

AUXIN RESPONSE FACTOR8 Is a Negative Regulator of Fruit Initiation in *Arabidopsis*^W

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Fruit and seed formation in plants is normally initiated after pollination and fertilization, and, in the absence of fertilization, flowers senesce. In the *Arabidopsis thaliana* mutant *fruit without fertilization*, a mutation in *AUXIN RESPONSE FACTOR8* (*ARF8*) results in the uncoupling of fruit development from pollination and fertilization and gives rise to seedless (parthenocarpic) fruit. Parthenocarpy was confirmed in two additional recessive alleles and was caused by mutations within the coding region of *ARF8*. Genetic experiments indicate that *ARF8* acts as an inhibitor to stop further carpel development in the absence of fertilization and the generation of signals required to initiate fruit and seed development. Expression of *ARF8* was found to be regulated at multiple levels, and transcriptional autoregulation of *ARF8* was observed. Analysis of plants transformed with a transcriptional $P_{ARF8}:\beta$ -glucuronidase (*GUS*) construct or a translational *ARF8:GUS* fusion construct displayed distinct developmental regulation of the reporter in floral tissues involved in pollination and fertilization and in the carpel wall. After fertilization, the level of *GUS* activity declined in the developing seed, while in unfertilized ovules that are destined to senesce, *ARF8:GUS* expression spread throughout the ovule. This is consistent with a proposed role for *ARF8* in restricting signal transduction processes in ovules and growth in pistils until the fruit initiation cue.

INTRODUCTION

Fruit development and seed set in flowering plants normally occur in a coordinated manner following pollination of the stigma and subsequent double fertilization in the ovule of the flower (Gillaspy et al., 1993). When the egg and central cell of the female gametophyte are not fused with sperm cells, they remain in a quiescent state and eventually degrade as the flower undergoes senescence (O'Neill and Nadeau, 1997). This has led to the interpretation that signaling processes are required to activate development of the fertilization products leading to the initiation of seed and fruit development (Raghavan, 2003).

Various phytohormones, including gibberellins, cytokinin, and auxin, are involved in signaling processes following pollination and fertilization as a prerequisite for further growth and development of seeds and fruits (Nitsch, 1952, 1970; Coombe, 1960; Garcia-Martinez and Hedden, 1997; Fos et al., 2000, 2001). Developing seeds appear to be essential for fruit growth and development because they are sources of phytohormones, and there may be a requirement for phytohormones continuously throughout seed and fruit formation (Nitsch, 1970; Eeuwens and

Schwabe, 1975; Archbold and Dennis, 1985; Talon et al., 1990a; Garcia-Martinez et al., 1991; Ben-Cheikh et al., 1997; Swain et al., 1997; Ozga et al., 2002).

Fruit development can be uncoupled from fertilization and seed development, and the fruits, termed parthenocarpic, are seedless (Talon et al., 1992; Fos and Nuez, 1996; Robinson and Reiners, 1999; Varoquaux et al., 2002). Parthenocarpy has a genetic basis (Pike and Peterson, 1969; Lin et al., 1984; de Menezes et al., 2005) and has been exploited by farmers and plant breeders for the production of seedless fruits (Sykes and Lewis, 1996). Elevated endogenous phytohormone levels have been observed during parthenocarpic fruit set (George et al., 1984; Talon et al., 1990b, 1992), suggesting that increased supply of phytohormones to fruits from sources other than seeds may be sufficient to induce fruit growth. Accordingly, parthenocarpy can be induced in *Arabidopsis thaliana* and in diverse agricultural species by the exogenous application of auxins, cytokinins, or gibberellins (Gillaspy et al., 1993; Vivian-Smith and Koltunow, 1999) or by expression of auxin biosynthesis genes in ovaries and ovules (Rotino et al., 1997; Carmi et al., 2003; Mezzetti et al., 2004). Nonetheless, the molecular events directly involved in the initiation of fruit development and their link to plant hormone signal transduction processes remain unknown.

We have isolated the parthenocarpic *fruit without fertilization* (*fwf*) mutant from *Arabidopsis* in which fruit development and growth are uncoupled from pollination and fertilization events (Vivian-Smith et al., 2001). When fertilization is prevented in *fwf* by removal of floral organs surrounding the carpel, a seedless, dehiscent fruit or silique develops. In this article, we show that the *fwf* mutant contains a lesion in *AUXIN RESPONSE FACTOR8* (*ARF8*), a member of the *ARF* transcription factor family. Although *ARF* transcription factors, including *ARF8*, have been

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the focus of many earlier molecular studies, the characterization of the *fwf* mutant allele of *ARF8* together with other T-DNA insertion alleles demonstrates that *ARF8* is an important regulator of fruit initiation and that the disruption of its normal function induces parthenocarpy in *Arabidopsis*. The involvement of *ARF8* potentially provides a molecular link between the process of fruit initiation and the auxin signal transduction pathway.

RESULTS

fwf Contains a Lesion in *ARF8*

Unfertilized *Arabidopsis* carpels elongate slightly by cellular expansion (Figure 1), but they do not form dehiscent siliques, which is a defining feature of parthenocarpy in *fwf* (Vivian-Smith et al., 2001). *fwf* was originally characterized in the Landsberg *erecta* (*Ler*) background and was mapped to chromosome five (Vivian-Smith et al., 2001). Further fine mapping (see Methods) positioned *fwf* in a genomic interval of ~110 kb between the genetic markers *PhyC* and *SO191* (Figures 2A and 2B). Twenty-four annotated genes were present in this region (*Arabidopsis* Genome Initiative, 2000; Figure 2C), and the *ARF8* gene was a candidate based on the phenotypic data previously obtained (Vivian-Smith et al., 2001; Vivian-Smith, 2001). *ARF8* was amplified by PCR from both mutant and wild-type plants in the *Ler* background and sequenced. We identified a transition from G to A in the predicted translation initiation codon in the PCR product isolated from *fwf* plants, which converted the predicted start Met ATG to ATA. This base substitution created a cleaved-amplified polymorphic sequence (CAPS) marker (see Methods; Konieczny and Ausubel, 1993) that fully correlated with the *fwf* phenotype in segregating populations (Vivian-Smith, 2001).

Other Mutations in *ARF8* Can Induce Parthenocarpy

Although other mutations in *ARF8* have been identified and published, none have been reported to show parthenocarpic fruit

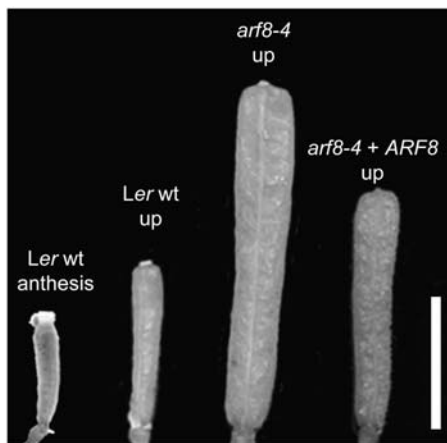


Figure 1. Pistils from Emasculated Flowers.

Comparison of wild-type anthesis carpels and unpollinated (up) pistils from *Ler*, *arf8-4*, and *arf8-4+ARF8* plants 7 d after emasculating. Bar = 3 mm.

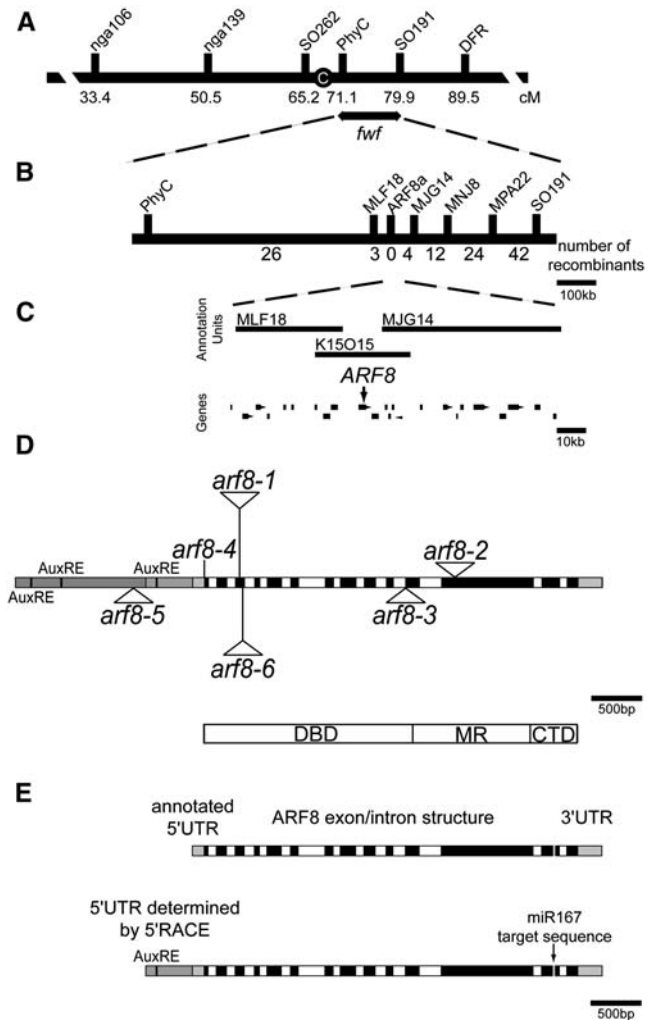


Figure 2. Physical and Genetic Mapping of *fwf*.

