ORIGINAL ARTICLE



EXPANSIN15 is involved in flower and fruit development in Arabidopsis

Judith Jazmin Bernal-Gallardo¹ · Karla L. González-Aguilera¹ · Stefan de Folter¹

Received: 4 June 2023 / Accepted: 14 December 2023 © The Authors 2024

Abstract

Key message EXPANSIN15 is involved in petal cell morphology and size, the fusion of the medial tissues in the gynoecium and expansion of fruit valve cells. It genetically interacts with SPATULA and FRUITFULL.

Abstract Cell expansion is fundamental for the formation of plant tissues and organs, contributing to their final shape and size during development. To better understand this process in flower and fruit development, we have studied the *EXPANSIN15* (*EXPA15*) gene, which showed expression in petals and in the gynoecium. By analyzing *expa15* mutant alleles, we found that EXPA15 is involved in petal shape and size determination, by affecting cell morphology and number. EXPA15 also has a function in fruit size, by affecting cell size and number. Furthermore, EXPA15 promotes fusion of the medial tissues in the gynoecium. In addition, we observed genetic interactions with the transcription factors SPATULA (SPT) and FRUITFULL (FUL) in gynoecium medial tissue fusion, style and stigma development and fruit development in Arabidopsis. These findings contribute to the importance of *EXPANSINS* in floral and fruit development in Arabidopsis.

Keywords Expansin15 · Flower · Gynoecium · Fruit · Transcription factors · Petal

Introduction

The size and shape of plant organs are regulated by proliferation and subsequent expansion, contributing to cell morphology in each organ (Ramachandran et al. 2000; Marshall et al. 2012; Guerriero et al. 2014; D'Ario et al. 2021). Cell wall expansion is a process mediated by EXPANSIN proteins, these enzymes are plant cell wall-loosening proteins that loosen cell walls by weakening the binding of polysaccharide polymers to each other, contributing to cell enlargement and thus to cell growth and shape (Sampedro and Cosgrove 2005).

According to expression patterns in Arabidopsis and other species, it is known that EXPANSIN proteins are expressed in floral tissues, such as sepals, petals, stamens and carpels, as well as in floral meristems (Brummell et al. 1999b; Armezzani et al. 2018; Liu et al. 2021). Expansins

Communicated by Gabriela Pagnussat .

Published online: 29 January 2024

play an important role in the development of leaves, fruit, pollen tube and roots, as well as in defense against different stresses, in Arabidopsis and other species (Reinhardt et al. 1998; Brummell et al. 1999a; Chen and Bradford 2000; Cho and Cosgrove 2000; Wrobel and Yoder 2001; Pezzotti et al. 2002; Zenoni et al. 2004; Jones and McQueen-Mason 2004; Belfield et al. 2005; Balestrini et al. 2005; Giordano and Hirsch 2007; Tsuchiya et al. 2015; Muthusamy et al. 2020; Liu et al. 2021; Samalova et al. 2022, 2023). However, in Arabidopsis due to the redundancy of the EXPANSIN family, there are few examples of specific functions in Arabidopsis development (Sampedro and Cosgrove 2005; Marowa et al. 2016; Samalova et al. 2022).

On the other hand, specific transcription factor networks involved in Arabidopsis gynoecium and fruit development have been described (Chávez Montes et al. 2015; Herrera-Ubaldo and de Folter 2022; Herrera-Ubaldo et al. 2023). One of the transcription factors required for gynoecium development is SPATULA (SPT), which is known to be involved in the development of the medial tissues of the gynoecium, and the lack of its functions affects reproduction and consequently, fruit size (Alvarez and Smyth 1999; Heisler et al. 2001; Girin et al. 2011; Reymond et al. 2012; Reyes-Olalde et al. 2017a). Another transcription factor that affects fruit size is FRUITFULL (FUL) (Qing Gu et al. 1998;



Unidad de Genómica Avanzada (UGA-Langebio), Centro de Investigación y de Estudios Avanzados del Instituto Politécnico Nacional (CINVESTAV-IPN), 36824 Irapuato, GTO., Mexico

Ferrándiz et al. 2000). It has been reported that *SPT* through genetic interaction with *FUL*, participates also in fruit development (Groszmann et al. 2011), suggesting being involved in proliferation and expansion processes.

In this work, we focus on the enzyme encoding gene *EXPA15* in Arabidopsis flower and fruit development. In the flower, we found that *EXPA15* is involved in petal shape and size by affecting cell morphology, size and number. In addition, this enzyme is involved in fruit size by controlling expansion in fruit valve cells. Furthermore, we identified genetic interactions of *EXPA15* with the transcription factors *SPT* and *FUL*.

Results

Mutations in *EXPA15* affect petal cell morphology and number in Arabidopsis flowers

The enzyme encoding *EXPANSIN15* (*EXPA15*) gene has been described as an important developmental regulator in the meristem and is known to be expressed in the root (Armezzani et al. 2018; Samalova et al. 2023) but has not been described to play a role in floral development in Arabidopsis.

In this work, we analyzed the possible effects of the absence of EXPA15 in flower and fruit development. We observed that in the inflorescences of expa15-1, there is a malformation in the flower (Fig. 1A-C), in part also observed in two additional homozygous insertional mutants in the Col-0 background, expa15-2 and expa15-3 (Figs. S1, S2, S3). In detail, the petals of the null mutant expa15-1 mutant are narrower compared to the wild-type Ler (WT) (Fig. 1D). At the cellular level, scanning micrographs show that petal cells in WT, which are characterized by their conical shape, lost this morphology in the expa15-1 mutant, as the cells observed are flat and elongated (Fig. 1E-H). In addition, cell number is strongly reduced in the petal claw and blade (Fig. S4). Petal cell number is also reduced in the expa15-2 and expa15-3 mutants, though conical cell shape in petals is maintained (Figs. S2, S3, S4).

