this study, we defined a genetically controlled allometric space that captured most of the variation in leaf and petal shape and size among MAGIC lines. Among the loci identified, with the exception of the ER locus, the other QTLs were not identical to previously identified shape- and size-associated loci. Additionally, in these QTL confidence regions, many cell proliferation- and cell expansion-associated
genes were isolated with a unique allele according to the accession distribution. Furthermore, we checked the candidate gene expression data and compared their promoter sequences in the 19 founder accessions of MAGIC lines and found that two candidate genes, *ERECTA* (*AT2G26330*) and *AGO4* (*AT2G27040*), in the ER locus on chromosome 2 had specific variations in promoter sequences unique to the maximal effects of accession. In addition, the specific promoter sequence variation also changed the expression level significantly (Supplementary Figure 11). However, more work is needed to test these loci, such as by constructing a NIL population.

**The interaction of leaf and petal shape and size variation**

Interestingly, when we augmented the leaf and petal data to investigate the leaf and petal allometry covariation, the results showed that negatively correlated changes in leaf and petal size provided the largest component of allometric variation among MAGIC lines. The QTLs on chromosome 3 for Leaf4-Petal.PC1 overlapped with QTLs for Leaf4.PC1 with the smallest leaf size and the largest petal size in the *Can-0* accession allele. This indicated that the locus positively regulating fourth leaf size also negatively regulated petal size. The QTL on chromosome 2 for Leaf4-Petal.PC3 overlapped with Leaf4.PC2 and Petal.PC2 with the widest leaf and petal in the *Ler-0* accession allele, which indicated that the locus positively regulated both the leaf and petal width. The QTL on chromosome 2 for Leaf4-Petal.PC4 was identical to the QTL for Petal.PC2 with the same accession allele effects distribution. Moreover, the QTL on chromosome 5 was identical to the QTL for Petal.PC2 with uncorrelated allele effects distribution. In Leaf7-Petal.PC4, the QTL on chromosome 2 was identical to the QTL for Leaf7.PC2 and overlapped with the ER locus for Petal.PC2 with the maximum allele effects in the *Ler-0* accession. Two QTLs on chromosome 5 for Leaf7-Petal.PC4 overlapped with the QTL for Petal.PC2, and the overlapping QTLs LF7PE_4.6 and PE_2.9 both obtained the minimum values in the *Ws-2* accession. In addition to these overlapping QTLs with leaf or petal allometry, others did not overlap and may have been independent loci for leaf and petal covariation.

**Leaf and petal allometry coordinated with local adaption**

Additionally, life-history traits, such as days to bolting, days to flower, rosette leaf number, branch number, and stem height, were also measured in the MAGIC lines. Although many significant phenotype correlations were found (Figure 2), there were overlapping QTLs that could be used to explain the genetic correlation that was identified. After QTL mapping for these life-history traits (Figure 5, Supplementary Tables 8 and 9, Supplementary Figure 9), the QTLs for the life-history traits and for the leaf and petal allometry model were compared (Supplementary Figure 10). In Leaf4.PC1, QTL LF4_1.1 overlapped with one linked QTL, RN.6, for rosette leaf number and showed the same allele distribution with maximum values in the *Po-0* accession. In Leaf7.PC2, QTL LF7_2.1 on chromosome 2 (~11.2 Mb) overlapped with three linked QTLs (DF.4, DF.5, and DF.6) for days to flower, with the highest value found in the *Bur-0* accession allele. In Petal.PC1, two QTLs, PE_1.2 on chromosome 1 (~16.9 Mb) and PE_1.3 on chromosome 4 (~0.05 Mb), overlapped with the QTLs RN.1 and RN.9 separately for rosette leaf number with uncorrelated allele distribution. In Petal.PC2, there were four QTLs overlapping with the QTLs for
days to bolting. Among these QTLs, the QTLs PE_2.2 and PE_2.3 on chromosome 1 overlapped with QTL DB.1, with the maximum value in the Po-0 allele. Moreover, PE_2.9 on chromosome 5 overlapped with DB.6, with a maximum value in the Can-0 accession. Others showed uncorrelated allele distribution. There were also four QTLs for Petal.PC2 that overlapped with the QTLs for days to flower, and all showed uncorrelated allele distribution. The overlapping QTLs for leaf and petal allometry with life-history traits provided a genetic basis in the correlation analysis. This colocalization may have resulted from pleiotropy or tightly linked causal genes, which indicated genetic integration among all traits.

The advantage of using an allometry model was that the effect of each leaf and petal shape and size variation could be fully described quantitatively with a vector, and the allometric spaces could be captured based on PCs. Conversely, the model was restricted to 2D morphological traits and could neither capture 3D effects, such as curvature changes, nor capture the special shape of a leaf, such as a simple leaf with serrations. In addition, we only explored one environmental condition. Many environmental factors, such as abiotic stress, could have had an effect on leaf and petal shape and size. Analysis of MAGIC lines grown under different conditions could be an additional application of the method.

