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Additional Information

Developmental role of the tomato Mediator complex subunit MED18 in pollen

ontogeny

Fernando Pérez-Martín¹, Fernando J. Yuste-Lisbona¹, Benito Pineda², Begoña García-

Sogo², Iván del Olmo³, Juan de Dios Alché⁴, Isabel Egea⁵, Francisco B. Flores⁵, Manuel

Piñeiro³, José A. Jarillo³, Trinidad Angosto¹, Juan Capel¹, Vicente Moreno² and Rafael

Lozano^{1,*}

¹ Centro de Investigación en Biotecnología Agroalimentaria (BITAL). Universidad de

Almería, 04120 Almería, Spain,

² Instituto de Biología Molecular y Celular de Plantas, Universidad Politécnica de

Valencia-CSIC, 46022 Valencia, Spain.

³ Centro de Biotecnología y Genómica de Plantas, Universidad Politécnica de Madrid

(UPM) - Instituto Nacional de Investigación y Tecnología Agraria y Alimentaria

(INIA), Campus Montegancedo UPM, 28223 Pozuelo de Alarcón (Madrid), Spain

⁴ Departamento de Bioquímica, Biología Celular y Molecular de Plantas, EEZ-CSIC,

18008 Granada, Spain.

⁵ Departamento de Biología del Estrés y Patología Vegetal, CEBAS-CSIC, 30100

Espinardo-Murcia, Spain

* Corresponding author: rlozano@ual.es

Prof. Rafael Lozano

Dept. Biología y Geología

Edificio Científico-Técnico II-B

Universidad de Almería

04120 Almería, Spain

Tel. +34 950 01 5111

Fax +34 950 01 5476

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Summary

Pollen development is a crucial step in higher plants which not only makes possible plant fertilization and seed formation, but also determine fruit quality and yield in crop species. Here, we reported a tomato T-DNA mutant, pollen deficient1 (pod1), characterized by an abnormal anther development and the lack of viable pollen formation, which led to the production of parthenocarpic fruits. Genomic analyses and the characterization of silencing lines proved that pod1 mutant phenotype relies on the tomato SIMED18 gene encoding the subunit 18 of Mediator multi-protein complex involved in RNA polymerase II transcription machinery. The loss of SIMED18 function delayed tapetum degeneration, which resulted in deficient microspore development and scarce production of viable pollen. A detailed histological characterization of anther development proved that changes during microgametogenesis and a significant delay in tapetum degeneration are associated with a high proportion of degenerated cells and hence, should be responsible for the low production of functional pollen grains. Expression of pollen marker genes indicated that SIMED18 is essential for the proper transcription of a subset of genes specifically required to pollen formation and fruit development, revealing a key role of SIMED18 in male gametogenesis of tomato. Additionally, SIMED18 is able to rescue developmental abnormalities of the Arabidopsis med18 mutant indicating that most biological functions have been conserved in both species.

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Introduction

Pollination and fertilization of angiosperms are coordinated processes which allow the conversion of gynoecium into a seeded fruit and therefore, are essential events to ensure species survival and fruit yield (Gillaspy et al., 1993; Ozga and Reinecke, 2003; Carbonell-Bejerano et al., 2010, Lora et al., 2011). Thus, pollen development and maturation comprises multiple cellular changes mediated by a precisely orchestrated gene expression regulation (Honys and Twell, 2004; Pina et al., 2005; Wilson and Zhang, 2009; Feng et al., 2012; Rutley and Twell, 2015). Male gametogenesis takes place in anthers, where diploid archesporial cells divide into two cell layers with different fates: i) the primary parietal layer, which gives rise to four different cell layers by successive divisions, to form concentric layers of pollen sac wall, i.e. epidermis, endothecium, middle layer and tapetum; and ii) the primary sporogenous layer, which undergoes a small number of divisions to produce pollen mother cells (PMCs). These undifferentiated cells undergo meiosis leading to the formation of tetrads of haploid cells, which are released as free microspores. Finally, uninucleate microspores mature after an asymmetric mitotic division to produce pollen grains, which in turn enclose the vegetative and the generative cells (Scott et al., 2004).

Significant progress in understanding the genetic and molecular basis of pollen development has been made from the study of mutants in the model species Arabidopsis thaliana. Thus, mutations affecting several stages of pollen ontogeny have been identified, which have led to the isolation and functional analysis of several genes involved in pollen development. For instance, sporocyteless/nozzle (spl/nzz) mutant fails to form sporogenous tissue during early anther development. SPL/NZZ encodes a MADS-box transcription factor that plays a central role in regulating anther cell differentiation (Schiefthaler et al., 1999; Yang et al., 1999; Liu et al., 2009). In the case of switch1 (swi1) mutant, male gametogenesis is affected during meiosis of PMCs. SWI1 encodes a novel protein involved in sister chromatid cohesion and meiotic chromosome organization during both male and female meiosis (Mercier et al., 2003). Likewise, during microgametogenesis, the programmed cell death of the tapetum tissue is essential for proper pollen development, as it supplies nutrients to the microspores, as well as for regulating microspores release (Pacini, 2010). Consequently, mutations that disrupt tapetum ontology and promote aborted microgametogenesis causing male sterility have been reported, mainly extra sporogenous cells/excess microsporocytes1 (ems1/exs), tapetal determinant1 (tpd1), aborted microspores (ams) and male sterility1 (ms1) mutations. EMS1/EXS and TPD1 genes encode a putative LRR receptor kinase and a small putatively-secreted protein, respectively, both required for specifying tapetal identity (Canales et al., 2002; Zhao et al., 2002; Yang et al., 2003). On the other hand, AMS gene encodes a transcription factor belonging to the MYC subfamily of bHLH genes, which is required for tapetal cell development in Arabidopsis (Sorensen et al., 2003). Similarly, ms1 mutant pollen degenerates after microspore release due to an abnormal vacuolization of the tapetum (Wilson et al., 2001). Therefore, MS1, which encodes a PHD-finger class of transcription factors, is a key gene required for correct tapetum degradation (Yang et al., 2007).

In tomato, pollen development and the predetermined tapetum degeneration processes are quite similar to that of Arabidopsis (Polowick and Sawhney, 1993a,b; Brukhin et al., 2003; Wilson and Zhang, 2009). Tomato male sterility has been the subject of genetic research since it was first described by Crane (1915), whereupon more than 50 tomato male sterile mutants have been reported (Gorman and McCormick, 1997); however, in contrast to Arabidopsis, a small number of pollen development related genes have been identified so far. Among them, the tomato homologue to the Arabidopsis ECERIFERUM6 (CER6) gene, which encodes a β-ketoacyl-coenzyme A synthase, is involved in the regulation of timely tapetum degradation (Smirnova et al., 2013). Recently, it has been demonstrated that *Male sterile* 10³⁵ (Ms10³⁵) gene encodes a basic helix-loop-helix transcription factor, which participates in regulating both meiosis and programmed cell death of the tapetum during microsporogenesis (Jeong et al., 2014). Likewise, the glycine-rich protein LeGRP92 is essential for normal pollen function and survival as it facilitates the outer cell wall (or exine) formation (McNeil and Smith, 2010). Similarly, the LATE ANTHER TOMATO 52 (LAT52) gene, encoding a heat-stable glycosylated protein, plays a crucial role in pollen hydration and germination (Muschietti et al., 1994).

Given the relevance of male sterility and fruit set for tomato breeding, a collection of tomato T-DNA insertion lines generated by an enhancer trap was screened (Pérez-Martín *et al.*, 2017) aiming to identify new regulators involved in male fertility. This work describes the characterization of a tomato T-DNA mutant, *pollen deficient1* (*pod1*), which displayed a significant reduction of pollen viability that yielded parthenocarpic fruits. Functional analyses demonstrated that the loss of *MEDIATOR COMPLEX SUBUNIT 18* (*POD1/SIMED18*) function is responsible for the observed *pod1* phenotypic alterations. MED18 is a subunit of the MEDIATOR COMPLEX that

binds RNA Polymerase II, an evolutionarily conserved transcriptional regulatory complex of class II genes in all eukaryotes (Kornberg *et al.*, 2005; Bourbon, 2008). Depending on the species, MEDIATOR is a large multimeric protein comprising 25-34 subunits (Allen and Taatjes, 2015; Samanta and Thakur, 2015), which are organised in four modules: head, middle, tail and CDK/Cycline (Chadick and Asturias, 2005; Conaway *et al.*, 2005). Each subunit seems to be a specific regulator for defined gene sets related to different functions involved in gene transcription mediated by RNA Polymerase II, including transcription, initiation, and elongation, as well as RNA processing, chromatin spatial conformation and enhancer-promoter interaction (Buendía-Monreal and Gillmor, 2016).