(A) Genetic map of chromosome 5 showing genetic markers used in cloning *fwf* and the map position of *fwf*. C, centromere; cM, centiMorgan (B) Physical map position of *fwf* with the number of recombinants between the markers indicated below.

(C) BAC vectors spanning the *FWF* region and the annotated genes delineated within this region. The position of *ARF8* is indicated by the arrow.

(D) Diagram of the *ARF8* gene with annotation of mutant alleles used in this study. T-DNA insertion lines are indicated by triangular insertions and the *arf8-4* allele as a line, as it is an ethyl methanesulfonate mutation. Alleles and their ecotypes examined in this study were *arf8-1* (*Ws*; Tian et al., 2004), *arf8-4* (*Ler*; Vivian-Smith et al., 2001), *arf8-5* (*Col*; Alonso et al., 2003), and *arf8-6* (*Col*; <http://www.hort.wisc.edu/krysan/DS-Lox/>). Gray boxes indicate the promoter region and 3' untranslated region (UTR). Three putative auxin response elements (AuxREs) are found in the promoter region. Black boxes specify exons, and white boxes indicate introns. A schematic representation of the *ARF8* protein and its domains is represented below. DBD, DNA binding domain; MR, middle region; CTD, C-terminal domain.

(E) Structure of the *ARF8* gene. Top: *ARF8* gene with short 5'UTR as annotated in the database. Bottom: *ARF8* gene with long 5'UTR as determined by 5'-RACE. The positions of the putative AuxRE present in the promoter region and of the microRNA target sequence (miR167) present in exon 13 are indicated.

development. *arf8-1* (Figure 2D) exhibits altered hypocotyl elongation (Tian et al., 2004), while a phenotype for *arf8-2* has not been described in detail so far (Okushima et al., 2005). The *arf8-3* mutant has delayed stamen development and decreased fecundity (Nagpal et al., 2005). We will henceforth refer to *fwf* as *arf8-4* (Figure 2D).

The flowers of the *arf8-1* mutant together with two previously uncharacterized insertional mutants *arf8-5* (SALK_049954) and *arf8-6* (WiscDsLox324F09) (Figure 2D) were emasculated to test for parthenocarpy. Silique development in emasculated *arf8* mutant flowers was compared with emasculated flowers from the corresponding wild-type ecotype background as controls because there are differences in the degree of carpel expansion in unfertilized wild-type flowers from different ecotypes (Vivian-Smith and Koltunow, 1999; Vivian-Smith et al., 2001).

Floral emasculation of plants homozygous for *arf8-1* and *arf8-6* produced dehiscent, parthenocarpic siliques (Table 1). Both of these recessive alleles (data not shown) contain a T-DNA insertion that disrupts the coding region (Figure 2D). F1 plants derived from crosses between *arf8-4* in a near isogenic Columbia (Col) ecotype background (*arf8-4NIL*) and *arf8-6* showed that these mutations were allelic because parthenocarpy was observed in plants trans-heterozygous for both mutations (data not shown).

By contrast, emasculation of flowers from homozygous *arf8-5* plants, which contain a T-DNA insertion 942 bp upstream of the *ARF8* coding region, resulted in nonparthenocarpic, indehiscent pistils (Table 1). Furthermore, *arf8-5* plants did not show the other phenotypes found in *arf8-1*, *arf8-4*, and *arf8-6*, including reduced seed set in proximal silique regions, broader silique shape, and

precocious carpel growth prior to flower opening (Vivian-Smith et al., 2001; data not shown). Collectively, these data suggest that parthenocarpic fruit development is caused by loss-of-function mutations due to disruptions within the coding region of *ARF8*.

The Parthenocarpic Phenotype in *arf8-4* Is Partially Rescued by an *ARF8* Genomic Fragment

Complementation of the *arf8-4* allele (Ler background) was attempted by inserting a wild-type *ARF8* genomic fragment (as shown in Figure 2D) by plant transformation. Nine independent lines containing one to three copies of the introduced gene were recovered and analyzed. Both the introduced wild-type and endogenous mutant gene were expressed in all nine lines because a mixture of both mutant and wild-type transcript was detected using the CAPS marker following RT-PCR assays (data not shown). All homozygous transgenic lines produced shorter siliques following floral emasculation when compared with the *arf8-4* mutant (Figure 1, Table 1). However, these shorter fruit were genuine siliques because they dehisced (Table 1). Moreover, some of the other features associated with *arf8-4* mutants, including reduced seed set in proximal silique regions, broad silique shape, and precocious growth of carpels prior to floral bud opening, were also generally retained in these lines. This indicated that only partial complementation had been achieved. At least two possibilities may account for this result. The introduced genomic fragment may be lacking key sequence information, or the *arf8-4* mutant allele is generating an inhibitory or competitive effect at the RNA and/or protein level.

Table 1. Analysis of Elongation, Dehiscence, and *ARF8* Expression Ratios in Lines Used in This Study

Line	Silique Length ^a	Elongation ^b	Dehiscence ^b	mRNA Expression Ratio ^c
Ler	4.5 mm ± 0.5 mm	–	–	1.0
<i>arf8-4</i>	7.5 mm ± 0.8 mm	+++	++	1.6 ± 0.2
Col	3.7 mm ± 0.5 mm	–	–	1.0
<i>arf8-4NIL</i>	5.9 mm ± 0.7 mm	++	++	1.3 ± 0.1
<i>arf8-5</i>	4.0 mm ± 0.6 mm	+	–	2.4 ± 0.2
<i>arf8-6</i>	5.2 mm ± 0.2 mm	++	++	0.6 ± 0.2
Ws	3.8 mm ± 0.3 mm	–	–	1.0
<i>arf8-1</i>	5.0 mm ± 0.1 mm	++	++	0.2 ± 0.1
<i>arf8-4</i> + <i>ARF8</i> #1	5.4 mm ± 0.8 mm	+	++	4.1 ± 1.2
<i>arf8-4</i> + <i>ARF8</i> #2	5.1 mm ± 0.6 mm	+	++	1.0 ± 0.1
<i>arf8-4</i> + <i>ARF8</i> #3	4.3 mm ± 0.7 mm	–	++	ND
<i>arf8-4</i> + <i>ARF8</i> #4	5.6 mm ± 0.6 mm	++	++	1.2 ± 0.2
<i>arf8-4</i> + <i>ARF8</i> #5	5.0 mm ± 0.7 mm	+	++	3.0 ± 0.2
<i>arf8-4</i> + <i>ARF8</i> #6	4.9 mm ± 0.6 mm	+	++	ND
<i>arf8-4</i> + <i>ARF8</i> #7	5.4 mm ± 0.6 mm	+	++	5.9 ± 0.3
<i>arf8-4</i> + <i>ARF8</i> #8	5.7 mm ± 0.5 mm	++	++	ND
<i>arf8-4</i> + <i>ARF8</i> #9	6.0 mm ± 0.8 mm	++	++	ND
Ler + <i>ARF8</i> #1	3.8 mm ± 0.7 mm	–	–	ND
Ler + <i>ARF8</i> #2	4.3 mm ± 0.4 mm	–	–	ND
Ler + <i>ARF8</i> #3	4.6 mm ± 0.4 mm	–	–	ND

^a Pistil lengths measured 7 d after emasculation (±SD; a minimum of 40 flowers was emasculated and measured for each line).

^b –, no elongation/dehiscence; +, some elongation; +++/++, very good/good elongation/dehiscence.

^c The expression ratio of *ARF8* mRNA was determined from flowers at anthesis. The ratio for each line is determined in relation to the expression in the corresponding wild-type line, which was set to 1.0 as the reference point. ND, not determined.

The *ARF8* Transcript Is Expressed in a Range of Tissues, and mRNA Levels Increase in the *arf8-4* Allele

RNA gel blot analysis of both wild-type and *arf8-4* plants detected mRNA of equal size in rosette and cauline leaves, inflorescence stems, flowers, and siliques. The size of the *ARF8* mRNA transcript found in all wild-type and *arf8-4* plant tissues examined was 3.4 kb in length (Figure 3). This was longer than the predicted 2.8-kb transcripts described in annotated databases. Rapid amplification of cDNA ends (5'-RACE) performed on RNA isolated from *arf8-4* and wild-type siliques and subsequent sequencing of the PCR products confirmed the presence of the mutation in the expressed RNA from *arf8-4* plant tissue and revealed that the mRNA in wild-type and *arf8-4* mutants extends a total of 627 bases 5' to the predicted translation initiation codon (Figure 2E), accounting for the observed 3.4-kb size. Thus, a stable full-length mRNA containing a mutation in the predicted translation start site is produced in *arf8-4* plants.