EXPA15 contributes to the development of the fruit by participating in cell expansion of the valves

Another phenotype observed in the *expa15* mutants is in fruit development. The *expa15-1* plants produce statistically significant shorter fruits (Fig. 2C), and moreover, the fruits have valves that are irregularly shaped, with bulging regions, something not observed in the WT (Fig. 2A, B). Furthermore, neither seed number nor seed area (Fig. 2D, E)

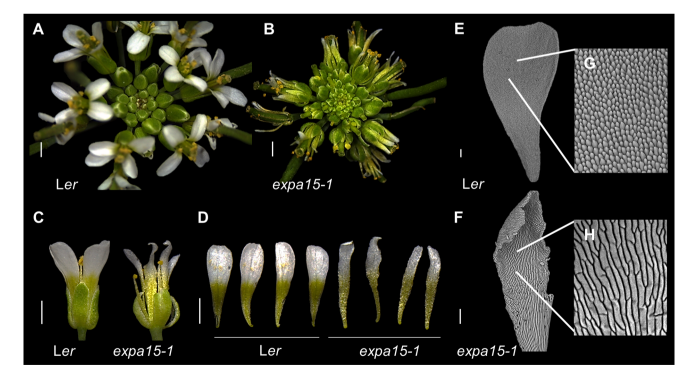


Fig. 1 *EXPA15* affects petal cell morphology.**A–B** Inflorescence of Ler and the *expa15-1* mutant. **C** Flower of wild-type Ler and *expa15-1*. **D** Narrow petals in *expa15-1* compared to Ler. **E–H**) Scanning

microscopy images of Ler and expa15-1 in adaxial part of petals. **G** and **H** are magnifications of **E** and **F**. Scale bars: 1 mm in **A-D**; 200 µm in **E**, **F**



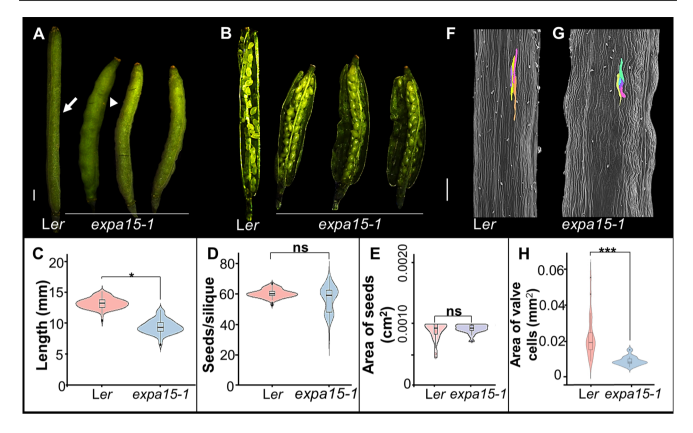


Fig. 2 The mutation in the *EXPA15* gene affects fruit length due to a decrease in the expansion of valve cells. **A–B** Images of fruits of Ler and *expa15-1*. **C–D** Analysis on fruits and seeds of Ler and *expa15-1*. **C** Fruit length. **D** Number of seeds per silique. **E** Area of seeds. **F–G** Scanning microscopy images of Ler and *expa15-1* fruits. **H** Analysis

of fruit valve cell area in Ler and expa15-1. Statistical analyses were performed using a Wilcoxon test in C, D, H and Student's t-test in E. n=80 in C-D, n=30 in E, n=5 in H: *p<0.01, ****p<0.0001, ns: not significant, p>0.05. Scale bars: 1 mm in A-B; 200 μ m in F, G

are affected in expa15-1 compared with WT. Probably, the bulging of the valves is caused by the seeds that are packed in a smaller volume, something similar has been observed in the mutant for the FRUITFULL (FUL) gene (Gu et al. 1998). To better understand the valve phenotype, we used scanning electron microscopy (SEM) and found that fruit valve cells are smaller in the expa15-1 mutant compared to WT, which is reflected in the decreased valve cell area (Fig. 2F-H). Similar alterations were noted in fruit length in expa15-2 and expa15-3. Notably, although fruit length is reduced in all three alleles due to reduced cell size, the number of valve cells is increased (Fig. 2, Figs. S2, S3, S4). However, this increased number of cells does not compensate for overall fruit length. In addition, in the expa15-2 and expa15-3 alleles, a reduction in seed number is observed (Figs. S2, S3). In summary, the results indicate that EXPA15 contributes to cell expansion in fruit valves.

The lack of EXPA15 can affect carpel fusion

It is worth mentioning that the patterning of most gynoecia in the *expa15-1* mutant is normal compared to WT, though

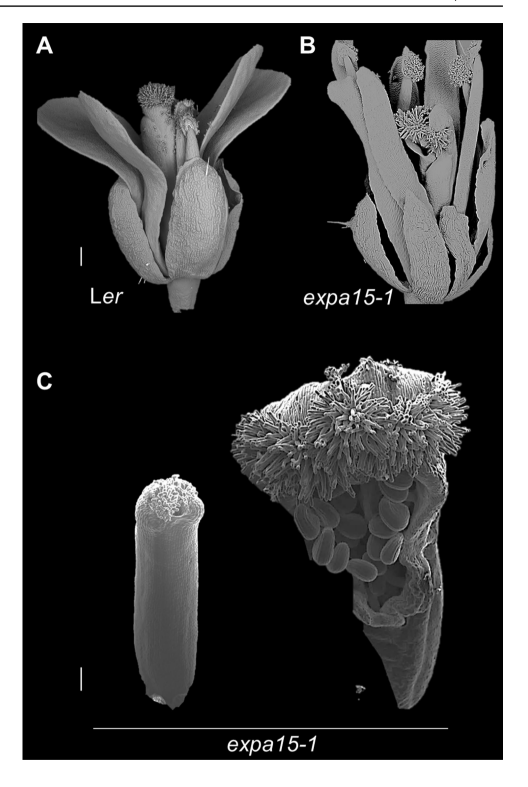
in a small frequency (around 5%), gynoecia with three valves have been observed (Fig. S5). In addition, we observed that some of the gynoecia of the *expa15-1* mutant showed affected carpel fusion (very low frequency; in around the first 8–10 flowers in the complete plant) (Fig. 3), which is a similar phenotype that has been reported for mutations in the *SPATULA* (*SPT*) gene (Alvarez and Smyth 1999; Heisler et al. 2001). This might suggest that there is a genetic interaction between *SPT* and *EXPA15*.