In Arabidopsis, *MED18* is involved in the transcriptional response to different physiological and cellular processes, like plant immunity (Lai *et al.*, 2014), flowering time and floral organ identity (Zheng *et al.*, 2013). Recently, Wang *et al.* (2018) have reported the function of tomato *MED18* in regulating the development of leaf and stem. However, little is known about the role of *MED18* in reproductive development. This study reports a key function of *MED18* in pollen ontogeny, which is required to ensure a proper differentiation and maturation of pollen grains and tapetum degradation in tomato.

RESULTS

Phenotypic and genetic analysis of the pollen deficient 1 (pod1) mutant

The *pod1* mutant was isolated from the screening of a collection of enhancer trap lines in the tomato cv. Moneymaker. Mutant plants were mainly affected in flower development. At anthesis stage, flowers showed a significant decrease in the length of petals, stamens and pistils of *pod1* flowers (Figure 1b; Table S1). About 10% of *pod1* flowers displayed different degrees of homeotic changes, from near to wild-type (WT) to aberrant phenotypes, which affected floral organs of the second and mainly the third whorls, indicating incomplete penetrance and variable expressivity of *pod1* mutation. Indeed, scanning electron microscopy (SEM) analysis showed the development of trichomes on the adaxial surface of some petals, a feature never observed in WT petals, as well as increased size of epidermal cells (Figure S1a,b). Moreover, a range of homeotic phenotypic alterations was observed from normal stamens to full conversion of stamens into carpels in the third whorl (Figure S1c,d), where the latter ones showed dramatic changes in size and shape of their epidermal cells compared to WT ones

(Figure S1e,f). Thus, both types of *pod1* stamens, with full and without any homeotic conversion, were subjected to qRT-PCR analysis. Three tomato B-class identity genes were assessed, i.e, *STAMENLESS* (*SL*), *TOMATO MADS6* (*TM6*) and *TOMATO PISTILLATA* (*TPI*), and significant down-regulation of all of them was observed in *pod1* stamens showing homeotic changes (Figure S1g). However, the expression of *TOMATO AGAMOUS 1* (*TAG1*), a C-class identity gene, was not altered in stamens of *pod1* plants (Figure S1g).

Stamens developed by *pod1* flowers showed a significant reduction in the amount (~5-fold) and viability (~20-fold) of pollen grains as compared to WT ones (Figure 1c,d; Table S1). An *in vivo* pollen germination assay was also performed through reciprocal crosses, which showed that WT pollen grains germinated and developed normal pollen tubes on stigmas of *pod1* flowers (Figure 1e); however, *pod1* pollen grains were unable to form pollen tubes on WT stigmas (Figure 1f). WT pollen was used in cross-pollination assays which yielded normal-seeded fruits, indicating that *pod1* mutation did not affect ovule functionality. However, selfing of *pod1* plants gave rise to small and parthenocarpic fruits, which showed reduced axial and equatorial diameters as well as decreased fresh weight compared to WT fruits (Figure 1g; Table S1).

In addition, vegetative developmental traits were also altered in *pod1* mutant plants, mainly a significant reduction in fresh weight and leaf length compared to WT plants, as well as a decreased development of secondary and tertiary leaflets, and reduced length of petioles (Figure 1a; Table S1). Such alterations coincide with those reported by Wang et al. (2018).

A genetic analysis of *pod1* mutant phenotype was performed on 411 segregating plants from two different progenies, 96 T2 and 315 T3 plants. The mutant phenotype was observed in 25 out of 96 T2 plants (26.04%) and 81 out of 315 T3 plants (25.71%). The Chi-square statistic test confirmed that segregation ratios were consistent with a monogenic autosomal recessive mode of inheritance for the *pod1* mutant phenotype ($\chi^2 = 0.14$, P = 0.71).

Cloning and molecular characterization of the *pod1* mutant locus

Southern blot hybridization indicated that a single copy of T-DNA was inserted in the *pod1* genome (Figure 1h). Afterwards, with the aim to isolate the gene harbouring *pod1* mutation, anchor-PCR assays were performed to clone the genomic regions flanking the

T-DNA insertion. Results revealed that T-DNA was located on chromosome 06 (at position 1,860,352 bp; ITAG2.50), in the promoter region of two adjacent genes transcribed in opposite direction, the *MEDIATOR COMPLEX SUBUNIT 18* (*SIMED18*) gene (*Solyc06g008010*) and a tomato member of the Zinc Finger HIT-type (ZNHIT) transcription factor family (*Solyc06g008020*). Specifically, the T-DNA was inserted 482 bp upstream of the 5'-untranslated region of the *SIMED18* gene and 203 bp upstream of the translation start codon of the *ZNHIT* gene. The T-DNA insertion produced a deletion of 160 bp which affected the promoter region of both *SIMED18* and *ZNHIT* genes (Figure 1i).

To establish a possible correlation between the T-DNA insertion site in the genome with the *pod1* phenotype, a co-segregation analysis was performed by PCR in 411 plants from T2 and T3 progenies, which revealed that a total of 106 mutant plants (25.80%), 25 from T2 and 81 from T3, were homozygous for the T-DNA insertion, whereas 201 of the 305 WT plants (48.90%) were hemizygous and the remaining 104 WT plants (25.30%) were azygous for the T-DNA insertion (Figure 1j). Therefore, results of co-segregation analysis supported that the *pod1* phenotype was linked to the T-DNA insertion.

Given the genomic position of the T-DNA insertion, qRT-PCR experiments were carried out in *pod1* and WT plants to determine whether the *pod1* mutation affected the expression of *SlMED18* and *ZNHIT* genes. Results showed that both *SlMED18* and *ZNHIT* genes were down-regulated in all *pod1* tissues here analysed, i.e. root, stem, leaf, apex and flower at anthesis (Figure 1k,1).

Phenotype of SIMED18 silencing plants resembles pod1 mutant

To conclude which of the two candidate genes affected by the T-DNA insertion was responsible for *pod1* mutant phenotype, single and double RNAi silencing lines for *SIMED18* and *ZNHIT* genes were generated, being the latter ones used to evaluate the hypothesis that simultaneous down-regulation of both genes could be responsible for *pod1* mutant phenotype. qRT-PCR analysis proved that gene silencing specifically affected the gene (or genes) targeted in each type of RNAi line (Figure 2d; Figure S2a-c). Phenotypic characterization of representative RNAi lines revealed that developmental alterations of both RNAi *SIMED18* and double RNAi transgenic plants were similar to those of *pod1* plants (Table S1). Indeed, these transgenic flowers were also smaller displaying shortened stamens and narrow petals as occurred in *pod1* mutant

(Figure 2a). In addition, some flowers of *SIMED18* silenced lines showed homeotic alterations in the second and the third floral organ whorls similar to those observed in *pod1* flowers (Figure S3). Likewise, a small amount of non-viable pollen grains was developed from *SIMED18* repressed lines (Figure 2b). Regarding the fruits, there was a strong similarity among RNAi *SIMED18*, double RNAi and *pod1* plants, as all of them yielded parthenocarpic fruits with a decreased fresh weight (Figure 2c). On the contrary, RNAi *ZNHIT* transgenic plants displayed a similar phenotype to WT plants with no obvious alteration in reproductive developmental traits (Figure 2a-c). Moreover, offsprings of RNAi *ZNHIT* lines also showed a WT phenotype, supporting the hypothesis that *pod1* mutant phenotype was caused by the loss of function of *SIMED18*.

To further confirm that down-regulation of *SIMED18* is responsible for the *pod1* phenotype, a molecular complementation assay was performed by overexpressing *SIMED18* in *pod1* plants under the control of a 35S constitutive promoter. Reproductive development of 35S::*SIMED18 pod1* lines were similar to WT controls (Figure 2a-c; Table S1), indicating that the overexpression of *SIMED18* gene was able to rescue the *pod1* mutant phenotype.

Expression patterns of SIMED18 during tomato reproductive development

SIMED18 is expressed from floral buds to mature fruits, although the highest level of SIMED18 transcripts was detected in flowers at anthesis, and the lowest one was found in fruits at immature green stage (Figure 3a). In situ hybridization analysis of SIMED18 in developing flower buds showed that SIMED18 mRNA was located in the two inner whorls of floral buds at stage 5 (according to Brukhin et al., 2003), where stamen and carpel primordia were initiated (Figure 3b,c). Later, expression of SIMED18 was strongly detectable in pollen and ovules at stage 8 of flower development (Figure 3d). Additionally, as the binary vector pD991 used for generating the enhancer trap lines contained a minimal promoter fused to the uidA reporter gene, a histochemical GUS assay was performed assuming that GUS expression is due to the activity of endogenous regulatory elements that promote the transcription of the uidA gene. In pod1 flowers, GUS staining was detected in stamens, stigma and ovules (Figure 3e), supporting that SIMED18 gene is specifically expressed in the two innermost floral organs.

In addition, given the homeotic conversion of stamens into carpel organs found in a low number of *pod1* flowers, expression of *SIMED18* was analysed in the floral organs developed in the third whorl of *pod1* mutant flowers. All *pod1* mutant flowers

showed down-regulation of *SIMED18* regardless of the existence of stamen to carpel homeotic conversion (Figure S1g), suggesting that such developmental alterations may correspond to pleiotropic effects of *pod1* mutation rather than being a direct consequence of the loss of *SIMED18* function.