In wild-type plants, RNA gel blot analysis detected the highest levels of steady state *ARF8* mRNA in flowers at anthesis (Figure 3). In *arf8-4*, mRNA levels were higher in most examined tissues compared with wild-type *ARF8* levels (Figure 3). Quantitative real-time PCR (qRT-PCR) showed that *arf8-4* levels in anthesis flowers were approximately twofold higher than levels of *ARF8* mRNA in wild-type anthesis flowers (see Supplemental Table 1 online). The higher level of *arf8-4* mRNA persisted in the developing seeded siliques of *arf8-4* plants for the 2 d after fertilization where embryos were at early to mid globular stages, respectively (Figure 3).

In situ hybridization analysis showed the expression of *ARF8* was unevenly distributed in cells of a particular tissue. Expression varied with respect to the stage of flower development and silique growth (Figures 4A to 4D). Comparable expression patterns were found throughout flower development in both *Ler* and *arf8-4*. *ARF8* mRNA was detected in sepals, anthers, and carpels prior to anthesis in both *Ler* and *arf8-4* (Figures 4A and 4C). At anthesis, expression was strong within the mesocarp layers of the fruit (Figures 4B and 4D), and *ARF8* mRNA was also detected

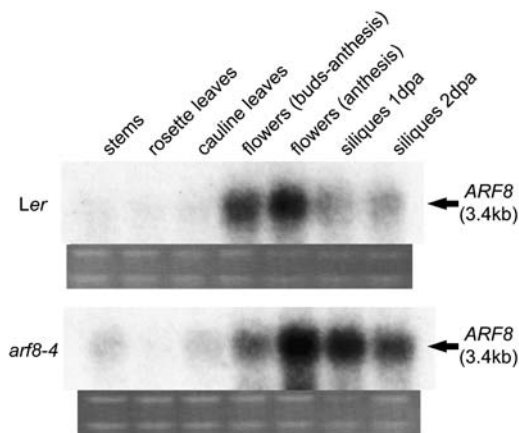


Figure 3. RNA Gel Blot Analysis.

Comparison of *ARF8* mRNA expression in various tissues from *Ler* and *arf8-4* plants. dpa, days postanthesis.

in the septum of the pistil and in the funiculi and integuments of both *Ler* and *arf8-4* ovules (Figures 4B and 4D). After fertilization, *ARF8* transcripts remained present in the integuments of both wild-type and *arf8-4* ovules 1 to 2 d after fertilization (see Supplemental Figure 1 online). *ARF8* mRNA was evident in the mesocarp of the fruit and the carpel septum during growth and was detectable 3 d after fertilization in both *Ler* and *arf8-4* (see Supplemental Figure 1 online). In situ hybridization is not reliable enough to use for the quantification of twofold changes in mRNA level, but we can conclude that the cellular distribution of both mutant and wild-type *ARF8* mRNA was diverse and spatially comparable at the light microscopy level in examined tissues.

ARF8 Shows Transcriptional Autoregulation

The increased levels of *arf8-4* mRNA in various tissues was further investigated by transforming *Ler* and *arf8-4* plants with a transcriptional fusion of the *ARF8* promoter fused to the β -glucuronidase (*GUS*) gene ($P_{ARF8}:GUS$; Jefferson et al., 1987). Three independent, homozygous lines containing $P_{ARF8}:GUS$ in each background were examined. Histochemical staining showed that the introduced transcriptional fusion was expressed at lower levels in all *Ler* tissues compared with transgenic *arf8-4* tissues at the same stage, although the spatial expression pattern was conserved. This was most obvious in seedlings (Figures 4E and 4F) but was also clearly evident in flowers and developing siliques. Two weeks after germination, *arf8-4* seedlings exhibited strong *GUS* activity throughout the hypocotyl, the inflorescence meristem, and the cotyledon margins. The true leaves had high *GUS* levels around the veins, the leaf margins, and the trichomes, and the root hairs were also intensely stained (Figure 4F). The *GUS* activity in transgenic *Ler* plants was significantly lower but showed the same pattern (Figure 4E). These observations are consistent with *arf8-4* mutants having an impaired ability to downregulate expression levels from the *ARF8* promoter in the transcriptional *GUS* fusion. Therefore, in wild-type plants, *ARF8* appears to be involved either directly or indirectly in the modulation of transcriptional expression from its own promoter.

To investigate *ARF8* regulation further, we examined mRNA levels in the nonparthenocarpic *arf8-5* allele (Col background) containing a T-DNA insertion between two putative auxin response elements in the predicted *ARF8* promoter (Figure 2D). We detected higher levels of *ARF8* mRNA in *arf8-5* flowers compared with that in wild-type Col-1 flowers of the same stage using qRT-PCR. The levels observed in *arf8-5* were also higher than those found in the *arf8-4* allele in the Col near isogenic line (*arf8-4NIL*; Table 1). The increased *ARF8* expression in the *arf8-5* plants suggests that the normal levels of expression might be altered by the presence of the T-DNA insertion within the promoter possibly influencing self-regulation.

$P_{ARF8}:GUS$ Expression during Parthenocarpic and Fertilization-Induced Silique Development in *arf8-4* and Wild-Type Plants

The expression patterns of the $P_{ARF8}:GUS$ construct were compared in flowers of wild-type and *arf8-4* mutant plants

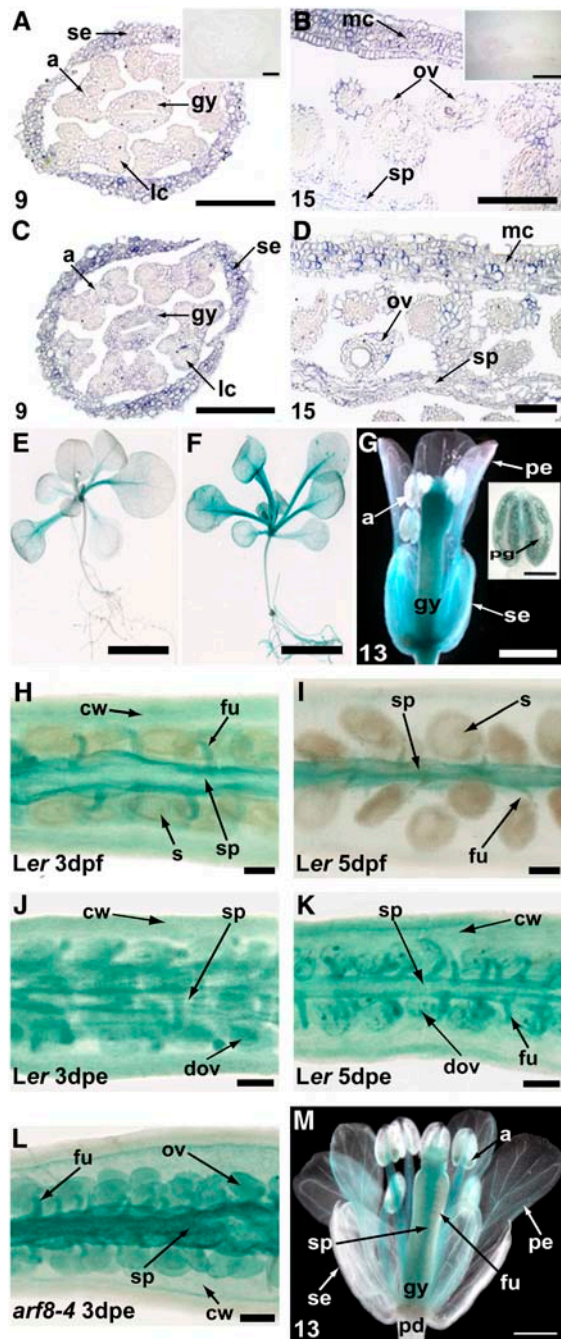


Figure 4. Expression of *ARF8*.

(A) to (D) In situ hybridizations on thin sections of developing *Ler* [(A) and (B)] and *arf8-4* [(C) and (D)] flowers [(A) and (C)] and siliques [(B) and (D)] hybridized with antisense-labeled *ARF8* RNA. Insets [(A) and (B)] show control hybridizations with sense-labeled *ARF8* RNA. The numbers at the bottom left indicate the stages of flower development (Smyth et al., 1990). Bars = 100 μ m.

(E) and (F) *ARF8* is involved in the modulation of *P_{ARF8}:GUS* expression. Seedlings of *Ler* (E) and *arf8-4* (F) plants transformed with the *P_{ARF8}:GUS* construct at 15 d after germination. Bars = 1 mm.

transformed with the *P_{ARF8}:GUS* construct. Throughout flower and fruit development, GUS activity was detected in a similar pattern to the mRNA distribution found in the in situ hybridization experiments. GUS activity was strong in carpels and pollen grains of *Ler* plants at anthesis, with comparably weaker activity in sepals and petals (Figure 4G). Further comparison of GUS activity in flowers after fertilization and after emasculation showed that 3 d after fertilization, GUS expression was visible within the carpel wall, the carpel septum, and the funiculi but not in the seeds (Figure 4H). Five days after fertilization, expression became restricted to the septum and the base of the funiculi (Figure 4I). After emasculation, there were changes in the observed expression pattern. Three days after emasculation of transgenic *Ler* plants, GUS activity was evident throughout the ovule in addition to the septum and carpel wall tissue (Figure 4J). Five days after emasculation, GUS activity had decreased in the carpel wall and in the septum but remained strong in the funiculi with some patchy GUS activity within the female gametophyte (Figure 4K).