SPATULA and EXPA15 have a genetic interaction to promote carpel fusion

To investigate whether there is a genetic interaction between SPT and EXPA15 in gynoecium formation, we generated the $spt-2 \times expa15-1$ double mutant. Interestingly, we observed that some phenotypes in the $spt-2 \times expa15-1$ double mutant were additive (Fig. 4). In the $spt-2 \times expa15-1$ double mutant, the petal phenotype observed in the expa15-1 single mutant can still be observed (Fig. S6). In the spt-2 mutant, carpel fusion defects can be seen at the apical part of the gynoecium



Fig. 3 The *expa15-1* mutant is affected in carpel fusion. A Flower of WT. B Flower of *expa15-1* with unfused carpels. C Gynoecium of the *expa15-1* mutant with unfused carpels in the apical part, and exposed ovules can be seen. Scale bars: 200 µm in A–B; 100 µm in C



(Fig. 4C), and internally, septum fusion and transmitting tract differentiation do not occur, as has been reported (Fig. 4E) (Alvarez and Smyth 1999, 2002; Heisler et al. 2001). The unfused carpel phenotype seems to be synergistically enhanced in the double mutant, evident from the early stages of gynoecium development (Fig. 4D). Furthermore, style and stigma development defects are also

synergistically enhanced, so much that the stigmatic papillae cells are hardly observed. Interestingly, a phenotype not observed in the single mutants is that in the double mutant gynoecia, sometimes the replum continues to grow and is longer than the valves (Fig. 4D, top image). In transverse gynoecia sections, no obvious defects in medial tissue development were seen in the *expa15-1* single mutant



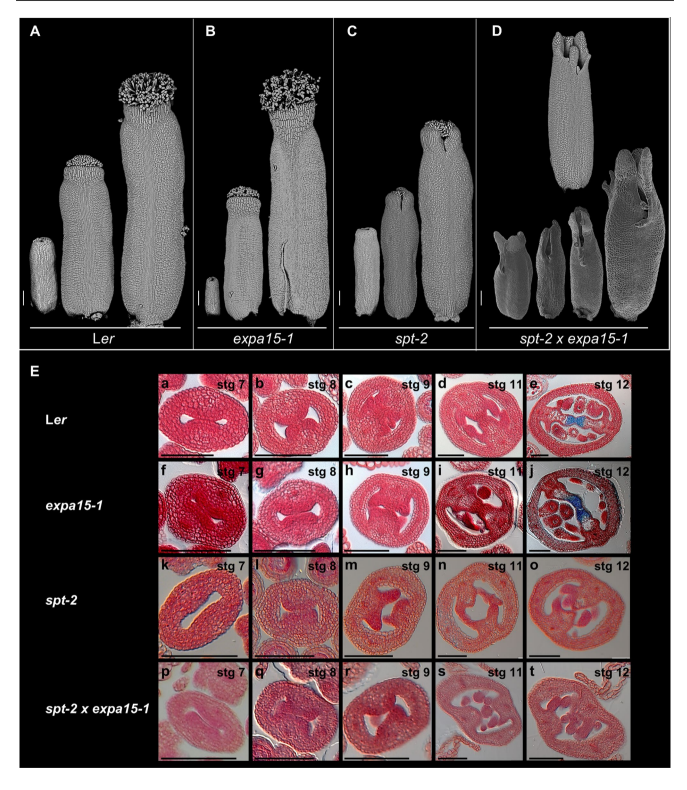


Fig. 4 The *spt expa15* double mutant shows developmental defects in gynoecium development. **A–D** Scanning microscopy images of gynoecium development in WT, *expa15-1*, *spt-2* and *spt-2 expa15-*

l double mutant. **E** Transverse gynoecia sections at different stages in WT, *expa15-1*, *spt-2* and *spt-2 expa15-1* mutant (**a–t**). Scale bars 100 μm in **A–D**; 10 μm in **E** (**a–t**)



(Fig. 4E). In the $spt-2 \times expa15-1$ double mutant, defects in medial tissue development and fusion are as in the spt-2 single mutant.

During fruit development, the fruits of the $spt-2 \times expa15-1$ double mutant do not develop seeds, in contrast to seed development in the single mutants (Fig. 5). Furthermore, the apical fusion defects seen in the gynoecium can still be seen in the fruit (Fig. 5). Different fruit morphologies were observed such as the development of an elongated split style in different degrees of severity, unfused carpels with a replum that continues to grow, as well as a large medial cleft, probably as a consequence of the synergistic effect of the single mutants.

In summary, the results suggest a genetic interaction between *SPT-EXPA15* is required for carpel fusion and coordination of apical gynoecium and fruit development. On the other hand, the results suggest that *EXPA15*

through an independent pathway contributes to petal cell expansion and valve cell expansion in the fruit.

SPT indirectly can affect EXPA15 expression during gynoecium development

To further understand the genetic relationship between *SPT* and *EXPA15*, we tested whether *EXPA15* expression could be affected by the *spt* mutation. First, it has already been described that *SPT* expression in the gynoecium is localized in the medial tissue during gynoecium development (Groszmann et al. 2010) (Fig. 6A–E). We studied the CSHL_ET6476 enhancer trap line, which carries a unique insertion of an enhancer trap (ET) transposable Ds element and a GUS reporter gene (Sundaresan et al. 1995). In this case, the line is a null mutant for *expa15* (Fig. S1) and serves as a reporter line for *EXPA15*. GUS expression in the homozygous line, reflecting *EXPA15* expression,

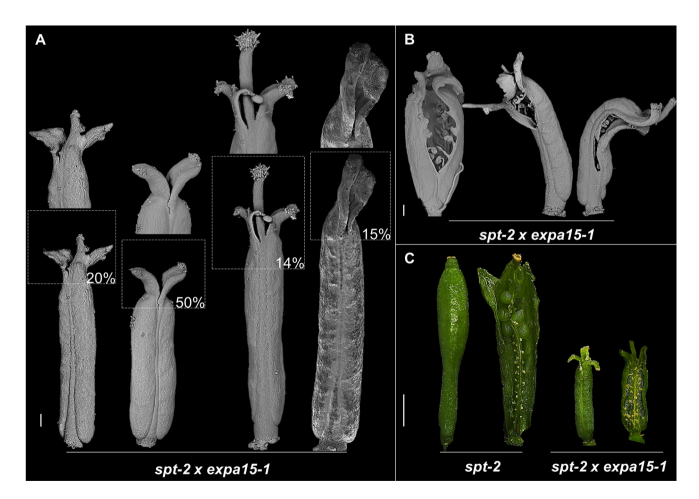


Fig. 5 The $spt-2 \times expa15-1$ double mutant produces a sterile fruit and an incorrect fusion in the apical part of the gynoecium. A Different fruits of the $spt-2 \times expa15-1$ double mutant with morphological phenotypes such as a split style, elongated replum formation and a reduced or lack of stigma formation. B Most severe phenotypes

seen in the *spt x expa15-1* double mutant; carpels largely unfused. **C** View of a fruit showing defects in medial tissue formation and seed-set in the *spt-2* single and *spt-2*×*expa15-1* double mutant. Scale bars: $200 \, \mu m$ in **A**, **B**; 1 mm in **C**