Down-regulation of *SIMED18* affects microgametogenesis and tapetum degradation

Flowers of pod1 and SIMED18 silenced lines developed a small proportion of viable pollen (Figure 2b); thus, a detailed study at the cellular level was performed to detect changes in pollen ontogeny promoted by the down-regulation of SIMED18 function (Figure 4). To rule out any effect of ZNHIT down-regulation, RNAi SIMED18 lines were used for this study instead of *pod1* mutant. PMCs prior to meiosis did not differ in their morphological features when observed in thin tissue sections. Indeed, the characteristic polyhedral shape, slightly stained cytoplasm, and visible nuclei with densely stained nucleolus were observed both in WT and RNAi SIMED18 plants (Figure 4a,d). Meiocytes at telophase I stage were also similar in shape, size and cytological characters, as well as the features of callose layer, and cytoplasm and chromatin staining properties (Figure 4b,e). These results pointed out to an equally canonical meiosis occurring during pollen development of the SIMED18 silenced plants. Callose-embraced microspores within the tetrad also showed no differences between both WT and silenced lines, thus, tetrad walls appear well defined and microspores show stained nuclei with nucleoli (Figure 4c,f). Moreover, at meiocyte and tetrad stages no appreciable differences were detected in the size and morphological features of the tapetum and the remaining layers of the anther wall (Figure 4a-f).

Subsequent stages of pollen development were also analysed and DAPI staining of squashed anther samples was performed in order to assess the corresponding microgametogenesis stage engaged, based on the presence and the position of nuclei. Vacuolated microspores showed a typical cytoplasmic distribution in WT anthers, with the occurrence of cytoplasmic vacuoles and a single nucleus clearly stained by DAPI (Figure 4g,j). However, RNAi *SIMED18* flowers developed some microspores which were smaller in size and displayed lower DAPI staining (Figure 4m,p), likely reflecting symptoms of chromatin disorganization. Differences between the WT and the RNAi *SIMED18* plants increased at the stages of young and mature pollen, with a progressive larger proportion of pollen grains with altered morphology, mainly small size,

differential cytoplasmic density, and the presence of empty pollen grains (Figure 4h,i,n,o). DAPI staining analysis of RNAi *SIMED18* anthers showed a reduced amount of apparently normal pollen grains with both the vegetative and the generative nuclei. Instead, a higher proportion of abnormal pollen grains either bearing degenerated nuclei or completely lacking nuclei were observed in anthers where *SIMED18* was silenced (Figure 4k,l,q,r).

Developmental differences in the timing and completion of tapetum degeneration were found throughout microgametogenesis of WT and RNAi SIMED18 anthers (Figure 5). Although histological features and relative size of the tapetum layer until the tetrad stage were identical in both types of plants, the subsequent tapetal degeneration that begins at the microspore stage in the WT plants (Figure 5a-c) was delayed in SIMED18 silenced plants (Figure 5d-f). Indeed, while tapetum tissue showed evident degradation symptoms and had almost disappeared at mature pollen stage in WT anthers (Figure 5c), it remains intact in RNAi plants (Figure 5f,g). Taken together, these observations indicated that the lack of SIMED18 function provoked significant changes during pollen ontogeny, which affected mainly tapetum degradation and pollen maturation. Such developmental abnormalities correlated with the lower percentage of pollen yielded by RNAi SIMED18 plants.

Silencing of SIMED18 modifies expression of genes involved in anther and pollen development

To investigate how the lack of *SIMED18* affects the expression of genes involved in anther and pollen ontogeny, a comparative qRT-PCR analysis was carried out in floral buds at five pivotal stages of anther development, i.e. PMCs, tetrads (Tds), young and vacuolated microspores (Mcs), young pollen (YP) and mature pollen (MP) (Figure 6). Thereby, the expression pattern of *SIMED18* and seventeen additional genes previously described as key regulators involved in tomato pollen and anther development was evaluated (Jeong *et al.*, 2014, Gómez *et al.*, 2015). Given that the asynchrony in the degradation timing of the tapetum suggested an abnormal RNAi *SIMED18* pollen formation, the considered genes were divided into two groups according to their functions in WT anthers: genes related to tapetum degradation and moreover, genes associated to pollen formation and maturation. In addition to *SIMED18*, the first group included *SISPL/HYDRA*, *MS10*³⁵, *AMS-like*, *AtMYB103-like*, *MS1-like*, *TGAS100*, *bHLH89/91*, *TA29*, *Cysteine protease*, *Aspartic proteinase* and *Arabinogalactan protein*

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(Figure 6), while the second one comprised *Sister chromatid cohesion*, *TomA108*, *LeGRP92*, *Endo-1,3-beta-glucanase*, *AtTDF1-like* and *pLAT52* (Figure S4). Most of the tapetum development related genes here analysed were down-regulated at some stage of pollen development in RNAi *SIMED18* lines (Figure 6c-f,i-l). However, *SISPL/HYDRA* and *bHLH89/91* were expressed without significant differences between WT and RNAi *SIMED18* anthers in all stages analysed (Figure 6b,h). In the case of *TGAS100*, the relative expression was up-regulated at PMCs stage, while it was down-regulated at YP stage (Figure 6g).

In the second group, analysed genes were involved in different functions linked to pollen formation (Figure S4), i.e. pollen meiosis (*Sister chromatid cohesion* and *TomA108*), callose degradation (*Endo-1,3-beta-glucanase* and *AtTDF1-like*), exine formation (*LeGRP92*) and pollen germination (*pLAT52*). All these genes showed significant down-regulation in RNAi *SIMED18* lines, except for *Sister chromatid cohesion*, which was up-regulated at PMC stage and down-regulated at Tds and MP stages (Figure S4a). Considering the alterations in gene expression due to the silencing of *SIMED18*, altogether these results showed that the lack of *SIMED18* function promoted a significant down-regulation of genes mainly related to anther and pollen development, which correlated with the defects observed in tapetum and pollen formation.

SIMED18 complements the phenotypic defects of Arabidopsis med18-1 mutant

In Arabidopsis, mutations at the *MEDIATOR SUBUNIT 18* (*AtMED18*) cause pleiotropic phenotypic alterations affecting inflorescence structure, flower morphology, silique size, and flowering time (Figure 7a-d), indicating an essential role for *AtMED18* in the control of these developmental processes. Besides, Arabidopsis *med18* plants also showed alterations in stamen development and pollen maturation (Zheng *et al.*, 2013). To assess whether *SIMED18* could complement the developmental defects observed in *med18-1* mutants, several Arabidopsis transgenic plants overexpressing the tomato *MED18* orthologue were generated by using a 35S constitutive promoter (Figure 7, Figure S5a). Under long-day conditions, *med18-1* mutants displayed a late flowering phenotype that was fully rescued by the overexpression of *SIMED18* (Figure 7a,d,e). Indeed, expression of *FLOWERING LOCUS T (FT)*, *SUPPRESSOR OF CONSTANS OVEREXPRESSION 1 (SOC1)* and *FLOWERING LOCUS C (FLC)* genes was restored to similar transcriptional levels, or even higher than those showed by WT plants (Figure

S5b). Furthermore, the altered floral organ number and the decreased silique size observed in *med18-1* were complemented by the expression of tomato *MED18* orthologue in the Arabidopsis mutant (Figure 7b,c,f). Thus, flower identity genes such as *SEPALLATA 3* (*SEP3*), *PISTILLATA* (*PI*) and *AGAMOUS* (*AG*), which were down-regulated in Arabidopsis *med18-1* mutant flowers, showed transcriptional levels similar to WT when *SIMED18* was over-expressed in *med18-1* plants (Figure S5c). These results indicated that *SIMED18* fulfil the functional roles exerted by the Arabidopsis *MED18* gene.