In *arf8-4* mutant plants containing *P_{ARF8}:GUS*, the GUS activity in the flowers at anthesis and in the developing seeds after fertilization was spatially similar to that in pollinated *Ler* plants (see Supplemental Figure 1 online). However, staining was stronger and was detected in higher levels over the 3 d following fertilization, consistent with the data obtained by RNA gel blot analysis (Figure 3). In parthenocarpic siliques of *arf8-4* plants containing *P_{ARF8}:GUS*, GUS activity was observed 3 d after emasculation in the carpel walls, the septum, the funiculi, and within the ovules (Figure 4L). GUS activity became restricted to the septum and the base of the funiculi at 7 d after emasculation (see Supplemental Figure 1 online), coincident with the onset of ovule senescence. The restricted expression of the *P_{ARF8}:GUS* marker in the septum and funiculi during parthenocarpic silique formation in *arf8-4* resembles that found in fertilization-induced *Ler* siliques and is distinct from that in *Ler* pistils undergoing senescence.

***ARF8:GUS* Expression Reveals Additional Levels of Regulation**

A translational gene fusion was made with the wild-type *ARF8* genomic fragment (shown in Figure 2D) and the *GUS* gene generating *ARF8:GUS*. *Ler* plants were transformed with the construct, and 14 independent lines were recovered and analyzed for GUS activity. Even though the introduced translational

(G) *Ler* flower transformed with *P_{ARF8}:GUS* at anthesis (bar = 500 μ m). Inset shows an anther with stained pollen (bar = 50 μ m).

(H) to (L) *P_{ARF8}:GUS* expression in parthenocarpic pistils resembles that of fertilization-induced silique development. *Ler* silique 3 d after fertilization (H); *Ler* silique 5 d after fertilization (I); *Ler* silique 3 d after emasculation (J); *Ler* silique 5 d after emasculation (K); *arf8-4* silique 3 d after emasculation (L). Bars = 50 μ m.

(M) *Ler* flower transformed with *ARF8:GUS* at anthesis. Bar = 500 μ m. a, anther; cw, carpel wall; dov, degenerating ovule; dpe, days post-emasculum; dpf, days postfertilization; fu, funiculus; gy, gynoecium; ov, ovule; pd, pedicel; pe, petal; pg, pollen grain; s, sepal; sp, septum; lc, locule; mc, mesocarp.

ARF8:GUS fusion contained the same *ARF8* promoter fragment used in *P_{ARF8}:GUS*, a more restricted pattern of expression was observed in flowers and siliques of transgenic wild-type plants at anthesis because expression was not evident in sepals or the pedicel (cf. Figures 4G and 4M).

Analysis of the expression of *ARF8:GUS* during floral development showed that it was first detected in flowers at stage 10 (Smyth et al., 1990) in the stigma (Figure 5A). During flower stages 11 and 12, GUS staining was also evident in the carpel septum, the anther filaments, and then in developing pollen at the tetrad stage after meiosis (Figure 5B). At pollination (flower stage 13), *ARF8:GUS* expression was evident in the ovule funiculi, the carpel wall, and in the petals (Figure 5C). After fertilization (Figure 5D, stage 15), the level of GUS activity declined and was barely visible in the stigmatic papillae, pollen, and silique walls. The altered pattern of expression of the *ARF8* translational fusion relative to the mRNA localization and the expression pattern described previously for plants containing the transcriptional fusion (*P_{ARF8}:GUS*) indicates that *ARF8* might be regulated at the posttranscriptional level.

***ARF8* Is Expressed in the Ovule and the Embryo Sac**

Before pollination (stage 12), expression of *ARF8:GUS* in the carpel was detected primarily in the septum tissue around the vasculature and also in the funiculus of the ovule (Figure 5E). After pollination, but before fertilization had occurred (stage 13), there was a lack of staining between the funiculus and carpel vasculature. GUS activity in the ovule was also evident in the chalazal region of the integuments adjacent to the embryo sac (Figure 5F). Expression in the mature embryo sac was evident in three discrete zones: one in the chalazal cytoplasm of the embryo sac, a second in the fused polar nuclei and surrounding cytoplasm, and the third in the egg apparatus comprising synergids and the egg cell (Figures 5F and 5G). After fertilization, the level of GUS activity in the embryo and endosperm compartments decreased relative to that seen in the embryo sac and by the globular embryo stage of seed development GUS was barely detectable (Figure 5H).

In unfertilized wild-type ovules that are destined to senesce, *ARF8:GUS* expression was evident throughout the ovule (Figure 5I), resembling the expression pattern observed in ovules of emasculated *Ler* siliques transformed with the *P_{ARF8}:GUS* construct. The difference in expression pattern between fertilized and senescing ovules reveals that *ARF8* expression is regulated differently in the ovule depending on whether or not fertilization occurs.

***ARF8:GUS* Expression Is Spatially and Temporally Altered in the *arf8-4* Background**

The *ARF8:GUS* construct was inserted into *arf8-4* mutant plants by plant transformation, and 14 independent lines were obtained. Analysis of the resulting expression pattern showed significant differences in the expression of *ARF8:GUS* in the *arf8-4* background compared with that observed in *Ler*. Expression was observed much earlier in the developing carpel and was evident in flower buds as early as stage 5 (see Supplemental

Figure 1 online). Expression continued throughout flower development, silique elongation, and early embryo development (Figures 5J to 5P). Expression in carpel and silique wall tissues was much higher than that found in the *Ler* background and somewhat masked the underlying expression in septum and funiculi (cf. Figures 5C and 5N). Septum and funiculi expression began earlier, at stage 9, and septum expression did not decline after fertilization but continued throughout silique growth in the *arf8-4* background until siliques turned yellow (Figures 5L to 5P). Expression of *ARF8:GUS* was also observed earlier in the embryo sac of the ovules, and a general distribution pattern was observed instead of the three discrete zones seen in the *Ler* background (cf. Figures 5F and 5J). GUS activity declined after fertilization, but in contrast with the *Ler* background, where it had disappeared by the early globular embryo stage, staining was still visible in the chalazal region and the funiculus of the ovule when the embryo was at the globular stage of development in the *arf8-4* background (cf. Figures 5H and 5K). GUS activity continued to decrease and was not visible at the early heart stage of embryo development (see Supplemental Figure 1 online). There were also differences in staining intensities in some tissues between the *Ler* and *arf8-4* backgrounds, with decreased *ARF8:GUS* activity in petals and filaments in *arf8-4* (Figures 5L to 5N) compared with *Ler* (Figures 5A to 5C). These data indicate that *ARF8:GUS* is being regulated differently in the *arf8-4* mutant. The temporally extended expression agrees with the different level of expression of *P_{ARF8}:GUS* in the *arf8-4* background and the higher levels of mRNA evident in developing *arf8-4* siliques.

***ARF8:GUS* Induces the Formation of the Dehiscence Zone in Transgenic *Ler* Plants**

We tested *Ler* and *arf8-4* plants containing *ARF8:GUS* for parthenocarpy by floral emasculation. Surprisingly, the transgenic *Ler* plants developed genuine siliques that dehisced but did not elongate significantly over that observed in unfertilized, expanded carpels (Table 2). This contrasted with the indehiscent carpels (Table 1) observed previously in *Ler* plants containing the identical genomic fragment lacking the *GUS* gene fused in frame to the 3' end of the *ARF8* gene. This indicates that the introduced *ARF8:GUS* is compromising the activity of the endogenous *ARF8* gene to prohibit dehiscence zone formation in the absence of fertilization. Introduction of *ARF8:GUS* into the *arf8-4* background resulted in partial complementation of the parthenocarpic phenotype because elongation was reduced but the dehiscence zone still formed (Table 2). This was consistent with the inability to complement the *arf8-4* mutant with the genomic fragment (Table 1), further supporting the possibility that the *arf8-4* allele can interfere with the activity of the endogenous *ARF8* gene.