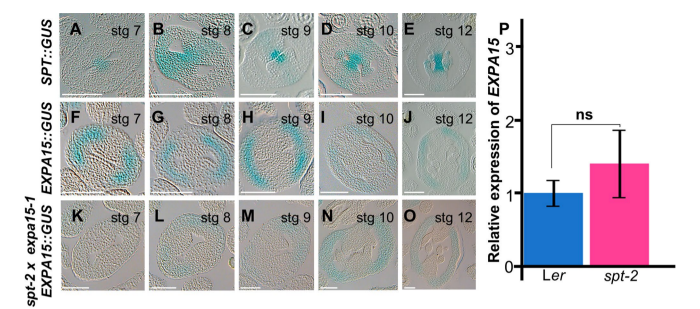


Fig. 6 SPT could regulate *EXPA15* in early stages of gynoecium development. **A–E** *SPT::GUS*, **F–J** *EXPA15::GUS* and **K–O** *EXPA15::GUS* spt-2×expa15-1 expression patterns in transverse gynoecia sections at different stages. **P** RT-qPCR expression data of

the *EXPA15* gene in inflorescences of Ler and spt-2. Student's t-test was used to evaluate significant differences between WT and spt-2. Significant values are indicated as follows: *p<0.05; ns: not significant, p>0.05. Scale bar 10 μ m

was observed very weakly at gynoecium stage 6-7 in the medial domain and strongly in the lateral domain. In later stages, strong GUS expression is observed specifically in the lateral domain of the gynoecium (Fig. 6F–J, Fig. S7). EXPA15 expression is limited to gynoecium development, since no GUS signal is observed during fruit development, only in the fruit pedicel (Fig. S7). Interestingly, EXPA15 expression in the medial domain at gynoecia stages 6 and 7 is almost absent in the $spt-2 \times expa15-1$ double homozygous mutant background, and a disperse GUS signal in the medial tissues at stage 8. As mentioned before, at later developmental stages, strong GUS signal is only observed in the lateral domain, however, in the double mutant background, the signal is decreased. Furthermore, it appears that this decrease in EXPA15 expression is affected from meristem formation onward, as almost no EXPA15 expression is observed in the double mutant (Fig. S8).

Using a complementary approach to find out whether SPT regulates *EXPA15* expression, we performed RT-qPCR analyses. However, when looking at *EXPA15* expression in inflorescence tissue of the *spt-2* mutant background, no statistically significant change was observed. These data suggest, together with the lack of a clear overlap in expression patterns between *SPT* and *EXPA15*, that the effect of SPT on *EXPA15* expression in the gynoecium is indirect.

FRUITFUL and EXPA15 have a genetic interaction to promote fruit valve elongation and style morphology

The above results show a genetic interaction between SPT and EXPA15 in carpel fusion and apical gynoecium development. However, considering that EXPA15 promotes cell elongation in fruit valves, as well is expressed in the valves, and expa15-1 mutant fruits have an appearance resembling ful-2 mutant fruits, we considered a possible relationship with the transcription factor FUL, which is one of the master regulators of fruit development (Gu et al. 1998). Therefore, we generated the ful- $2 \times expa15$ -1 double mutant, to find out if there is a genetic interaction between FUL and EXPA15. First, the ful-2 mutant is characterized by shorter valves and a zig-zag morphology of the replum, and depending on the allele, style morphology is affected (Gu et al. 1998; Ferrandiz et al. 2000). In particular, the short valve length phenotype is exacerbated in the $ful-2 \times expa15-1$ double mutant, as valve size is shorter than the single mutants (Fig. 7A, G). Interestingly, we observed that the fruit in the double mutant $ful-2 \times expa15-1$ developed a longer style than the ful-2 single mutant, something similar has been reported for other ful alleles (Fig. 7A) (Ferrandiz et al. 2000). It is worth mentioning that the cells of the double mutant ful- $2 \times expa15-1$ are similar to those of the ful-2 single mutant,



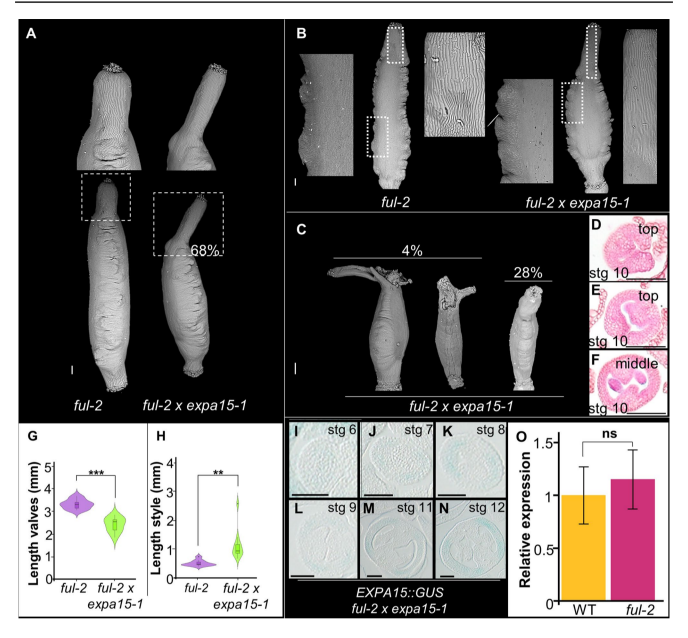


Fig. 7 EXPA15 and FUL contribute to valve and medial tissue development and EXPA15 contributes to style development in the fruit. A Scanning microscopy images of ful-2 and ful-2×expa15-1 double mutant fruits. B Magnifications of valve and style cells in ful-2and ful-2×expa15-1 double mutant fruits. C Severe phenotype in the ful-2×expa15-1 double mutant fruit. D-F Transverse gynoecia sections