DISCUSSION

Transcriptional activity of *SIMED18* is essential for tapetum degradation and pollen development

The *pod1* tomato T-DNA mutant is severely affected in flower and pollen development. Molecular cloning of the tagged gene proved that POD1 encodes the Mediator of RNA polymerase II transcription subunit 18 (MED18) supporting a functional role for this gene in reproductive development of tomato. GUS expression was detected in the stigma and pollen sacs (Figure 3e), which agrees with the spatial expression pattern detected by in situ hybridization (Figure 3c,d). These results support that transcriptional activity of SIMED18 is required during male gametogenesis. Although no morphological alterations were observed at early stages of pollen development of plants lacking POD1/SlMED18, microscopy analysis revealed changes during microgametogenesis (Figure 4), which involved a significant delay in tapetum degeneration as compared to WT (Figure 5). Moreover, from microspore stage onwards, defective anthers yielded a low amount of mature pollen grains, most of them being degenerated cells (near to 83%). Adequate sporophytic cell layer development is necessary to give rise to functional mature pollen in plants (Ma, 2005; Yuan et al., 2009; Zhou et al., 2011), being the tapetum the most important layer, since it supplies nutrients required for pollen development. Indeed, male sterility is normally associated with abnormal tapetum development, as occurs in the Arabidopsis ems1/exs and tpd1 mutants (Canales et al., 2002; Zhao et al., 2002; Yang et al., 2003), as well as in the tomato mutants affected either in SICER6 or Ms10³⁵ genes (Smirnova et al., 2013; Jeong et al., 2014). All of these mutants showed early tapetum degradation, the opposite effect to that observed in RNAi SIMED18 plants where tapetum degenerated later than in WT plants. However, the consequence in both cases was an abnormal pollen formation, which corroborates that timely coordinated degradation of the tapetum is a crucial step during microgametogenesis, and that this process requires the *SIMED18* function. Therefore, *SIMED18* may act as a link between sporophytic (tapetum tissue) and gametophytic (pollen development) tissues, an hypothesis that would be in agreement with the expression pattern of *SIMED18*, whose transcript levels were mainly found in anther primordia and pollen grains.

Recently, a genetic pathway including *DYT1-TDF1-AMS-bHLH89/91-MYB80* transcriptional cascade has been proposed to regulate tapetum ontogeny in Arabidopsis (Li *et al.*, 2017). In addition, *MS10*³⁵, the tomato *DYT1* homologue, has been suggested as an upstream regulator of that transcriptional cascade, whose function is necessary for proper meiosis and tapetum development (Jeong *et al.*, 2014). *MS10*³⁵ was significantly repressed in RNAi *SlMED18* anthers suggesting that *SlMED18* may promote tapetum development and degradation through direct or indirect regulation of *MS10*³⁵. In accordance with this, several tapetum development-related genes were down-regulated in RNAi *SlMED18* plants as occurred with *AtTDF1-like*, *AMS*-like, *Cysteine protease*, *Aspartic proteinase*, *TA29* and *MS1-like*, which support that tapetum functionality should be compromised by the lack of *SlMED18*.

Together, the results reported here provide strong evidence about the function of *SIMED18* in the transcriptional regulation of a subset of genes specifically required to develop mature pollen properly. In addition, hormones play a central role in male gametogenesis, and the function of *SIMED18* in the modulation of genes involved in hormonal pathways cannot be ruled out. Nevertheless, further research is required to better understand the hierarchical and functional relationships among the genes integrating the regulatory pathway involved in pollen development and their link with hormone pathways.

Functional divergence of SIMED18 gene

The Mediator complex is recognised as a central player in eukaryotic gene regulation. In Arabidopsis, several functions have been reported for the different subunits integrating this complex (Zheng *et al.*, 2013; Samanta and Thakur, 2015; Fallath *et al.*, 2017). Although such functions must be interconnected, subunits that integrate the same Mediator module seem to participate in the same biological process (Davoine *et al.*, 2017). *MED18* belongs to the head-module Mediator complex that was originally identified as a general transcription factor that stimulates basal RNA Polymerase II

transcription in yeast (Kornberg, 2005; Larivière et al., 2006). Later, yeast MED18 was described as a key element for proper elongation (Lee et al., 2013) and termination of transcription of a subset of genes (Mukundan and Ansari, 2011). In Arabidopsis, it has been proposed that MED18 regulates flowering time and floral organ formation through regulation of FLC and AG, respectively (Zheng et al., 2013). Expression levels of the floral repressor FLC were found up-regulated in med18-1 plants, which also agreed with the late flowering phenotype of these mutants. Concomitantly, decreased expression of FT and SOC1 floral integrators were detected in med18 (Zheng et al., 2013). In addition, the number of petals increased and the number of stamens decreased in *med18* plants, two features which were reminiscent of the floral phenotype of ag mutants (Chuang and Meyerowitz, 2000). In accordance, epistasis was observed in the med18-1 ag-1 double mutants and AG expression was down-regulated in med18-1 plants (Zheng et al., 2013). Interestingly, constitutive expression of SIMED18 in Arabidopsis med18 background rescues all the developmental defects displayed by med 18-1 plants related to flowering time and floral organ identity. Furthermore, SIMED18 also complements the altered pattern of expression of the key genes involved in the regulation of these processes (Figure S5). Thus, expression levels of AG, PI and SEP3 increased in complemented med18 plants, supporting that the MED18 tomato orthologue is able to regulate the expression of these identity genes in Arabidopsis and rescue the defects in flowering time and floral organ number observed in *med18* plants.

Overall results indicated that *SIMED18* shares significant biological function with Arabidopsis *MED18*. In addition to the altered floral organ number, Arabidopsis *med18* mutants also showed a delay both in stamen development and pollen maturation, which led to a reduction in seed set (Zhang *et al.*, 2013). These results indicate that *MED18* is also involved in the genetic control of male gametogenesis for this model species; although a detailed study of pollen ontogeny, similar to that conducted here with tomato, would be necessary in Arabidopsis to draw a final conclusion on the maintenance or divergence of MED18 function in these species. Nevertheless, in contrast to Arabidopsis *med18* mutants, *pod1* plants and *SIMED18* silenced lines showed no alterations in flowering time and reduced penetrance and variable expressivity of floral homeotic changes. As described for the Arabidopsis *med18* mutant, the occasional homeotic changes observed in the third floral whorl of *pod1* and RNAi *SIMED18* plants could be the consequence of the down-regulation of the tomato B-class identity key genes *TPI*, *TM6* and *SL* (Figure S1g). Altogether, these

observations suggest certain functional divergence between tomato and Arabidopsis MED18 orthologues. Interestingly, a single *MED18* gene has been found in the Arabidopsis genome (*AT2G22370*), whereas in tomato two paralogous *MED18* genes have been annotated (i.e, *Solyc03g046360* and *Solyc03g046370*) apart from the orthologous *SIMED18* gene (*Solyc06g008020*) reported here (Figure S6), indicating a different evolution dynamics for this family gene in these species.

Experimental procedures

Plant material

Tomato *pod1* mutant was isolated from a collection of T-DNA insertion lines generated by the enhancer trap vector pD991 in the tomato cv. Moneymaker (Pérez-Martín *et al.*, 2017). All tomato plants were grown under greenhouse conditions using standard practices with regular addition of fertilizers. The *med18-1* mutant in Columbia (Col) genetic background was kindly provided by Dr. David Oppenheimer (Zheng *et al.*, 2013).

Histochemical GUS staining

A GUS assay was carried out following the method described by Atarés *et al.* (2011). The resulting GUS-stained tissues were examined under a zoom stereomicroscope (MZFLIII, Leica). Three replicates of each sample were analysed.

Analysis of pollen viability

In vitro pollen viability was determined by the Tetrazolium staining method (Cottrell, 1948). Pollen grains from more than 30 flowers of each genotype were stained with 0.5% 2, 3, 5-triphenil tetrazolium chloride (TTC) (w/v) in 0.5 M sucrose in a wet chamber for 2h at 50°C in darkness (Viéitez Cortizo, 1952). Subsequently, pollen was visualised with a Nikon OPTIPHOT-2 optical microscope.

To evaluate *in vivo* pollen viability the Johansen's (1940) staining method for fluorescence microscope was assayed. Ten flowers from each WT and *pod1* plants were self-pollinated and reciprocally crossed. Two days after pollination, these flowers were collected and fixed in FAE (10% formaldehyde, 5% acetic acid and 50% absolute ethanol) for at least 24h, washed in water over night at 4°C, softened with NaOH 0.8 N during 6h and washed again in water over night at 4°C. Pollen tubes were stained with 0.1% aniline blue (w/v) in K₃PO₄ 0.1 N for 2h in darkness. Fluorescence was visualised

with a Nikon OPTIPHOT-2 optical microscope associated to HB-10101AF Mercury Lamp (Nikon).

Molecular cloning procedures

The DNA-blot hybridization was carried out following the protocol described by Yuste-Lisbona *et al.* (2016). Hybridization was performed with a chimeric probe constituted by the fused coding sequence of two genes, *NEOMYCIN PHOSPHOTRANSFERASE II* (*NPTII*) and *FALSIFLORA* (*FA*), the later was used as hybridization positive control.

The sequences flanking T-DNA insertion sites were isolated by a modified anchor-PCR according to the protocol described by Pérez-Martín *et al.* (2017). The cloned sequences were compared with SGN Database (http://solgenomics.net/) to assign the T-DNA insertion site on tomato genome.

Co-segregation of the T-DNA insertion site with the *pod1* phenotype was evaluated by PCR using i) the specific genomic forward and reverse primers (619a_genot_F/_R) to amplify the WT allele (without T-DNA insertion) and ii) one specific genomic primer (619a_genot_R) and the specific T-DNA right border primer (RB_pD991_F) to amplify the mutant allele (carrying the T-DNA insertion). The sequences of anchor-PCR and genotyping primers used are listed in Table S2.