DISCUSSION

***ARF8* Is a Negative Regulator of Fruit Initiation and Growth**

The parthenocarpic *fwf* phenotype (Vivian-Smith et al., 2001) is caused by a lesion in *ARF8*. *ARF8* is a member of a family of 23 transcription factors in *Arabidopsis* (Guilfoyle and Hagen, 2001;

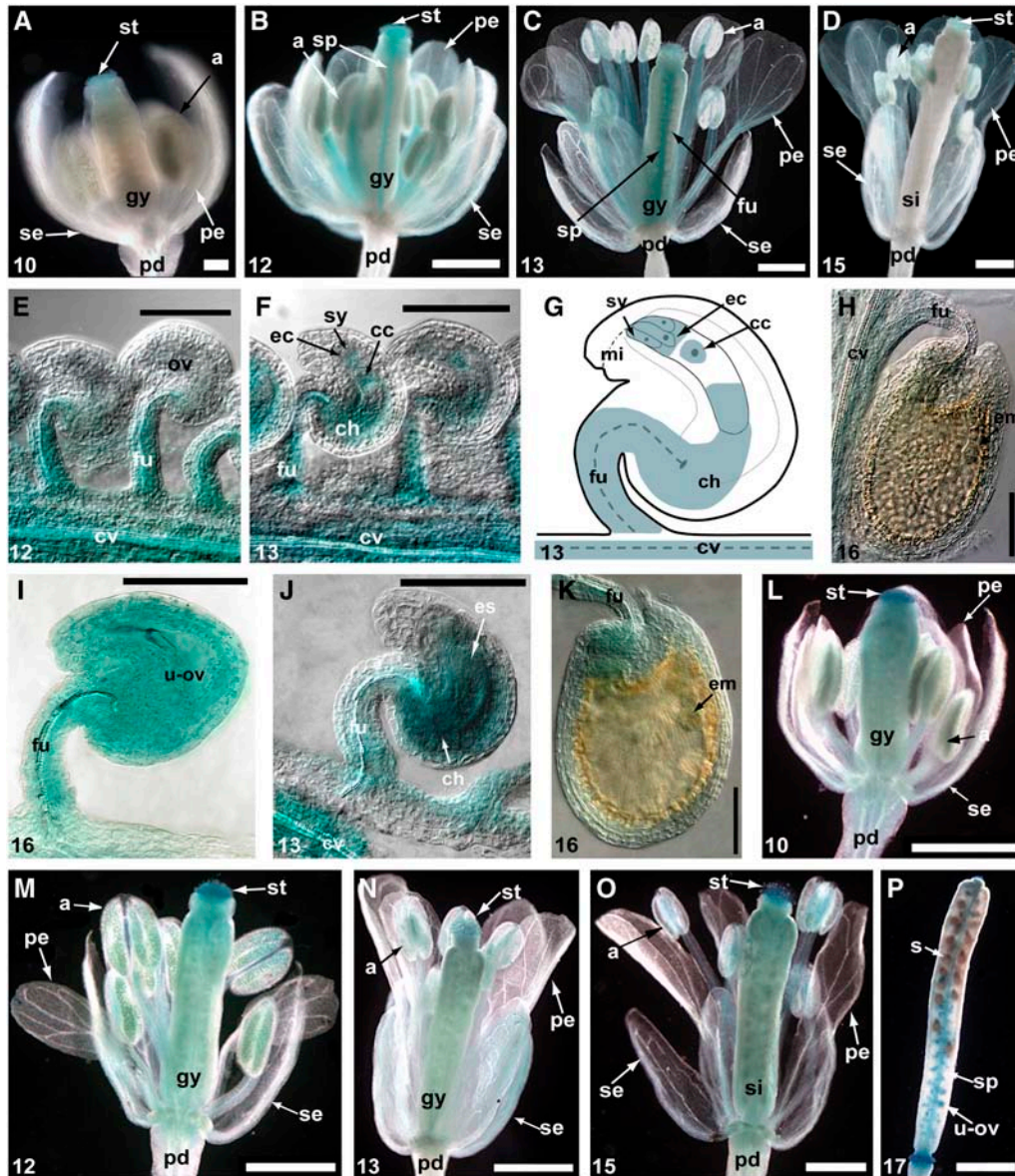


Figure 5. Expression of Translational ARF8:GUS Fusion Protein.

(A) to (D) GUS staining patterns in *Ler* flowers and siliques transformed with *ARF8:GUS* at different developmental stages. *Ler* flower bud at stage 10 (A); *Ler* flower around stage 12 (B); *Ler* flower at anthesis (stage 13) (C); *Ler* silique after fertilization (stage 15) (D).

(E) to (I) GUS staining patterns in *Ler* and *arf8-4* ovules between anthesis and globular embryo stage.

(E) Staining in septum tissue and funiculi before anthesis.

(F) After pollination, GUS staining is evident in the chalazal region and in the mature embryo sac around the fused polar nuclei and the egg apparatus.

(G) Schematic representation of GUS staining in ovules after pollination.

(H) At the globular embryo stage, no staining is visible in fertilized ovules.

(I) Unfertilized ovules show strong GUS expression throughout the ovule.

(J) GUS expression was observed earlier and in a general distribution pattern in the embryo sac of *arf8-4* ovules transformed with *ARF8:GUS*.

(K) In the *arf8-4* background, staining in the chalazal region is still visible at the globular embryo stage in fertilized ovules.

(L) to (P) GUS staining patterns in *arf8-4* flowers and siliques transformed with *ARF8:GUS* at different developmental stages.

(L) *arf8-4* flower at stage 10.

(M) *arf8-4* flower at stage 12. Expression is visible in the septum and within the ovules.

(N) *arf8-4* flower at anthesis (stage 13).

(O) After fertilization (stage 15), staining remains visible in the carpel walls, the septum, and the ovules.

(P) In growing siliques (stage 17), ARF8:GUS is expressed in the septum and in unfertilized ovules.

The numbers at the bottom left indicate the stages of flower development (Smyth et al., 1990). a, anther; cc, central cell; ch, chalazal end; cv, carpel vasculature; ec, egg cell; em, embryo; es, embryo sac; fu, funiculus; gy, gynoecium; mi, micropylar end; ov, ovule; pd, pedicel; pe, petals; se, sepals; sp, septum; si, silique; st, stigma; sy, synergids; u-ov, unfertilized ovule. Bars = 1 mm in (A) to (D) and (L) to (P) and 100 μ m in (E) to (K).

Table 2. Analysis of Elongation and Dehiscence in the Translational ARF8:GUS Lines

Line	Silique Length ^a	Elongation ^b	Dehiscence ^b
<i>Ler</i>	4.1 mm ± 0.4 mm	–	–
<i>Ler</i> + <i>ARF8:GUS</i>	3.9 mm ± 0.8 mm	–	++
<i>arf8-4</i>	7.2 mm ± 0.6 mm	+++	++
<i>arf8-4</i> + <i>ARF8:GUS</i>	4.9 mm ± 0.7 mm	+	++

^a Pistil lengths measured 7 d after emasculation (±SD; a minimum of 37 flowers was emasculated and measured).
^b –, no elongation/dehiscence; +, some elongation; +++/++, very good/good elongation/dehiscence.

Liscum and Reed, 2002; Remington et al., 2004) and has previously been linked to light-stimulated hypocotyl elongation, root growth, and auxin homeostasis (Tian et al., 2004). *ARF8* has also been shown to be involved in stamen filament elongation and anther dehiscence (Nagpal et al., 2005). Our analysis demonstrates that *ARF8* is also a negative regulator of fruit initiation because loss-of-function *arf8* alleles allow parthenocarpy and alter carpel mesocarp cell division (Vivian-Smith et al., 2001). The degree of carpel elongation in unfertilized flowers and the extent of parthenocarpic fruit growth was found to be dependent on the ecotype background, with the strongest silique elongation observed in the *Ler* background and significantly weaker elongation found in mutants in the *Col* and Wassilewskija (*Ws*) ecotypes (Vivian-Smith et al., 2001; this study).

Partial Complementation of *arf8-4* May Be Due to Inhibitory or Competitive Effects

Our attempt to complement the *arf8-4* mutation by the introduction of a wild-type genomic *ARF8* fragment only reduced the extent of parthenocarpic silique length, while other *arf8-4* characteristics, including the formation of the dehiscence zone without seed set, were still observed. Similarly, full complementation was not achieved in *arf8-1*, as the defective elongation in light-grown hypocotyls was incompletely restored (Tian et al., 2004). One possibility is that other genomic elements necessary for full *ARF8* function were absent in the fragments used for complementation. Alternatively, the *arf8-1* and *arf8-4* alleles may produce aberrant transcripts or proteins that prevent full complementation by the introduced wild-type copy of the gene. Consistent with this suggestion, both the *arf8-1* (Tian et al., 2004; data not shown) and *arf8-4* alleles produce transcripts with the potential to encode truncated ARF8 proteins that might interfere with wild-type ARF8 protein function. While further experiments are necessary to determine if a protein fragment can indeed be produced in these mutants, the ability of the ARF8:GUS transgene to induce parthenocarpic phenotypes in wild-type plants (albeit very short but dehiscent siliques) suggests that altered ARF8 proteins can have dominant-negative effects.

ARF8 Expression Is Regulated at Multiple Levels

Our analysis in both the wild-type and *arf8-4* backgrounds has shown that the expression of *ARF8* is regulated at multiple levels.

RNA gel blot analysis, RNA in situ hybridizations, and the observed expression pattern of a transcriptional *P_{ARF8}:GUS* construct showed that the expression of both mutant and wild-type mRNA was spatially comparable in developing flowers and during fertilization-induced silique growth. However, differences were found in the expression patterns of *P_{ARF8}:GUS* during parthenocarpic silique formation in *arf8-4* compared with that in *Ler* pistils undergoing senescence after emasculation. The parthenocarpic expression pattern resembled more that found in fertilization-induced *Ler* siliques, indicating that initiation of fruit development alters the expression of *ARF8*. Furthermore, both RNA gel blot analysis and qRT-PCR found higher levels of mRNA in the *arf8-4* plants, which also persisted in developing siliques. Experiments using the transcriptional *P_{ARF8}:GUS* fusion construct suggest that *ARF8* is involved in transcriptional autoregulation. The transcriptional regulation of *ARF8* and genes under control of the *ARF8* promoter is disturbed in the *arf8-4* mutant, leading to higher expression levels of those genes.