Conclusions

This is the first report on the genetic basis of allometry variation of leaves and petals and their interaction under the incorporated framework by using MAGIC lines. PCA for the MAGIC lines indicated that size variation was a major component of allometry variation and revealed negatively correlated changes in leaf and petal size. In this study, five QTLs for the fourth leaf, 11 QTLs for the seventh leaf, and 12 QTLs for petal size and shape were identified. These QTLs were not identical to those previously identified, with the exception of the ER locus. This indicated that the allometry variation was not simply the combination of organ width, length, and size. In addition, after QTL analysis of the leaf and petal integrated model, 12 QTLs were identified in association with the fourth leaf and petal allometry covariation, and eight QTLs were identified to be associated with the seventh leaf and petal allometry covariation. The QTL overlap explained the allometry correlation within different leaves and the homologous organs leaf and petal. However, some specific QTLs between Leaf4 and Leaf7 may explain the leaf allometry divergence, which may be associated with leaf developmental constraints. Additionally, the correlation of life history traits with leaf and petal allometry and the QTL overlap hinted at the genetic integration and the interaction of organ allometry with local adaptation.

Methods

Plant material and growth conditions

The large population of 527 RILs (Kover et al., 2009) was obtained from the University of Oxford in the United Kingdom and then propagated at Shandong Normal University in China. Seeds were sterilized for 10 minutes in 75% ethanol, washed in 95% ethanol four to six times, and then suspended in 0.1% agar. All lines were grown separately in 1/2 Murashige and Skoog medium. The seeds were
subsequently grown under the following conditions in a plant incubator (Percival Scientific, Inc): 22°C/18°C (Day/Night) and a long photoperiod of 16 hours/8 hours (Day/Night) after treatment for four days at 4°C in the dark for stratification. After they were grown in the medium for seven days, the seedlings were transplanted into soil when the true leaves could be seen. For each line, we planted eight seedlings, with four seedlings per pot, which were randomly assigned to a tray. The trays were rotated throughout the incubator every week.

Leaf and petal collection

After the first flower of the plant had opened, the fourth and seventh leaves of each plant were picked, flattened, and then glued onto paper for scanning to record the leaf shape. In total, the fourth leaves were obtained from 232 lines, and the seventh leaves were obtained from 215 lines (Supplementary Table 1). Because the leaves are more susceptible to environmental influences during growth, we calculated the average area of all obtained leaves in the same line and retained leaves with a difference in the range of +/- 20% for further analysis. To measure the shape of the petals, we picked and dissected the floral buds using a stereomicroscope at fully reflexed petal stage 13 when the buds had fully opened and the petals were visible and in anthesis. All four petals, four sepals, six stamens, and one pistil were removed, placed on 1% agar on a plate, and photographed with a Leica camera. Only buds between bud positions 4 and 10 on the main stem were used. Two flowers were dissected per plant, and each line we collected had at least four plants. Since some of these lines did not bloom properly under current planting conditions, we finally obtained petals from 345 lines for model construction (Supplementary Table 1).

Modelling in leaves and petals

To generate a parameterized space that captured variation in leaf and petal shape and size, the outlined organs were obtained for intercrossing populations of Arabidopsis derived from MAGIC lines. To investigate the allometric variation in Arabidopsis leaf and petal, plants were grown together in a glasshouse, and their fully expanded leaves and petals were flattened and imaged. For each independent line, digital images were taken from 8 independent, mature (after the first flower flowered) fourth and seventh true leaves. A digital image was made of a flattened leaf taken from the fourth and seventh nodes from the base of each plant, and 25 points were placed around the leaf outline using the leaf (Le) template. The resulting leaf shapes were aligned through translation and rotation (Procrustes alignment, Goodall, 1991) to generate a data set in which the outline of a leaf was represented by the Cartesian coordinates of its 25 points, each expressed in standard deviation from the mean position of the point within the collection of leaves. An equivalent procedure was used to generate a petal data set using a 25-point petal (Pe) template.

After all of the lines of the fourth and seventh leaves and the petals were prepared, the leaves were properly oriented (with the tip always pointing to the right and with good horizontality) using Photoshop CS5 software (Adobe). We used MATLAB R2007b (MathWorks) and the AAM Toolbox (Bensmihen et al., 2008) to construct the model of each individual leaf and petal separately. After placing the landmarks, the secondary landmark points were evenly spread along the leaf outline. This was achieved by taking
two primary points and fitting a spline between them using the secondary points, a spline being a special function that was defined piecewise by polynomials. The secondary points were then rearranged to be equidistant from each other along the spline. Models including the fourth leaf (Leaf4), the seventh leaf (Leaf7), and petals were separately generated using the AAM Toolbox (version 6.5). Points placed around each leaf and petal outline were plotted to show the pattern of allometry in the data set. Because the positions of the outline points were unlikely to change independent of one another, a PCA was used on the whole data set to identify trends in variation.