Generation of transgenic lines

An interference RNA (RNAi) approach was performed to down-regulate candidate genes. To generate the RNAi *SIMED18* construct, a 164 bp fragment of *SIMED18* cDNA was cloned in sense and antisense orientation into the vector pKannibal (Wesley *et al.*, 2001), which was digested with *Not*I and the resulting fragment was cloned into the binary vector pART27 (Gleave, 1992) following the method described by Helliwell and Waterhouse (2003). Likewise, a 248 bp fragment of the *ZNHIT* cDNA was used to generate the RNAi *ZNHIT* construct. In addition, both *SIMED18* and *ZNHIT* genes were simultaneously inhibited by a double RNAi construct. For this purpose, the 164 bp fragment of *SIMED18* was amplified using RNAi-doble_F and RNAiMED18_R primers, and the RNAiZn_F and RNAi-doble_R primers were used to amplify the 248 bp fragment of the *ZNHIT* gene. Thereupon, both fragments were used as template in a PCR using the primers RNAiZn_F and RNAiMED18_R to join the PCR products resulting from the above PCR amplifications. The resulting PCR product was finally cloned in pART27 as described above.

To generate the overexpression gene construct (35S::SIMED18), the complete open reading frame of SIMED18 was amplified from S. lycopersicum cv. Moneymaker cDNA using 35S-Med_F and 35S-Med_R primers. The SIMED18 cDNA was cloned into the binary vector pROKII (Baulcombe et al., 1986). The overexpression construct 35S::SIMED18 was also used for genetic complementation of pod1 plants. The sequences of primers used in the generation of silencing and overexpression constructs are shown in Table S2.

Genetic transformation experiments were performed using *A. tumefaciens* (strain LBA4404) as described by Ellul *et al.* (2003). The ploidy levels in transgenic plants were evaluated by flow cytometry according to the protocol described by Atarés *et al.* (2011). Thus, diploid RNAi *SlMED18* (6 lines), RNAi *ZNHIT* (5 lines), double RNAi (3 lines) and 35S::*SlMED18* (3 lines) transgenic plants were selected for further phenotypic and expression analyses.

Microscopy analysis

Anther sections at key stages of microsporogenesis in the WT and transgenic genotypes were processed for light microscopy according to Jimenez-Lopez *et al.* (2016). Sections (7 μ m) were stained with a mix of toluidine blue/methylene blue, and observed in a Nikon Eclipse Ti-U microscope.

For the study of nuclei, pollen grains were released on a slide by squash from at least 20 anthers at four different stages and stained with 4′,6′-diamidino-2-phenylindole (10 ng·ml⁻¹, DAPI) in McIlvaine buffer (0.1 M citric acid, 0.2 M Na₂HPO₄, 1% Triton X-100, pH 4.0) according to the method previously described by Coleman and Goff (1985). Samples were incubated at room temperature for 15 min in darkness and examined using using an epifluorescence microscope Nikon OPTIPHOT-2 associated to HB-10101AF Mercury Lamp (Nikon).

Scanning electron microscopy (SEM) analyses were carried out as previously described by Lozano *et al.* (1998). Flowers from *pod1* mutant and WT were fixed in FAEG (10% formaldehyde, 5% acetic acid, 50% absolute ethanol and 0.72% glutaraldehyde) and stored in 70% ethanol. Critical point dried with liquid CO₂ in a critical point drier Bal Tec (Liechtenstein) CPD 030 was performed after dehydration of samples. Gold coat was applied in a Sputter Coater (Bal Tec SCD005). Samples were visualised using the scanning electron microscopy Hitachi S-3500N at 10kV.

Gene expression analysis

Total RNA was isolated using TRIZOL (Invitrogen) following the manufacturer's instructions from flowers at 5 different stages in relation to pollen cells development: PMCs (floral buds of 1-2 mm); Tds (floral buds of 3-4 mm); Mcs (flowers of 5-6 mm); YP (flowers of 7-8 mm); and MP (flowers at anthesis). cDNA was synthesized from 500 ng of total RNA using the M-MuLV reverse transcriptase (Fermentas Life Sciences) with a mixture of random hexamer and oligo(dT)₁₈ primers. Expression analyses were performed with three biological and two technical replicates. qRT-PCRs were performed with the SYBR Green PCR Master Mix (Applied Biosystems) kit using the 7300 Real-Time PCR System (Applied Biosystems). The housekeeping *Ubiquitine3* gene was used as control in all gene expression analyses. Specific primer pairs for each evaluated gene were described in Table S3. Results were expressed using the $\Delta\Delta$ Ct calculation method (Winer *et al.*, 1999) in arbitrary units by comparison with a data point from the WT samples.

In situ hybridization assays, tissue preparation, sectioning and transcript detection were carried out as described by Lozano et al. (1998). A POD1/SIMED18 probe was prepared using cDNA as template (200-pb fragment of the 3'UTR from the Solyc06g008010 gene). Antisense probe was synthesized using the DIG RNA labelling mix (Roche Applied Science). As negative control, sense RNA probe was hybridized with the same sections and no signals were observed under the hybridization and detection conditions used.

Genetic complementation of the Arabidopsis *med18-1* mutant

The overexpression construct 35S::SIMED18 was transformed in the Arabidopsis med18-1 plants (Zheng et al., 2013) by A. tumefaciens (strain C58C1) mediated transformation using the floral-dip method (Clough and Bent, 1998). The resulting 35S::SIMED18 med18-1 transgenic lines were selected on MS-glucose kanamycin-containing media plates. At least 10 independent transformants were evaluated phenotypically for different reproductive traits.

Controlled environmental conditions were provided by walk-in growth chambers at 22°C and 65% relative humidity. For the *in vitro* experiments, seedlings were cultured on agar-solidified MS medium. Plants were illuminated with cool-white fluorescent lights (120 µmoles/m²/s); Long-day (LD) conditions consisted of 16h light/8h dark and short-day (SD) conditions were 8h light/16h dark.

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Conflict of interest

The author(s) declare that they have no competing interests

Supporting information

- **Figure S1.** Scanning electron microscopy analysis and expression of B- and C-class genes in wild-type and *pod1* plants.
- **Figure S2.** Relative expression of *SIMED18* and *ZNHIT* genes in transgenic lines.
- **Figure S3.** Floral homeotic changes displayed by *pod1* mutants and *SIMED18* silenced lines.
- **Figure S4.** Relative expression of pollen marker genes in floral buds of wild-type (WT) and RNAi *SIMED18* plants.
- **Figure S5.** Relative expression levels of flowering time and floral organ identity genes are restored in 35S::SIMED18 med18-1 plants of Arabidopsis thaliana.
- **Figure S6.** Phylogenetic analysis of *MED18* homologue genes.
- **Table S1**. Relevant phenotypic traits measured in the wild-type, *pod1* mutant, and transgenic lines.
- Table S2. Primer sequences used for anchor-PCR, genotyping and transgenic constructs
- **Table S3.** Primer sequences used for qRT-PCR analyses

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Figure legends

Figure 1. Phenotypic and molecular characterization of *pod1* insertional mutant. (a,b) Compared to WT plants (left), both leaf (a) and flower (b) development were reduced in pod1 plants (right). (c,d) Stamens of WT plants yielded a normal amount of viable pollen stained with TTC (c), while stamens of pod1 plants produced little amount of viable pollen (d). (e,f) In vivo pollen germination analysis. WT pollen grains developed normally on pod1 stigmas allowing to complete pollination (e), however, pod1 pollen grains were unable to form pollen tubes on cross-pollinated WT flowers (f). (g) Tomato fruits yielded by pod1 plants were seedless (parthenocarpic) and displayed a significant reduction in size. Scales bars: 10 cm in (a); 0.5 cm in (b); 200 µm in (c) and (d); 100 µm in (e) and (f); and 1 cm in (g). (h) Southern blot analysis of genomic DNA digested with EcoRI (E) and HindIII (H) and hybridized with a chimeric probe including NPTII and FA genes (the latter used as hybridization positive control). (i) Genomic organization of the SIMED18 and ZNHIT genes. The T-DNA insertion event produced a 160 bp deletion (Δ_{160}) on the *POD1* genomic region of the mutant. Exons of *SIMED18* and ZNHIT are depicted as white and grey boxes, respectively. Promoter region of both genes is shown as a bold line between transcription start sites (ATG). (j) Co-segregation analysis of the T-DNA insertion and the pod1 mutant phenotype in 16 plants of the T2 population. T2 plants heterozygous (3, 4, 6, 7, 9, 10, 14 and 16) and homozygous for the WT allele (2, 8, 11 and 12) showed WT phenotype, while T2 plants homozygous for the mutant allele (1, 5, 13 and 15) displayed *pod1* mutant phenotype. Dashed circles indicate pod1 plants displaying mutant phenotype. (k) Quantitative real-time PCR assay for SIMED18 gene. (1) Quantitative real-time PCR analysis of ZNHIT gene. Asterisk denotes significant differences (Student's t-test, P < 0.05).