The analysis of a translational ARF8:GUS construct revealed an additional level of regulation. Expression of the translational ARF8:GUS fusion protein was altered in the *arf8-4* mutant in several ways compared with the wild-type pattern. ARF8:GUS levels in *arf8-4* appeared to be reduced in petals and especially anther filaments in the mutant. ARF8 activity in these tissues may be connected to the inhibitory effects of outer floral whorls on silique elongation (Vivian-Smith et al., 2001), and a role for ARF8 in stamen filament elongation has been demonstrated in the *arf6-2 arf8-3* mutant (Nagpal et al., 2005). Moreover, a temporal change in expression was also observed, especially in the carpel walls, the septum, the funiculi, and the ovules, where GUS activity was detected earlier and persisted in the mutant.

ARF8 translation may also be regulated via the 5'-transcript leader region. The *ARF8* mRNA is 629 bp longer than previously reported, and the translational leader region contains 10 upstream AUGs that the ribosomal translational complex can potentially act on before reaching the presumed *ARF8* initiation codon. These upstream open reading frames (uORFs) may regulate ARF8 translation, as similar uORFs have been identified in nine other ARF genes (Nishimura et al., 2004, 2005) where they have been shown to affect translation of the main ORF. Short uORFs in the 5'-transcript leader region of *ETTIN* (*ARF3*) and *MONOPTEROS* (*ARF5*) repress the expression of the downstream main ORFs at the translational level (Nishimura et al., 2005). Short upstream ORFs have also been reported in other genes where they significantly regulate translation of the main ORF (Geballe and Sachs, 2000; Morris and Geballe, 2000; Kawaguchi and Bailey-Serres, 2002).

Potential also exists for regulation of ARF8 by short-interfering RNAs, as recent research has shown that many ARFs are the target of microRNAs (miRNAs; Mallory et al., 2005) and trans-acting silence-inducing RNAs (Allen et al., 2005; Williams et al., 2005). The *ARF8* gene contains a miRNA target sequence (miR167; see Figure 2E) also present in *ARF6* (Rhoades et al., 2002; Bartel and Bartel, 2003; Kasschau et al., 2003). The levels of miR167, which are not regulated by auxin, appear to set a homeostatic level of ARF translation (Mallory et al., 2005) and might therefore be involved in the proposed posttranscriptional regulation or transcriptional autoregulation of *ARF8*. However,

the role of the miRNA in the modulation of *ARF8* expression and in the regulation of hypocotyl and root growth, flower development, or fruit initiation has not yet been investigated.

Signal Transduction Events in the Ovule May Regulate Fruit Growth

Fertilization is critical for initiating wild-type fruit formation. The expression pattern of *ARF8* in the ovule and female gametophyte around fertilization suggests that *ARF8* might act in these tissues to regulate fruit initiation.

ARF8:GUS expression is switched off soon after fertilization has occurred in wild-type plants, indicating that a fertilization signal deactivates *ARF8*. The removal of *ARF8* activity after fertilization might abolish a developmental block that represses fruit growth and allows initiation of seed and fruit developmental programs.

ARF8:GUS expression is spatially and temporally altered in the parthenocarpic *arf8-4* background, indicating a disturbed regulation of the introduced gene in the *arf8-4* mutant. This may result from an interaction between the *ARF8*:GUS gene and a mutant *arf8-4* protein product, as we have observed transcriptional autoregulation. However, we have yet to examine *arf8-4*:GUS expression and determine if a mutant protein is produced. We speculate that all of the examined *arf8* mutant alleles are unlikely to produce a functional *ARF8* protein, and parthenocarpy may simply occur because fruit development is not blocked.

In unfertilized ovules, expression of the *ARF8*:GUS marker persists and staining is observed throughout the ovule, indicating that the negative regulation through *ARF8* is kept active. These expression patterns are therefore consistent with *ARF8* acting as a negative regulator of fruit initiation, and collectively they indicate a central role for the ovules in mediating positive and negative signals involved in fruit development.

We observed previously that the *ats* mutation, which changes ovule integument structure, enhances the parthenocarpy phenotype when combined with *arf8-4* (Vivian-Smith et al., 2001). In emasculated *ats arf8-4* plants, seedless siliques formed that are comparable in length to those induced after fertilization. The *ats* mutation also counteracts the inhibitory effects of surrounding floral whorls on silique elongation (Vivian-Smith et al., 2001). *ATS* is *KANADI4*, and it is specifically expressed in the initiating ovule integument (McAbee et al., 2006), suggesting that modifications to the ovule integument influence parthenocarpic fruit growth in *ats arf8-4* plants. The importance and contribution of different structural components of the ovule to parthenocarpic fruit development can be examined genetically using a suite of ovule mutants in the *arf8-4* background. This should provide further information concerning the relationship of signal transduction events in the ovule and fruit growth and the role of *ARF8* in these processes.

Model for the Role of *ARF8* in Fruit Development

Figure 6 shows a model for the role of *ARF8* during the transition from carpel to fruit growth. Our data suggest that *ARF8* represses fruit development, and in the simplest model, *ARF8* may do so by directly activating genes that themselves repress fruit

development. Alternatively, *ARF8* may invoke repression by being a member of a complex of proteins. Protoplast transformation experiments support the concept that the transcriptional activity of the *ARF8* protein is regulated by heterodimerization with auxin/indole-3-acetic acid (*Aux/IAA*) proteins that inhibit this activity (Guilfoyle et al., 1998; Ulmasov et al., 1999a, 1999b; Guilfoyle and Hagen, 2001; Rogg and Bartel, 2001; Liscum and Reed, 2002; Tiwari et al., 2003). Physical interactions between *ARF8* and members of both the *Aux/IAA* repressor and *ARF* protein families have been demonstrated (Hardtke et al., 2004; Tatematsu et al., 2004). While studies have shown that *ARF8* transcription is not regulated by auxin per se (Ulmasov et al., 1999a; Pufky et al., 2003; Okushima et al., 2005), auxin indirectly regulates *ARF* activity by promoting turnover of *Aux/IAA* proteins (Ulmasov et al., 1999b; Gray et al., 2001; Tiwari et al., 2001; Zenser et al., 2001). This allows the *ARFs* to become active and impose their regulatory influence on the expression of auxin-responsive genes (Gray et al., 2001; Kepinski and Leyser, 2004). Consistent with an essential role for the closely related *ARF6* and *ARF8* proteins in flower development, several studies have demonstrated that *ARF8* can mediate auxin-induced gene activation (Ulmasov et al., 1999b; Tiwari et al., 2003; Nagpal et al., 2005), and several auxin-responsive genes have been identified as candidates for direct regulation by these two *ARFs* in microarray experiments (Nagpal et al., 2005). An increase in total auxin levels within flower buds during various developmental stages was not observed (Nagpal et al., 2005), suggesting that localized changes in auxin levels may be more important in regulating *ARF8* function.

In addition to its role with *ARF6* in flower development, we propose that before pollination and fertilization, *ARF8* is bound to the promoters of a range of primary auxin-responsive genes that play an essential role in fruit initiation and development. Transcription of these fruit initiation genes is repressed at this stage by a protein complex that includes *ARF8* bound to *Aux/IAA* proteins, which functions as a repressor (Figure 6A). This extends the model described above to include an active repressor role of the *ARF8*-*Aux/IAA* complex rather than merely a lack of gene activation by *ARF8*. The existence of a functional repressor function of the *ARF8*-*Aux/IAA* complex is required to explain the observation that reduced *ARF8* function allows fruit initiation. The identity of the *Aux/IAA* protein(s) proposed to act with *ARF8* is not known, but a likely candidate is the protein encoded by the *Arabidopsis* ortholog of tomato (*Solanum lycopersicum*) *IAA9* (Wang et al., 2005). As described above, this repressor function of *ARF8* may occur in ovules.

Pollination is known to induce increases in both auxin and ethylene levels in floral organs, and this correlates with subsequent growth or senescence of these organs (O'Neill, 1997). In wild-type flowers, a fertilization-induced auxin burst could induce the degradation of the *Aux/IAA* protein through a proteolytic pathway (Gray et al., 1999, 2001; Rogg and Bartel, 2001; Hellmann and Estelle, 2002; Dharmasiri and Estelle, 2004; Jenik and Barton, 2005). This would abolish the repression of crucial auxin-responsive fruit initiation genes by the *ARF8*-*Aux/IAA* protein complex (Figure 6A).