of $ful-2 \times expa15-1$ affected in medial tissue fusion. **G**, **H** Quantification of valve and style length of ful-2 and $ful-2 \times expa15-1$ double mutant. Statistical analyses were performed using a Wilcoxon test in G and Student's t test in **H**, **O**. n = 10: ***, p < 0.001, **p < 0.01. Scale bars 200 μ m in **A–C**; 100 μ m in **D–F**, **I–N**

likewise, the cells of the style are more elongated, causing the style in the double mutant to be longer (Fig. 7A, B, H). In addition, we observed in a lower frequency fruit with an unfused apex, a seemingly one-side of a split style structure and reduced stigmatic tissue (Fig. 7C). The effect of the lack of fusion of the medial tissue can be observed from the formation of the gynoecium (Fig. 7D–F). Furthermore, looking at the pattern of EXPA15 expression in the $ful-2 \times expa15-1$ double mutant, based on the GUS

signal this expression is decreased in the lateral domain (Fig. 7I–N). In addition, we have performed RT-qPCR analysis to determine whether *EXPA15* expression is regulated by FUL (Fig. 7O). However, we did not observe a change in *EXPA15* expression in the *ful-2* mutant. These data confirm that there is a genetic interaction between *FUL-EXPA15* to promote fruit valve elongation and style development, but we cannot conclude that FUL regulates *EXPA15* during gynoecium development.



Discussion

EXPA15 is an enzyme that contributes to cell morphology in petals

Cell expansion contributes to the shape and size of tissues and organs in floral organs and fruit (Alvarez-Buylla et al. 2010; Marshall et al. 2012; Ripoll et al. 2019; Herrera-Ubaldo and de Folter 2022). One of the enzyme groups that are involved in cell expansion or elongation are expansins (Samalova et al. 2022). In this work, we focused on the study of the expansin encoding gene *EXPA15*.

In general, in most flowering species, petals are characterized by having typical conical cells on their surface, though some variations can exist (Whitney et al. 2011). Based on our analyses, EXPA15 contributes to the morphology of petal cells, since in the expa15-1 null mutant (Ler background) petal cells lost their typical conical morphology, being observed elongated and flat. However, their differentiation seems to be unaffected since the blade of the petals remain white, characteristic of the Arabidopsis petal (Irish 2008). In two additional alleles (expa15-2 and expa15-3 in the Col-0 background), overall petal shape and size was also affected. All three alleles displayed a reduced number of cells in the petal. Notably, the alleles in the Col-0 background maintained the conical cell shape. In Petunia, it has been reported that the expansin PhEXP1 is involved in petal size by controlling expansion in petal cells, however, no overall morphological changes were observed in cell shape when PhEXP1 was downregulated (Zenoni et al. 2004). In conclusion, results support that in Arabidopsis, EXPA15 functions in petal shape and size by controlling cell elongation and cell number, though apparently there is an accession related effect.

SPATULA and EXPA15 together participate in carpel fusion in Arabidopsis

On the other hand, the Arabidopsis gynoecium is characterized by a fusion of two carpels at early stages of development. Cell proliferation and expansion is key for medial domain tissues of the gynoecium to fuse and subsequently differentiate into septum tissue, transmitting tract and subsequent formation of a style and stigma (Bowman et al. 1999; Alvarez-Buylla et al. 2010; Reyes-Olalde et al. 2013). Expansion during gynoecium development has been described to be a stage and tissue-specific phenomenon during development (Gómez-Felipe et al. 2023). Interestingly, in *expa15-1* mutants, some of the gynoecia showed incorrect carpel fusion as seen in the *spt* mutant (Alvarez

and Smyth 1999, 2002; Heisler et al. 2001), and this effect is enhanced in the $spt-2 \times expa15-1$ double mutant.

Based on the localization of *SPT* and *EXPA15* expression, in the gynoecium there is no or hardly any overlap, meaning that the effect of SPT on *EXPA15* is likely indirect. Only in the floral meristem, there is overlap in gene expression, though further studies would be necessary to explore if there is a direct interaction.

On the other hand, the phenotype of a split style and absence of stigma in the double mutant $spt-2 \times expa15-1$ has been observed in other double mutants of SPT and its target genes HOMEOBOX ARABIDOPSIS THALIANA 3 (HAT3) and ARABIDOPSIS THALIANA HOMEOBOX 4 (ATHB4) that contribute to the apical radialization of the style (Reymond et al. 2012; Carabelli et al. 2021), as well as in mutants of transcription factors of the NGATHA family (Trigueros et al. 2009), affecting the signaling or biosynthesis of auxins and cytokinins. It is known that EXPA15 can be regulated by type B Arabidopsis Response Regulators (type B ARRs) of the cytokinin signaling pathway, since its promoter contains DNA binding sites of this family of transcription factors (Samalova et al. 2023). In addition, SPT is known to contribute to cytokinin signaling by regulating ARR1 and ARR12 (Reyes-Olalde et al. 2017a, b), which could be the regulatory pathway of SPT on EXPA15 and contribute to medial tissue fusion and gynoecium apical tissue development.

A genetic interaction between FUL and EXPA15 affects fruit development

The genetic interaction between EXPA15 and FUL in fruit development is reflected in the additive effect of valve size reduction in the $ful-2 \times expa15-1$ double mutant. However, style elongation seems to be dependent on the participation of this enzyme, variations in style elongation have also been observed independently in other ful mutant alleles (Ferrándiz et al. 2000). Furthermore, lack of medial tissue fusion and a split style have also been observed in the $spt-2 \times ful-2$ double mutant (Groszmann et al. 2010), reflecting the role of EXPA15 together with SPT and FUL in medial tissue formation.

Although, it is unclear by RT-qPCR whether FUL regulates *EXPA15*, it cannot be ruled out that there is a relation with FUL, as we observed a decrease of *EXPA15* signal in the background of the *ful-2*×*expa15-1* double mutant in the lateral tissue of the gynoecium.