Figure 2. SIMED18 silencing plants phenocopy the pod1 mutant phenotype. (a-c) Phenotypic variation of reproductive traits observed in transgenic plants. Morphological features of flowers (a), pollen grains (b) and fruits (c) developed by RNAi SIMED18 and double RNAi silencing lines were similar to those of pod1 plants. In contrast, no differences were observed in RNAi ZNHIT silencing lines and in pod1 plants overexpressing SIMED18 (35S::SIMED18 pod1) compared to wild-type (WT) ones. (d) Comparison of the average relative expression of SIMED18 and ZNHIT genes in leaves from WT and transgenic lines. Scale bars: 1 cm in (a) and (c); and 100 μm in (b).

Figure 3. Expression of *SIMED18* in reproductive floral organs. (a) Time-course of *SIMED18* gene expression during flower and fruit development. (b-d) *In situ* hybridization assay of the *SIMED18* gene in tomato flower buds. While no signal was found with sense probe hybridization (b), transcript accumulation signals were found in the two inner whorls of early developed flowers (stage 5, according to Brukhin *et al.*, 2003) (c); later, *SIMED18* transcripts were mainly detected in pollen and ovules in flower buds at stage 8 (d). (e) *GUS* expression was found in stamens, stigma, ovules and at the bottom of petals and carpel of *pod1* flowers, but not in WT ones. S, sepal primordium; P, petal primordium; St, stamen primordium, C, carpel primordium; Ov, ovules; Pg, pollen grains. Scale bars: 100 μm in (b), (c) and (d); and 1 mm in (e).

Figure 4. Microscopy analysis of microsporogenesis and microgametogenesis in wild-type and RNAi *SIMED18* plants. (a-f) Morphological and histological features of anther sections during microsporogenesis of wild-type (a-c) and RNAi *SIMED18* (d-f) plants. No differences were detected at the following developmental stages: pollen mother cells prior to the onset of meiosis (a,d), meiocytes at telophase I stage (b,e), and tetrads (c,f). (g-r) Morphological and histological features of anther sections during microgametogenesis of wild-type (g-l) and RNAi *SIMED18* (m-r) plants; tissue sections were stained either with aniline blue (g-i and m-o) or DAPI (j-l and p-r). Comparative analyses were performed at the following developmental stages: vacuolated microspores (g,j *versus* m,p), young pollen grains (h,k *versus* n,q), and mature pollen grains prior to anther dehiscence (i,l *versus* o,r). Aw, anther wall; Ca, callose; Chr, chromatin; dN, degenerated nucleus; GN, generative nucleus; N, nucleus; PMC, pollen mother cell; T, tapetum; V, vacuole; VN, vegetative nucleus; asterisk, degenerated/altered microspore or pollen grain. Scale bars: 10μm.

Figure 5. Tapetum development is altered during microgametogenesis of RNAi *SIMED18* plants. (a-f) Morphological and histological features of anther sections focused on the tapetal layer at three stages before to anther dehiscence, i.e. vacuolated microspore (a,d), young pollen (b,e) and mature pollen (c,f). (g) Morphometric quantification of the tapetum tissue area (measured as a percentage of the whole anther locule) indicated a higher area of tapetum tissue RNAi *SIMED18* plants. Asterisks

denote significant differences (Student's t-test, P < 0.05). Aw, anther wall; T, tapetum. Scale bars: 10 μ m.

Figure 6. Expression of tapetum marker genes in wild-type and RNAi SIMED18 plants. qRT-PCR assay for SIMED18 (a), SISPOROCYTLESS/HYDRA (b), $MS10^{35}$ (c), AMS-like (d), AtMYB103-like (e), MS1-like (f), TGAS100 (g), bHLH89/91 (h), TA29 (i), $Cysteine\ protease$ (j), $Aspartic\ proteinase$ (k), and $Arabinogalactan\ protein$ (l) genes. The results show the averages and standard errors of three independent biological experiments and three technical replicates. Asterisks denote significant differences (Student's t-test, P < 0.05).

Figure 7. Expression of the tomato *SIMED18* rescues the phenotype defects displayed by Arabidopsis *med18-1* plants. Compared to wild-type plants (Columbia), no visible differences were found in the Arabidopsis *med18-1* plants overexpressing tomato *SIMED18* (35S::*SIMED18 med18-1*) with respect to flowering time (a), identity and number of floral organs (b), fruit size (c,f), and number of leaves before flowering (d,e).

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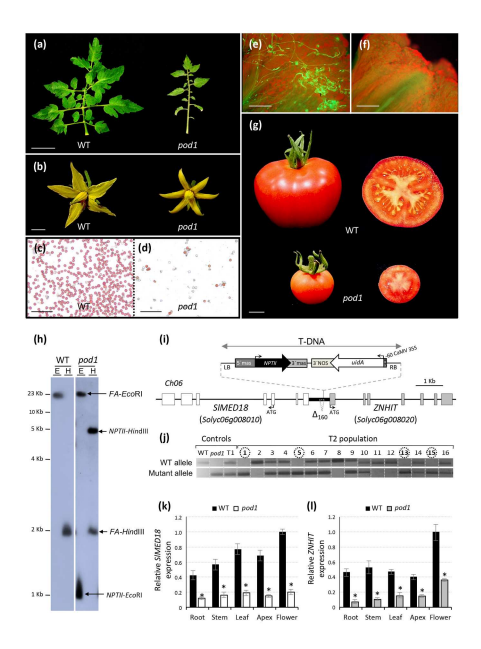


Figure 1. Phenotypic and molecular characterization of pod1 insertional mutant $161 x 219 mm \; (300 \; x \; 300 \; DPI)$

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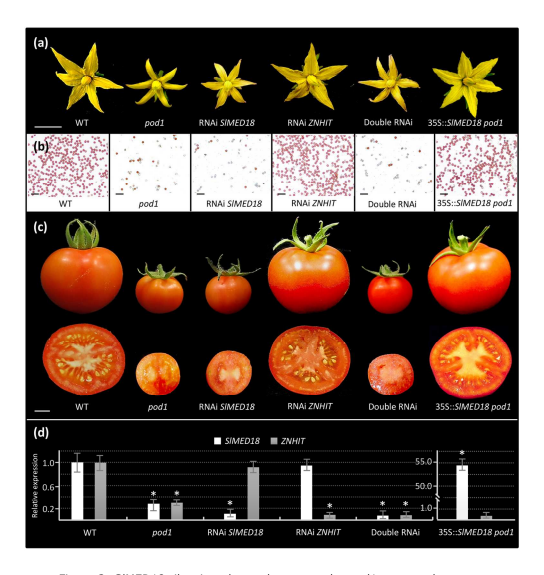


Figure 2. SIMED18 silencing plants phenocopy the pod1 mutant phenotype $136 \times 144 \text{mm} \ (300 \times 300 \ \text{DPI})$

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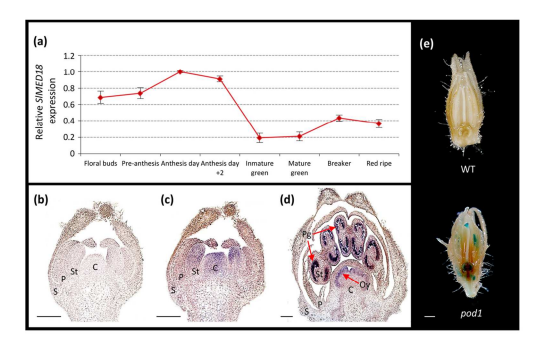


Figure 3. Expression of SIMED18 in reproductive floral organs 104x66mm~(300~x~300~DPI)

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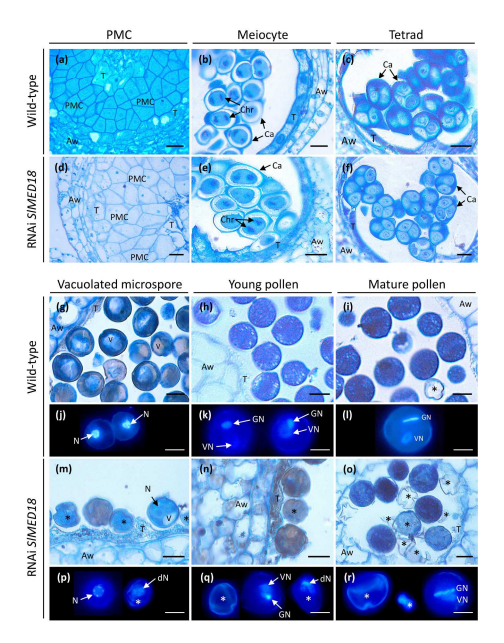


Figure 4. Microscopy analysis of microsporogenesis and microgametogenesis in wild-type and RNAi SIMED18 plants

221x289mm (300 x 300 DPI)