In the *arf8* mutant lines, this mechanism of repression is impaired. If no *ARF8* protein is made, the inhibitory complex

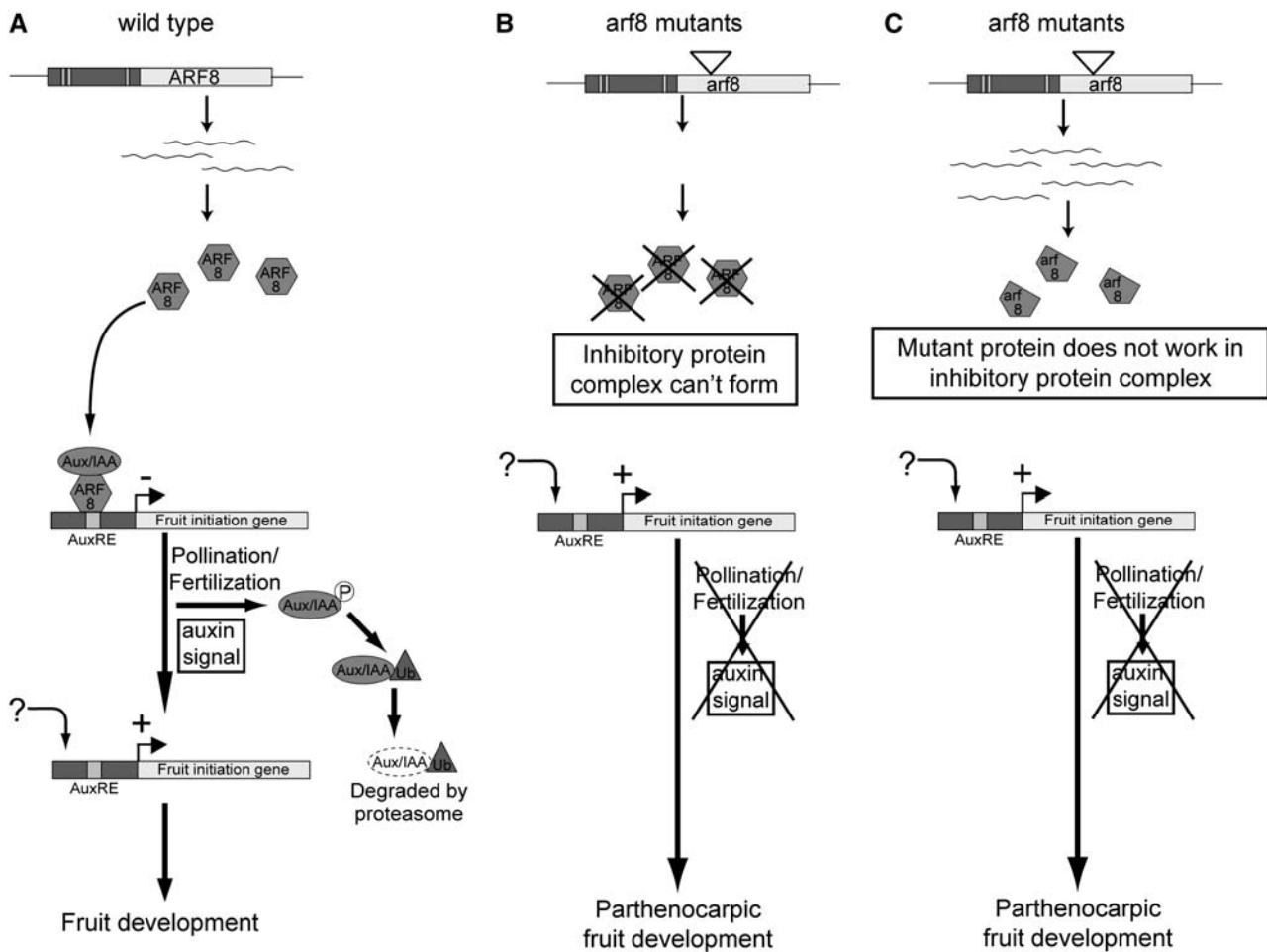


Figure 6. Model for the Role of *ARF8* in Fruit Development.

(A) Model showing *ARF8* action in restricting the expression of auxin-responsive genes in a complex together with an Aux/IAA repressor protein. The auxin signal induced by pollination leads to phosphorylation and subsequent proteolytic degradation of the Aux/IAA protein, and unknown factors subsequently activate expression of the fruit initiation genes.

(B) In *arf8* mutants in which no *ARF8* protein can be formed (null mutants), the inhibitory complex of *ARF8* with the Aux/IAA protein cannot form. Therefore, the expression of the fruit initiation genes can be activated in the absence of pollination and fertilization, leading to parthenocarpic fruit development.

(C) In *arf8* mutants that can produce a mutated version of the *ARF8* protein (*arf8*), this mutant protein fragment can either not bind to the Aux/IAA protein or does not bind the promoter of the fruit initiation genes. Therefore, the inhibitory complex is not correctly formed, and the expression of the fruit initiation genes can be activated in the absence of pollination and fertilization, leading to parthenocarpic fruit development.

cannot form or is unable to bind to the promoters of the auxin-responsive fruit initiation genes (Figure 6B). Alternatively, the formation of a functional inhibitory complex can be prevented by a mutant version of the *ARF8* protein that cannot bind to either the Aux/IAA protein or other components of the protein complex, or to the promoters of fruit development genes (Figure 6C). As a result, repression of fruit initiation gene transcription does not occur, and parthenocarpic results by allowing fruit initiation to occur in the absence of fertilization. Once the proposed *ARF8*-Aux/IAA complex is removed, either *ARF8* is replaced by other transcription factors, presumably other ARFs, or *ARF8* itself functions as a transcriptional activator. If the latter hypothesis is correct, the capacity to initiate fruit development in *arf8* mutants

implies that *ARF8* is functionally redundant in this activation process, presumably with *ARF6*.

The notion that biological processes actively restrict the carpel from forming a fruit was first proposed by Nitsch (1952), who from various studies held the view that auxin was a key component of this process. Many subsequent studies have suggested that developing seeds are important sources for phytohormones, such as auxins, needed for fruit development (Nitsch, 1970; Eeuwens and Schwabe, 1975; Archbold and Dennis, 1985; Talon et al., 1990a; Garcia-Martinez et al., 1991; Ben-Cheikh et al., 1997; Swain et al., 1997; Ozga et al., 2002). The discovery that *FWF* is *ARF8* supports these suggestions and means that we can now begin to dissect the developmental pathways controlling this crucial process.

METHODS

Plant Growth and Scoring Parthenocarpy

Arabidopsis thaliana seeds were surface-sterilized and grown as described before (Vivian-Smith and Koltunow, 1999). Parthenocarpy was assessed by flower emasculatation (Vivian-Smith and Koltunow, 1999; Vivian-Smith et al., 2001). Only siliques above flower position 20 were used, collected, and photographed, and their lengths were determined using the Scion Image Beta 4.02 program (http://www.scioncorp.com/frames/fr_scion_products.htm). Plants producing siliques that significantly and reliably elongated more than the corresponding wild-type ecotype plants and that formed a dehiscence zone were scored as parthenocarpic.

Fine Mapping of the *fwf* Mutation and Map-Based Cloning of *fwf*

The map position of the *fwf* lesion was described on chromosome V between the markers AthPhyC and AthSO191 (Vivian-Smith et al., 2001). Fine mapping was performed using recombinant populations from crosses between a Col-4 female parent and the *fwf* mutant (*Ler* ecotype) as pollen donor together with a population that consisted of F2 individuals from a cross between Col-4 and the recessive *ats fwf* double mutant (*Ler* ecotype) that was linked in coupling phase (Vivian-Smith et al., 2001). Recombinants between *fwf* and *ats* facilitated mapping, and both populations were used to generate informative recombinants in the AthPhyC and AthSO191 interval through a PCR screening method. Plants were assessed by the absence or presence of siliques exhibiting parthenocarpy when emasculated.

Plant DNA was extracted for PCR screening in microtitre plates essentially as described by Langridge et al. (1991). Multiplex PCR reactions were made with primers to the AthPhyC and AthSO191 CAPS markers. A total of 121 recombinant chromatids were identified from 2442 plants (Vivian-Smith, 2001). Each recombinant plant was transferred to soil, grown, and scored for parthenocarpic silique elongation. Genomic DNA was extracted from leaf tissue (Edwards et al., 1991) and AthPhyC and AthSO191 checked. New simple sequence length polymorphisms (Bell and Ecker, 1994) and new CAPS (Konieczny and Ausubel, 1993) molecular markers were created for the region spanning AthPhyC and AthSO191 (Vivian-Smith, 2001). Markers surrounding the ARF8a CAPS marker had the least number of recombinants. This marker is PCR amplified with ARF8+1478F (5'-GAGCTCCTTTAAGACAGCAGTTTGT-3') and ARF8+2817R (5'-CCTAGGAAAGTTAGTTACCCTGAGAC-3') and cut with *AccI* (New England Biolabs). The nearest centromeric marker was MLF18 (#2), a marker dominant in Col-1, amplified by MLF18#2F (5'-TTTGTCATGTTGGGTTCCG-3') and MLF18#2R (5'-GGGAGACGGGTGAGACAAAT-3'). The nearest telomeric marker was MJG14 (#2) amplified by MJG14#2F (5'-GGATCGTTAGGCAATGGGAT-3') and MJG14#2R (5'-TGGATACGATGGGGACAAA-3') and cut with *VspI* (New England Biolabs). The latter two markers defined 110 kb spanning part of the P1 vector MLF18 on the left and MJG14 on the right and the whole TAC vector K15O15.