To conclude, according to our results, the regulation of *EXPA15* appears to be complex, as it might depend on the stage and tissue in which *EXPA15*-mediated expansion is required. This complexity of expansion regulation has been described, variable outcome of the function of an EXPANSIN has been observed depending on the amount of EXPANSIN present, i.e., despite cell expansion, at high



EXPANSIN concentration, cell expansion is reduced (Choi et al. 2003; Goh et al. 2014).

Nevertheless, this work contributes to the knowledge of the importance of cell expansion in floral tissues and fruit. One of the observed phenotypes is fruit size, which is an agronomic trait that can be altered without affecting the number of seeds. The modulation and specific localization of EXPANSINS could contribute to an interesting agronomic phenotype (Marowa et al. 2016; Cosgrove 2021). Future work will be necessary to better understand how these EXPANSIN-type enzymes contribute to reproductive tissue and organ development in the plant.

Materials and methods

Plant material and growth conditions

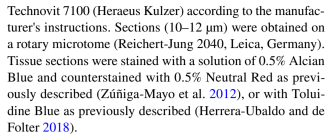
Arabidopsis thaliana plants used in this study was the expa15-1 mutant obtained from Jan Traas (Armezzani et al. 2018; CSHL_ET6476; ABRC CS25610), expa15-2 (ABRC GABI_556F03, expa15-3 (ABRC CS921359), spt-2 (Alvarez & Smyth 1999), pSPT-6253:GUS (Groszmann et al. 2010), ful-2 (Ferrandiz et al., 2000), expa15 and spt-2 are in the Ler background, and ful-2, expa15-2, expa15-3 in the Col-0 background and the wild type accessions Col-0 and Ler. The $spt-2 \times expa15-1$ double mutant was generated by crossing spt-2 with expa15-1, and the $spt-2 \times expa15-1$ double mutant was generated by crossing $spt-2 \times expa15-1$. Double homozygous plants $spt-2 \times expa15-1$.

Scanning electron microscopy

During reproductive development, the different lines were scanned using a Zeiss EVO40 environmental scanning electron microscope (Carl Zeiss; Oberkochen, Germany) with 25 kV beam, and the signal was collected using the SE or the BSD detector. Each plant tissue was collected and directly observed in the microscope.

Histology and microscopy analyses

For thin tissue section analysis, inflorescences of *expa15*, *spt-2*, *spt expa15* and Ler were collected, and the tissue was fixed in FAE solution (3.7% formaldehyde, 5% glacial acetic acid and 50% ethanol) with vacuum (15 min, 4 °C), and afterward incubated for 60 min at room temperature. The material was rinsed with 70% ethanol and incubated overnight at 4 °C in 70% ethanol, followed by dehydration in a series of ethanol dilutions (70, 85, 95 and 100% ethanol) for 60 min each. Inflorescences were embedded in



The *expa15-1* mutant was generated with an enhancer trap construct that contains a Ds transposon carrying a glucuronidase reporter gene (Sundaresan et al. 1995). We used the *expa15-1* line as a GUS reporter line to reflect the *EXPA15* gene expression. The inflorescences were collected and stained as previously described (Marsch-Martínez et al. 2014), with the following modifications: 30 min of vacuum and after 1.5 h in substrate at 37 °C, we proceeded with the dehydration with ethanol. The GUS-stained inflorescences were fixed, dehydrated as described above and embedded in Technovit 7100; 10–12 μm thick sections were analyzed. Subsequently, the samples were observed using a Nomarski Leica DM4000 microscope with DIC function.

For statistical analysis, data were first tested for normality using the Shapiro–Wilk test. Then, means were compared pair-wise using either Student's t-test, Wilcoxon test or one-way ANOVA *t* test. All calculations were performed in R (R Core Team 2022).

Quantitative real-time RT-PCR

For RT-qPCR analysis, the spt-2, ful-2 and Ler lines were collected inflorescences with floral buds only. Three biological replicates were sampled. After collection, total RNA was extracted using the Direct-zol RNA Mini Prep Plus Kit (Zymo Research, USA). Reverse transcription and amplification were performed in triplicate with a SyGreen 1-Step Go Hi-ROX qPCR kit (PCR BIOSYSTEMS, USA). RTqPCR was performed using a real-time Open qPCR machine (Model: A1005, CHAIBIO, USA). Expression levels of target genes were normalized with TUA2. Data were analyzed using the $2^{-\Delta\Delta CT}$ method (Livak and Schmittgen 2001). The following primers were used: EXPA15 (AT2G03090) F:5'-CTTCTGTAGGAAACAGGGACAAC-3' and R: 5'- CCT CCGCTTCATCATTTCGATC-3', and TUA2 (AT1G50010) F: 5'-GGTTCCAGGTTTGTCACTCGTT-3' and R: 5'-CCG AGAAGGTAAGCATCATGCG-3'.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00497-023-00493-4.

Acknowledgements We thank Victor M. Zuñiga-Mayo and Erik Cruz-Valderrama for discussions. We thank the Consejo Nacional de Humanidades, Ciencias y Tecnologías (CONAHCYT) for



a PhD fellowship to JJB-G. This work in the SdF laboratory was financed by the CONAHCYT grants FC-2015-2/1061 and CB-2017-2018-A1-S-10126.

Author Contributions JJBG and SdF conceived and designed research. JJBG conducted most experiments. KLGA performed the RT-qPCR experiments. JJBG, KLGA and SdF analyzed data. JJBG and SdF wrote the manuscript. All authors read and approved the manuscript.