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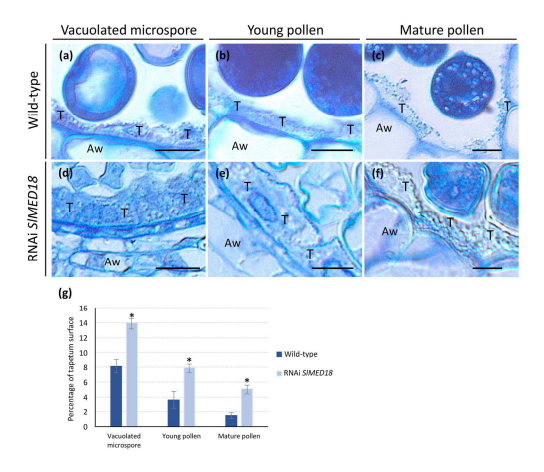


Figure 5. Tapetum development is altered during microgametogenesis of RNAi SIMED18 plants 147x129mm (300 x 300 DPI)

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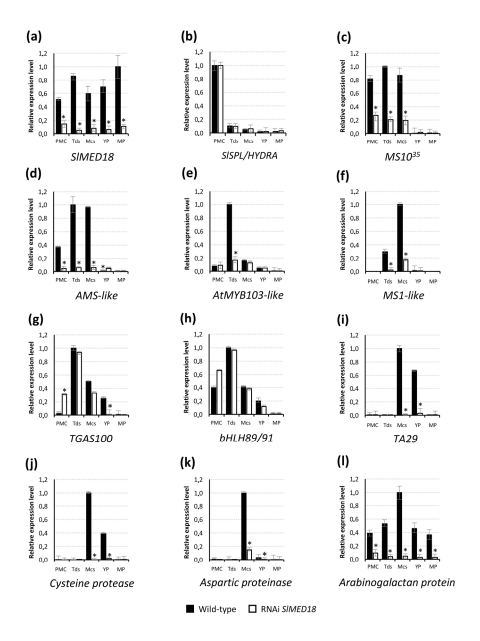


Figure 6. Expression of tapetum marker genes in wild-type and RNAi SIMED18 plants $179x241mm (300 \times 300 DPI)$

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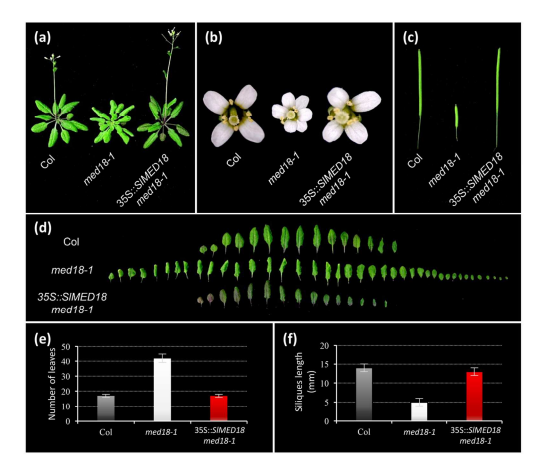
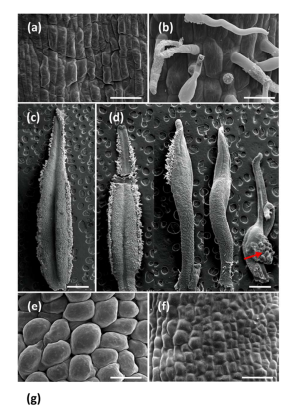


Figure 7. Expression of the tomato SIMED18 rescues the phenotype defects displayed by Arabidopsis med18-1 plants

120x104mm (300 x 300 DPI)

Significance statement

Pollination is a key development process in the life cycle of flowering plants. Genetic and molecular characterization of a tomato mutant have led to the identification of *POD1* gene encoding the Mediator complex subunit MED18 whose function is required for tapetum tissue degeneration, a crucial step for pollen development. Furthermore, we show that MED18 fulfils an essential role in tomato, ensuring proper gene regulation during pollen ontogeny.



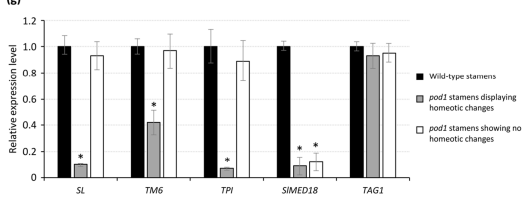


Figure S1

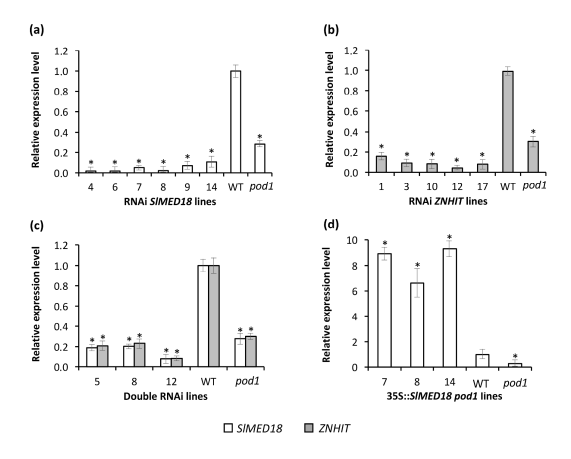


Figure S2

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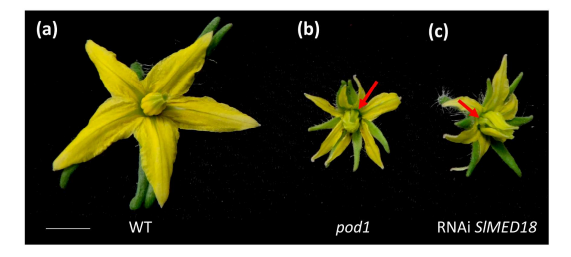


Figure S3

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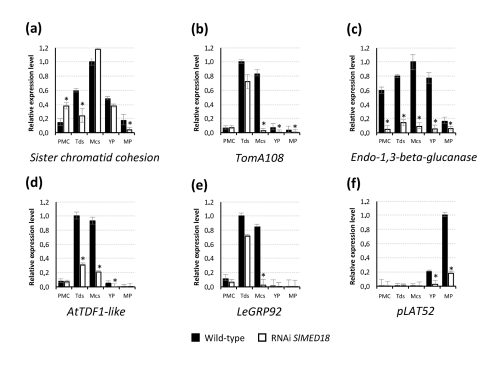
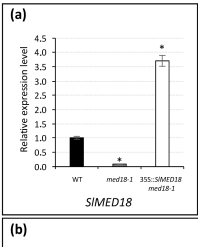
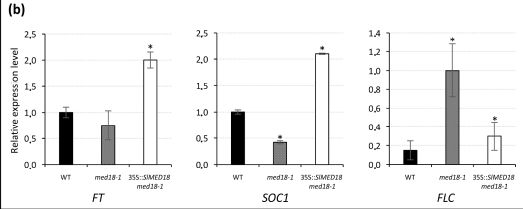


Figure S4





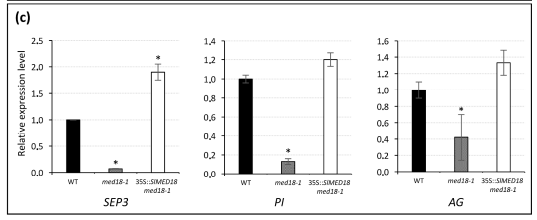


Figure S5



Figure S6

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Table S1

Category	Wild-type	pod1	RNAi ZNHIT	RNAi <i>SlMED18</i>	Double RNAi	35S::SIMED18 pod1
Fresh weight of leaf (g)	54.31 ± 12.92^{a}	19.68 ± 5.97^{b}	50.98 ± 11.03^{a}	21.33 ± 7.01^{b}	20.16 ± 6.45^{b}	55.12 ± 9.14^{a}

Values are expressed as the mean \pm standard deviation. Values followed by the same letter (a , b , or c) are not statistically different (Least significant difference test, P < 0.01).