**Statistical analyses**

Broad sense heritability ($H^2$) for each trait was estimated as the ratio of the variance among lines to the total variance. To determine phenotype correlations between traits, pairwise Pearson correlations between the line means were calculated. QTL analyses were then performed using the R (version 3.4.4) software package HAPPY as described by Kover et al. (2009). Briefly, this approach used a hidden Markov model to make a multipoint probabilistic reconstruction of the genome of each MAGIC line as a mosaic of the founder haplotypes. Thus, at each marker, a probability of being derived from each of the parental accessions was assigned to each line, and our hypothesis that there would be no QTL was evaluated by fitting a fixed-effect linear model with up to 18 degrees of freedom. We performed QTL analysis for the line average of Leaf4, Leaf7, petal PCs, argument of Leaf4-Petal PCs, argument of Leaf7-Petal PCs, and life-history traits. Two QTLs located less than 1 Mb apart were considered overlapping QTLs reflecting genetic pleiotropy (Gnan et al., 2014).

**Abbreviations**

MAGIC: Multiparent advanced generation intercross

PCA: Principal component analysis

PCs: Principal components

QTL: Quantitative trait loci

**Declarations**

**Ethics approval and consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Availability of data and materials**
The raw data used for this research and the supplementary material are available in Figshare (https://figshare.com/s/90c637df9f8965f346c8). Dataset 1, which is used for PCA, contains all the cropped images and point models for each plant of the MAGIC lines. The values of each PC and the life history traits for each plant in the MAGIC lines are listed in Dataset 2. Dataset 3 is the record of correspondence between planting ID and the MAGIC line ID. File 1 contains the R source code files for QTL mapping in MAGIC lines.

**Competing interests**

The authors declare that they have no competing interests.

**Funding**

This work was supported by programmes from the National Natural Science Foundation of China (under grant nos. 31470286 and 31801387). The funding organizations played no role in the design of the study and the collection, analysis, and interpretation of data or in writing the manuscript.

**Authors' contributions**

SXY and XZF conceived the project and designed this work. XL, CXW and QS performed the experiments, and YHZ and XL analysed the data. SXY and XZF wrote the manuscript.

**Acknowledgements**

We are very grateful to Professor Enrico Coen of the John Innes Center for supplying seeds of the 527 MAGIC lines.

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Supplementary Material

Additional file 1: Figure S1-S11.

Figure S1. PCA was applied to the Leaf4, Leaf7, and petal data sets to identify trends in shape and size variations among MAGIC lines.

Figure S2. Range of PC values obtained for the leaf and petal allometric model.

Figure S3. Variations along each PC for Leaf4, Leaf7 and Petal allometry within the MAGIC lines.

Figure S4. QTL scan of the PCAs for the fourth leaf allometry among MAGIC lines.

Figure S5. QTL scan of the PCAs for the seventh leaf allometry among MAGIC lines.

Figure S6. QTL scan of the PCAs for petal allometry among MAGIC lines.

Figure S7. QTL scan of the PCAs for the fourth leaf and petal allometry covariation among MAGIC lines.

Figure S8. QTL scan of the PCAs for the seventh leaf and petal allometry covariation among MAGIC lines.

Figure S9. QTL scan of the life-history traits among MAGIC lines.

Figure S10. The genetic correlation between life-history traits and the allometry model.

Figure S11. The promoter sequence alignment and expression analysis of candidate genes.

Additional file 2: Table S1. The phenotype data for all traits used for QTL analysis.

Additional file 3: Table S2. Significant QTL detected for the leaf and petal allometry models.

Additional file 4: Table S3. The estimated value for each of the 19 parental alleles at each detected QTL for leaf and petal allometry models.

Additional file 5: Table S4. The candidate genes account for leaf and petal allometry variation.

Additional file 6: Table S5. Significant QTL detected for the leaf and petal allometry covariation.

Additional file 7: Table S6. The estimated value for each of the 19 parental alleles at each detected QTL for leaf and petal allometry covariation.

Additional file 8: Table S7. The candidate genes account for leaf and petal allometry covariation.

Additional file 9: Table S8. The significant QTL detected for life-history traits in MAGIC lines.
Additional file 10: Table S9. The estimated value for each of the 19 parental alleles at each detected QTL for life-history traits.

Tables

**Table 1. Phenotypic variation among MAGIC lines for leaf and petal allometry models**

| Trait      | Min     | Max     | Mean ± SD          | $H^2$ |
|------------|---------|---------|--------------------|-------|
| Leaf4.PC1  | -103.05 | 101.52  | 1.68 ± 31.9        | 0.83  |
| Leaf4.PC2  | -39.84  | 42.86   | -0.22 ± 13.4       | 0.67  |
| Leaf7.PC1  | -226.53 | 270.35  | -2.54E-09 ± 70.0   | 0.87  |
| Leaf7.PC2  | -81.51  | 68.12   | 1.08E-08 ± 26.4    | 0.77  |
| Leaf7.PC3  | -47.81  | 72.20   | 2.95E-09 ± 14.7    | 0.62  |
| Petal.PC1  | -86.65  | 56.59   | 0.01 ± 24.1        | 0.83  |
| Petal.PC2  | -20.83  | 30.18   | -0.001 ± 6.7       | 0.74  |

Minimum (Min) and maximum (Max) phenotypic values for each trait, as well as the phenotypic means plus or minus their standard deviation (SD) and their broad-sense heritability ($H^2$), are shown.