Declarations

Conflict of interest The authors declare that they have no competing interests

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

- Alvarez J, Smyth DR (1999) CRABS CLAW and SPATULA, two Arabidopsis genes that control carpel development in parallel with AGAMOUS. Development 126:2377–2386
- Alvarez J, Smyth DR (2002) CRABS CLAW and SPATULA genes regulate growth and pattern formation during gynoecium development in Arabidopsis thaliana. Int J Plant Sci 163:17–41. https:// doi.org/10.1086/324178
- Alvarez-Buylla ER, Benítez M, Corvera-Poiré A et al (2010) Flower Development. Arabidopsis Book 8:e0127. https://doi.org/10.1199/tab.0127
- Armezzani A, Abad U, Ali O et al (2018) Transcriptional induction of cell wall remodelling genes is coupled to microtubule-driven growth isotropy at the shoot apex in Arabidopsis. Development. https://doi.org/10.1242/dev.162255
- Balestrini R, Cosgrove DJ, Bonfante P (2005) Differential location of α-expansin proteins during the accommodation of root cells to an arbuscular mycorrhizal fungus. Planta 220:889–899. https://doi. org/10.1007/S00425-004-1431-2
- Belfield EJ, Ruperti B, Roberts JA, McQueen-Mason S (2005) Changes in expansin activity and gene expression during ethylene-promoted leaflet abscission in Sambucus nigra. J Exp Bot 56:817– 823. https://doi.org/10.1093/JXB/ERI076
- Bowman JL, Baum SF, Eshed Y et al (1999) 4 Molecular genetics of gynoecium development in Arabidopsis. Curr Top Dev Biol 45:155–205. https://doi.org/10.1016/S0070-2153(08)60316-6
- Brummell DA, Harpster MH, Civello PM et al (1999a) Modification of expansin protein abundance in tomato fruit alters softening and cell wall polymer metabolism during ripening. Plant Cell 11:2203–2216. https://doi.org/10.1105/TPC.11.11.2203
- Brummell DA, Harpster MH, Dunsmuir P (1999b) Differential expression of expansin gene family members during growth and ripening of tomato fruit. Plant Mol Biol 39:161–169

- Carabelli M, Turchi L, Morelli G et al (2021) Coordination of biradialto-radial symmetry and tissue polarity by HD-ZIP II proteins. Nat Commun. https://doi.org/10.1038/s41467-021-24550-6
- Chávez Montes RA, Herrera-Ubaldo H, Serwatowska J, de Folter S (2015) Towards a comprehensive and dynamic gynoecium gene regulatory network. Curr Plant Biol 3–4:3–12. https://doi.org/10.1016/j.cpb.2015.08.002
- Chen F, Bradford KJ (2000) Expression of an expansin is associated with endosperm weakening during tomato seed germination. Plant Physiol 124:1265. https://doi.org/10.1104/PP.124.3.1265
- Cho HT, Cosgrove DJ (2000) Altered expression of expansin modulates leaf growth and pedicel abscission in Arabidopsis thaliana. Proc Natl Acad Sci U S A 97:9783–9788. https://doi.org/10.1073/PNAS 160276997
- Choi D, Lee Y, Cho HT, Kende H (2003) Regulation of expansin gene expression affects growth and development in transgenic rice plants. Plant Cell 15:1386. https://doi.org/10.1105/TPC.011965
- Cosgrove DJ (2021) Expanding wheat yields with expansin. New Phytol 230:403–405. https://doi.org/10.1111/NPH.17245
- D'Ario M, Tavares R, Schiessl K et al (2021) Cell size controlled in plants using DNA content as an internal scale. Science 372:1176–1181. https://doi.org/10.1126/science.abb4348
- Ferrándiz C, Gu Q, Martienssen R, Yanofsky MF (2000) Redundant regulation of meristem identity and plant architecture by FRUITFULL, APETALA1 and CAULIFLOWER. Development 127:725–734. https://doi.org/10.1242/DEV.127.4.725
- Giordano W, Hirsch AM (2007) The Expression of MaEXP1, a melilotus alba expansin gene, is upregulated during the sweet-clover-sinorhizobium meliloti interaction. MPMI 17:613–622. https://doi.org/10.1094/MPMI.2004.17.6.613
- Girin T, Paicu T, Stephenson P et al (2011) INDEHISCENT and SPATULA interact to specify carpel and valve margin tissue and thus promote seed dispersal in Arabidopsis. Plant Cell 23:3641–3653. https://doi.org/10.1105/tpc.111.090944
- Goh HH, Sloan J, Malinowski R, Fleming A (2014) Variable expansin expression in Arabidopsis leads to different growth responses. J Plant Physiol 171:329–339. https://doi.org/10.1016/j.jplph.2013.09.009
- Gómez-Felipe A, Marconi M, Branchini E et al (2023) Competing differentiation gradients coordinate fruit morphogenesis. BioRxiv. https://doi.org/10.1101/2023.01.19.524793
- Groszmann M, Bylstra Y, Lampugnani ER, Smyth DR (2010) Regulation of tissue-specific expression of SPATULA, a bHLH gene involved in carpel development, seedling germination, and lateral organ growth in Arabidopsis. J Exp Bot 61:1495–1508. https://doi.org/10.1093/jxb/erq015
- Groszmann M, Paicu T, Alvarez JP et al (2011) SPATULA and ALC-ATRAZ, are partially redundant, functionally diverging bHLH genes required for Arabidopsis gynoecium and fruit development. Plant J 68:816–829. https://doi.org/10.1111/j.1365-313X. 2011.04732.x
- Gu Q, Ferrándiz C, Yanofsky MF, Martienssen R (1998) The FRUIT-FULL MADS-box gene mediates cell differentiation during Arabidopsis fruit development. Development 125:1509–1517
- Guerriero G, Hausman J-F, Cai G (2014) No stress! relax! mechanisms governing growth and shape in plant cells. Int J Mol Sci 15:5094–5114. https://doi.org/10.3390/ijms15035094
- Heisler MG, Atkinson A, Bylstra YH et al (2001) SPATULA, a gene that controls development of carpel margin tissues in Arabidopsis, encodes a bHLH protein. Development 128:1089–1098. https://doi.org/10.1105/tpc.9.10.1859
- Herrera-Ubaldo H, de Folter S (2018) Exploring cell wall composition and modifications during the development of the gynoecium medial domain in Arabidopsis. Front Plant Sci 9:454. https://doi.org/10.3389/fpls.2018.00454