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Length of leaf (mm)	491.83 ± 39.54^{a}	347.5 ± 46.58^{b}	477.91 ± 42.22^{a}	360.1 ± 37.11^{b}	355.8 ± 54.22^{b}	489.25 \pm
						41.69 ^a
Length of petiole (mm)	58.40 ± 1.08^{a}	37.77 ± 3.20^{b}	56.16 ± 2.23^{a}	36.16 ± 3.84^{b}	38.01 ± 2.99^{b}	54.10 ± 1.56^{a}
Length of secondary petiole (mm)	27.01 ± 2.40^{a}	12.50 ± 1.20^{b}	25.94 ± 3.56^{a}	11.85 ± 3.72^{b}	14.31 ± 3.02^{b}	24.98 ± 3.11^{a}
Length of sepal (mm)	9.81 ± 0.55^{a}	9.46 ± 2.04^{a}	9.47 ± 1.26^{a}	9.77 ± 1.85^{a}	9.97 ± 2.05^{a}	9.63 ± 1.67^{a}
Length of petal (mm)	14.80 ± 1.30^{a}	13.16 ± 1.57^{b}	14.56 ± 0.73^{a}	12.85 ± 1.60^{b}	13.04 ± 1.16^{b}	14.80 ± 1.30^{a}
Length of stamen (mm)	8.46 ± 1.60^{a}	6.64 ± 1.30^{b}	8.50 ± 0.49^{a}	7.71 ± 1.03^{c}	7.67 ± 0.83^{c}	8.18 ± 1.00^{a}
Length of pistil (mm)	7.18 ± 0.55^{a}	5.26 ± 0.50^{b}	7.68 ± 0.83^{a}	6.07 ± 1.09^{c}	6.19 ± 1.31^{c}	7.34 ± 0.91^{a}
Diameter of peduncle (mm)	0.71 ± 0.06^{a}	1.20 ± 0.20^{b}	0.75 ± 0.23^{a}	1.00 ± 0.16^{c}	0.96 ± 0.20^{c}	0.68 ± 0.15^{a}
Diameter of carpel (mm)	1.63 ± 0.24^{a}	1.63 ± 0.26^{a}	1.58 ± 0.32^{a}	1.60 ± 0.22^{a}	1.66 ± 0.27^{a}	1.59 ± 0.28^{a}
Axial diameter of fruit (mm)	47.97 ± 2.82^{a}	28.75 ± 2.06^{b}	52.50 ± 1.32^{a}	27.12 ± 1.30^{b}	28.68 ± 2.10^{b}	50.07 ± 3.05^{a}
Equatorial diameter of fruit (mm)	59.44 ± 4.28^a	31.40 ± 5.83^{b}	62.33 ± 2.88^{a}	30.88 ± 1.52^{b}	32.03 ± 1.79^{b}	61.28 ± 3.87^{a}
Fresh weight of fruit (g)	130.75 ± 19.09^{a}	18.90 ± 5.90^{b}	134.59 ± 17.92^{a}	19.61 ± 1.95^{b}	20.07 ± 1.89^{b}	133.05 \pm
						18.51 ^a
Relative amount of pollen (%)	98.30 ± 5.20^{a}	17.60 ± 9.10^{b}	98.20 ± 7.30^{a}	14.30 ± 6.20^{b}	16.30 ± 6.40^{b}	93.90 ± 5.80^{a}
Pollen viability (%)	97.60 ± 3.60^{a}	28.70 ± 12.40^{b}	96.20 ± 4.10^{a}	18.70 ± 8.80^{b}	15.70 ± 9.90^{b}	96.50 ± 3.60^{a}

Table S2

(a) Primers used for anchor-PCR					
Primer name	Primer sequence 5`-3`				
Ad1	CTAATACGACTCACTATAGGC				
Ad2	CTATAGGGCTCGAGCGGC				
Ad3	AGCGGCGGGAGGT				
ARB-1	ACAGTTTTCGCGATCCAGAC				
ARB-2	GGTCTTGCGAAGGATAGTGG				
ARB-3	CTGGCGTAATAGCGAAGAGG				
ALB-1	TTGGCGTGTCAGCGTATCTA				
ALB-2	ATCGGTCTCAATGCAAAAGG				
ALB-3	ATAATAACGCTGCGGACATCTAC				
(b) Primers used for PCR ge	(b) Primers used for PCR genotyping analysis				
Primer name	Primer sequence 5`-3`				
619a_genot_F	TCAACAGTAAAAACGAGCCAAA				
619a_genot_R	GGATGAAGCAATTGGGACAC				
RB_pD991_F	TACAACGTCGTGATGGGAAA				
(c) Primers used for generation of transgenic lines					
Primer name	Primer sequence 5`-3`				
RNAiMED18_F	tctagactcgagTTCTTTGGCTCGTTGTTATCG				
RNAiMED18_R	atcgatggtaccTTGAACAGTTGAAGCAATCTCA				
RNAiZn_F	tctagactcgagTGGAGAAGTTCAACAGGATACG				
RNAiZn_R	atcgatggtaccAATGGAAGCAGAAGCAGAGG				
RNAi-doble_F	gctagtcgcgaggcTTCTTTGGCTCGTTGTTATCG				
RNAi-doble_R	gcctcgcgactagcAATGGAAGCAGAAGCAGAGG				
35S-Med_F	ggtaccAAAAATCTCTCTTTGGCTCGTTT				
35S-Med_R	gagctcCTGCTGTGCTTTGTTTTTCG				

^{*} In lowercase is shown the endonuclease site introduced in the primer sequence

Table S3

Locus	Forward primer sequence 5'- 3'	Reverse primer sequence 5'- 3'	SGN id
SlAMS-like	TGCAGAGATGTTATGTTTCAGCATC	TCGTCTCTGTCTCTTTCTCCTTCTG	Solyc08g062780
SlMS1-like	TTGTGTCAATGGATCATTGGAAAC	AACCTCTTGCCTAGACACCCATC	Solyc04g008420
Slendo13bGl	GATCCAATGTGGGGAAGAAA	CCACAAATCAAAGCACCTCA	Solyc03g046200
SlpLAT52	AAGGTGTGACTGATAAAGATGGC	AACCCAACTCATCAAGAGCTTC	Solyc10g007270
SlbHLH89/91	TCCATGGATGGTAGTGATGC	TCGACAATCCGAACATCAAC	Solyc01g081100
SlCysProt	ATTGGTGTCGATTGGAGGAAG	CAAATGCACTTTCCATAAACCC	Solyc07g053460
SlAspProt	GTGATATTAATTGGCTTCAATGTGAACC	ATACTCGCCGGAACCTGTAACATC	Solyc06g069220
SITA29	AAGATTTTAACCATGAACTTCTTC	ACATTCTTCAGTGTCACATACATC	Solyc02g078370
SlTomA108	ATGCAATTAGGAGCCTTGATTC	CAGTTCCAGTTCCTGTTCCG	Solyc01g009590
SlTGAS100	TATATAGACATGGCAATGAAATGGC	AGTCAAGACAACGATCAAGAATGC	Solyc06g064480
SlSisterCC	CATTGGCTTTCAGAGCTTCC	GCAGCAGAAAGCGAAATTCT	Solyc03g116930
LeGRP92	ATGCAATTAGGAGCCTTGATTC	CAGTTCCAGTTCCTGTTCCG	Solyc02g032910
SlMS10-35	AGATCTCTCTGATTCGATTAGCTTCAG	TCTTGAAATGGAAGCAACTCAGG	Solyc02g079810
SlTDF1-like	GGTAATTGGGCAACCATGTC	TTGAGGCGTAAAGCTGTCCT	Solyc03g113530
SlMYB103-like	TGCTGAGGAAGATGCAAAAA	GGTCCATCTCAGCCTACAGC	Solyc03g059200
SlArabinogalProt	CCTTTTCATTCTGGGGTGAC	CGTCACTAACAACCTTTGAACG	Solyc11g072780
619a ZFinger	AGCTGTGTAAGGCGTGCTCT	ACAGCTATATCGATACACTTCGTTT	Solyc06g008020
SIMED18	TCTCTGATGTCTGATGGTGGA	GAAGGAGAATGGCGAAATAC	Solyc06g008010
AtMED18	CGAACCCACATGGACGGTTAAA	AGATGAAACAGCAGCAGCGACT	AT2G22370
FT	CATCGTGTCGTGTTTATATTGTTTCG	CCTCCGCAGCCACTCTCC	AT1G65480
SOC1	ACTCTTGGGAGAAGGCATAGGA	TGGGCTACTCTCTTCATCACCT	AT2G45660
FLC	TCACCTTCTCCAAACGTCGCAA	TGAGTTCGGTCTTCTTGGCTCT	AT5G10140
SEP3	ACGCCTTACAGAGAACCCAAAGGA	TTTGTCTCAGTCAGCATGCGTTCC	AT1G24260
PI	ACCAATGCTCCTCTTCTTGTTCTTC	ACTCTGTTGTTTGCGTTCTCTATCC	AT5G20240
AG	CGAGTATAAGTCTAATGCCAGGAG	GAGTAATGGTGATTGTTAGGTTGC	AT4G18960
TM6	GGAAAAATTGAGATCAAGAAG	TCAGGAGAGACGTAGATCAC	Solyc02g085480
TPI	TGGGGAGAGGTAAAATAGAG	GTAGATTTGGCTGCATTGGC	Solyc06g059970
SL (AP3)	ATGGCTCGTGGTAAGATCCAG	TCAACCTAGAGCAAAAGTAG	Solyc04g081000
UBC21	CTTGGACGCTTCAGTCTGTG	TGAACCCTCTCACATCACCA	AT5G25760
Ubiquitine3	CACACTTCACTTGGTCTTGCGT	TAGTCTTTCCGGTGAGAGTCTTCA	Solyc01g056940