ARF8 T-DNA Mutants

The *arf8-1* mutant was published as a hypocotyl elongation mutant allele (Tian et al., 2004) in the *Ws* background. Seeds were kindly donated by Kotaro Yamamoto (Hokkaido University, Sapporo, Japan).

The Col plant line SALK_049954 was designated *arf8-5*. The line was identified from the collection of SALK lines (Alonso et al., 2003) and obtained from the ABRC. The position of the T-DNA insertion was verified by sequencing, and it was found to be 431 bp 5' to the annotation published in The Arabidopsis Information Resource database (Figure 2D). The Col plant line WiscDsLox324F09 (University of Wisconsin T-DNA

lines; <http://www.hort.wisc.edu/krysan/DS-Lox/>) was designated *arf8-6* and was identical in phenotype to the *fwf* allele in Col-1 (*arf8-4NIL*; Vivian-Smith, 2001).

Cloning of ARF8 and Complementation of *fwf*

The *ARF8* gene, including 1.8 kb of 5' upstream promoter sequence and the complete coding region including the 3'UTR, was amplified from *Ler* wild-type and *fwf* mutant plants via PCR using the eLONGase amplification system (Invitrogen) according to the manufacturer's instructions with the primers ARF8-1830F (5'-CTCGAGTGAGAAGTCTATGATG-3') and ARF8+2817R (5'-CCTAGGAAAGTTAGTTACCCTGAGAC-3'). The PCR products were cloned using the pGEM-T Easy Vector System I (Promega), and sequencing revealed that the *fwf* fragment contained the transition from G to A in the predicted translation initiation codon (Vivian-Smith, 2001). The identified mutation in *ARF8* from *fwf* plants creates a CAPS marker. Amplification with the primers ARF8-143F (5'-AGGAG-ATGGAGAAAGACGAG-3') and ARF8+48R (5'-CTCTCCTTCATGACCC-TGTTG-3') and subsequent digest with *Hsp92II* (Promega) resulted in bands of 142 bp + 41 bp + 8 bp from *Ler* wild-type plants, whereas 183 bp + 8 bp bands were present in *fwf* plants.

The wild-type version of *ARF8* was subcloned into the pBIN19 vector (Bevan, 1984) and transformed into *Agrobacterium tumefaciens* strain LBA4404. *Ler* and *fwf* plants were separately transformed with the construct via the floral dip method (Clough and Bent, 1998).

RNA Preparation and RNA Gel Blot Analysis

Total RNA was extracted from *Arabidopsis* plant tissues using TRIZOL LS reagent (Invitrogen) or the RNeasy plant mini kit (Qiagen). RNA gel blot analysis was performed with 10 µg of total RNA as described (Goetz et al., 2001) using a radioactive labeled *ARF8*-specific probe. The probe was labeled using the random primer Rediprime II DNA labeling system (Amersham Biosciences).

DNA Preparation and DNA Gel Blot Analysis

Genomic DNA was extracted from *Arabidopsis* plant tissues using a CTAB extraction protocol (Goetz et al., 2001). Ten micrograms of genomic DNA were digested with different restriction enzymes for 24 h and then subjected to electrophoresis in a 0.8% agarose gel. DNA was transferred to a nylon membrane (Biodyne B/Plus; PALL Gelman Laboratory), and hybridization and autoradiography were performed as for RNA gel blot analysis.

RT-PCR and qRT-PCR

One microgram of total RNA extracted from *Arabidopsis* plant tissues with the RNeasy plant mini kit and treated with an on-column RNase-free DNase protocol was used as template for cDNA synthesis with the ThermoScript RT-PCR system (Invitrogen). One microliter of the cDNA was then used for a standard PCR reaction using the primer pair NKE11F (5'-GCGGC-CGCGGTACCTTTCCTATGTATCCA-3') and ARF8+3956R (5'-GTGACCTAGAGATGGGTGCGGGTTTTCG-3') to amplify *ARF8* and the primer pair b-tubF (5'-GGGTGCTGGAACAATTGGGCTAA-3') and b-tubR (5'-ACTGCTCACTCACGCGCTAA-3') to amplify β-tubulin as a control.

For qRT-PCR, 3 µL of one-fifth diluted cDNA were used as a template in a 15 µL reaction with 300 nM of each primer and 7.5 µL of 2× Absolute QPCR SYBR Green ROX mix (ABgene, Integrated Sciences). Reactions were performed using the Rotor-Gene 3000 (Corbett Research, Adela Scientific) and analyzed with Rotor-Gene 6.0 software (Corbett Research, Adela Scientific). Primer pairs used for amplification of *ARF8* fragments during qRT-PCR were as follows: Q-ARF8-122F (5'-TTGTACTTCCGGAGCT-AAAGAGTT-3') and Q-ARF8-15R (5'-CAAGAAACCAACTTTGAAAACC-3'),

Q-ARF8+296F (5'-AACCATTTGACACCGGAGGAG-3') and Q-ARF8+481R (5'-GCTGCAGTGTGAATCCAATGG-3'), or Q-ARF8+2495F (5'-TGAAGTCGTTTCCACTCATCTTT-3') and Q-ARF8+2601R (5'-AAGTTTCAGGACCCATACCTACA-3'). The primer pair Q-UbceF (5'-AGGTACAGCGAGAGAAAGTAGCAGA-3') and Q-UbceR (5'-AACAGAAAAGCAAGC-TGAAAAACA-3') was used for amplification of a ubiquitin conjugating enzyme (At5g25760) fragment as a control and to normalize cDNA starting amounts.

GUS Constructs, Staining, and Microscopy

To create the transcriptional fusion construct $P_{ARF8}:GUS$, 1.8 kb of the 5' upstream promoter sequence of *ARF8* was amplified from *Ler* wild-type plants via PCR with the primers ARF8-1830F (5'-CTCGAGTGAGAAGTC-TATGATG-3') and ARF8-1R (5'-CTCGAGTTCAACTCAAGAA-3'). The PCR fragment was cloned into the vector pART7NAPX, which contains a promoterless GUS gene (Gleave, 1992). The expression cassette, containing the amplified *ARF8* promoter, the GUS gene, and the OCS terminator sequence, was excised from the vector with *NotI* and cloned into the binary vector pART27 (Gleave, 1992). This was transformed via *Agrobacterium* strain LBA4404 as described above (Clough and Bent, 1998).

The translational fusion constructs consisted of 5.8 kb of genomic sequence, including 1.8 kb of the 5' upstream promoter sequence and the entire *ARF8* coding region excluding the stop codon. This was amplified from wild-type plants and *arf8-4* plants with the primers A8-1823SalI (5'-GTCGACTGAGAAGTCTATGATGAG-3') and ARF8:G:g:Rev (5'-CATCCCTAGGGAGATGGGTCGGGTTTGC CGGAA-3'). The PCR fragment was cloned into the vector pGEM T-easy (Promega). *Sall* and *AvrII* fragments were released, allowing the *ARF8* fragment to be subcloned in frame with a promoterless GUS gene in the vector pCambia1381Xa (GenBank accession number AF234303). The constructs were transformed via floral dip as above (Clough and Bent, 1998).

Histological analysis of GUS enzyme activity in *Arabidopsis* tissue was done essentially as described by Jefferson et al. (1987). The stained tissues were postfixed in FAA (4% formaldehyde [v/v], 5% acetic acid [v/v], and 50% ethanol [v/v]) and cleared in 70% ethanol before being photographed. Whole-mount tissue samples were viewed with Stemi2000C or Axioplan microscopes (Carl Zeiss). Digital images were captured using a Spot II camera (Diagnostic Instruments Inc). Image processing and reproduction were performed with Auto Montage Essentials (Synchroscopy) and Photoshop 7.0 (Adobe Systems).

In Situ Hybridization

A 328-bp fragment from the 3'UTR of *ARF8* was amplified with primers ARF8+2481F (5'-GGTACCAGAAGATGTGCATCAAATGGG-3') and ARF8+2718R (5'-CCTAGGAAAGTTTAGTTACCCTGAGAC-3') and cloned into pGEM-T Easy. Plasmids were sequenced to verify identity and orientation of inserts. Probe preparation and in situ hybridizations were performed as described previously (Tucker et al., 2003), except that once probes were added to the formamide-based hybridization solution and cover slips applied, the slides were heated to 80°C for 2 min prior to hybridization overnight at 42°C.

Accession Numbers

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers AT5G37020 (*ARF8*) and AT1G30330 (*ARF6*).

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Table 1. Expression Ratio of *ARF8* mRNA in Various Tissues.

Supplemental Figure 1. Expression of *ARF8* and GUS Marker Constructs in *Ler* and *arf8-4* Plants.

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AUXIN RESPONSE FACTOR8* Is a Negative Regulator of Fruit Initiation in *Arabidopsis

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