- Herrera-Ubaldo H, Campos SE, López-Gómez P et al (2023) The protein-protein interaction landscape of transcription factors during gynoecium development in Arabidopsis. Mol Plant 16:260–278. https://doi.org/10.1016/J.MOLP.2022.09.004
- Herrera-Ubaldo H, de Folter S (2022) Gynoecium and fruit development in Arabidopsis. Development (cambridge). https://doi.org/10.1242/DEV.200120/274550
- Irish VF (2008) The Arabidopsis petal: a model for plant organogenesis. Trends Plant Sci 13:430–436. https://doi.org/10.1016/j.tplants.2008.05.006
- Jones L, McQueen-Mason S (2004) A role for expansins in dehydration and rehydration of the resurrection plant Craterostigma plantagineum. FEBS Lett 559:61–65. https://doi.org/10.1016/S0014-5793(04)00023-7
- Liu W, Xu L, Lin H, Cao J (2021) Two Expansin Genes, AtEXPA4 and AtEXPB5, are redundantly required for pollen tube growth and AtEXPA4 is involved in primary root elongation in Arabidopsis thaliana. Genes (basel) 12:1–16. https://doi.org/10.3390/ GENES12020249
- Livak KJ, Schmittgen TD (2001) Analysis of relative gene expression data using real-time quantitative PCR and the 2 C T method. Methods 25:402–408. https://doi.org/10.1006/meth.2001.1262
- Marowa P, Ding A, Kong Y (2016) Expansins: roles in plant growth and potential applications in crop improvement. Plant Cell Rep. https://doi.org/10.1007/S00299-016-1948-4
- Marsch-Martínez N, Zúñiga-Mayo VM, Herrera-Ubaldo H et al (2014) The NTT transcription factor promotes replum development in Arabidopsis fruits. Plant J 80:69–81. https://doi.org/10.1111/tpj. 12617
- Marshall WF, Young KD, Swaffer M et al (2012) What determines cell size? BMC Biol 10:1–22. https://doi.org/10.1186/1741-7007-10-101
- Muthusamy M, Kim JA, Jeong MJ, Lee SI (2020) Blue and red light upregulate α -expansin 1 (EXPA1) in transgenic Brassica rapa and its overexpression promotes leaf and root growth in Arabidopsis. Plant Growth Regul 91:75–87. https://doi.org/10.1007/S10725-020-00588-2
- Pezzotti M, Feron R, Mariani C (2002) Pollination modulates expression of the PPAL gene, a pistil-specific β-expansin. Plant Mol Biol 49:187–197. https://doi.org/10.1023/A:1014962923278
- R Core Team (2022) R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. https://www.R-project.org/
- Ramachandran S, Christensen HEM, Ishimaru Y et al (2000) Profilin plays a role in cell elongation, cell shape maintenance, and flowering in Arabidopsis. Plant Physiol 124:1637–1647. https://doi.org/ 10.1104/pp.124.4.1637
- Reinhardt D, Wittwer F, Mandel T, Kuhlemeier C (1998) Localized upregulation of a new expansin gene predicts the site of leaf formation in the tomato meristem. Plant Cell 10:1427–2143
- Reyes-Olalde JI, Zuñiga-Mayo VM, Chávez Montes RA et al (2013) Inside the gynoecium: at the carpel margin. Trends Plant Sci 18:644–655. https://doi.org/10.1016/j.tplants.2013.08.002
- Reyes-Olalde JI, Zúñiga-Mayo VM, Marsch-Martínez N, de Folter S (2017a) Synergistic relationship between auxin and cytokinin in the ovary and the participation of the transcription factor SPAT-ULA. Plant Signal Behav 12:e1376158. https://doi.org/10.1080/ 15592324.2017.1376158

- Reyes-Olalde JI, Zúñiga-Mayo VM, Serwatowska J et al (2017b) The bHLH transcription factor SPATULA enables cytokinin signaling, and both activate auxin biosynthesis and transport genes at the medial domain of the gynoecium. PLoSGenet. https://doi.org/10.1371/journal.pgen.1006726
- Reymond MC, Brunoud G, Chauvet A et al (2012) A light-regulated genetic module was recruited to carpel development in Arabidopsis following a structural change to SPATULA. Plant Cell 24:2812–2825. https://doi.org/10.1105/tpc.112.097915
- Ripoll JJ, Zhu M, Brocke S et al (2019) Growth dynamics of the Arabidopsis fruit is mediated by cell expansion. Proc Natl Acad Sci U S A 116:25333–25342. https://doi.org/10.1073/PNAS.1914096116
- Samalova M, Gahurova E, Hejatko J (2022) Expansin-mediated developmental and adaptive responses: a matter of cell wall biomechanics? Quant Plant Biol 3(e11):1–14. https://doi.org/10.1017/OPB.2022.6
- Samalova M, Melnikava A, Elsayad K et al (2023) Hormone-regulated expansins: expression, localization, and cell wall biomechanics in Arabidopsis root growth. Plant Physiol 00:1–20. https://doi.org/10.1093/plphys/kiad228
- Sampedro J, Cosgrove DJ (2005) The expansin superfamily. Genome Biol 2005(6):242. https://doi.org/10.1186/gb-2005-6-12-242
- Sundaresan V, Springer P, Volpe T et al (1995) Patterns of gene action in plant development revealed by enhancer trap and gene trap transposable elements. Genes Dev 9:1797–1810
- Trigueros M, Navarrete-Gómez M, Sato S et al (2009) The NGATHA genes direct style development in the arabidopsis gynoecium. Plant Cell 21:1394. https://doi.org/10.1105/TPC.109.065508
- Tsuchiya M, Satoh S, Iwai H (2015) Distribution of XTH, expansin, and secondary-wall-related CesA in floral and fruit abscission zones during fruit development in tomato (Solanum lycopersicum). Front Plant Sci 6:1–9. https://doi.org/10.3389/fpls.2015.
- Whitney HM, Bennett KMV, Dorling M et al (2011) Why do so many petals have conical epidermal cells? Ann Bot 108:609–616
- Wrobel RL, Yoder JI (2001) Differential RNA expression of α-expansin gene family members in the parasitic angiosperm Triphysaria versicolor (Scrophulariaceae). Gene 266:85–93. https://doi.org/10.1016/S0378-1119(01)00376-6
- Zenoni S, Reale L, Tornielli GB et al (2004) Downregulation of the petunia hybrida α-expansin gene PhEXP1 reduces the amount of crystalline cellulose in cell walls and leads to phenotypic changes in petal limbs. Plant Cell 16:295. https://doi.org/10.1105/TPC. 018705
- Zúñiga-Mayo VM, Marsch-Martínez N, de Folter S (2012) JAIBA, a class-II HD-ZIP transcription factor involved in the regulation of meristematic activity, and important for correct gynoecium and fruit development in Arabidopsis. Plant J 71:314–326. https://doi.org/10.1111/j.1365-313X.2012.04990.x

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

