CYTOLOGICAL OBSERVATION OF SOLANUM PIMPINELLIFOLIUM L. MICROSPORE DEVELOPMENT

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Abstract

Tomato (Solanum lycopersicum L.) is one of the most important horticultural crops worldwide, and has also been adapted as a model plant in a wide range of research disciplines. However, stamen development in tomato is less known than other model species, such as Arabidopsis thaliana and Oryza sativa L. (rice). In order to understand tomato flower development in more detail, the key goal of this study was to establish a precise correlation between visual morphological features and cytological events. To this end, we characterized a wild tomato species, S. pimpinellifolium (accession LA1585), to define stamen developmental stages using semi-thin sectioning. Based on distinct stages of flower development, S. pimpinellifolium anther/stamen development was divided into ten stages, and characteristic morphological and cytological features in stamens at each developmental stage were identified. In this way, flower characteristics and microspore development were correlated. In addition, specific cytological key events in S. pimpinellifolium were compared with those in A. thaliana and rice at corresponding stages, as well as with those in domesticated tomato (S. lycopersicum L.), and the differences are discussed.

Key words: Solanum pimpinellifolium, Cytological observation, Tomato

Introduction

In angiosperms, microsporogenesis occurs within the anther, which has a four-lobed structure (Goldberg et al., 1993), and microspore cell-specification and differentiation events within the anther give rise to mature pollen grains that contribute to male reproductive success. The formation of microspores, which develop into the male gametophyte in land plants, has been studied in detail in a number of model species, including Arabidopsis thaliana (Sanders et al., 1999), Zea mays (Barnabas et al., 1987), Oryza sativa (Zhang & Wilson, 2009; Zhang et al., 2011) and Nicotiana tabacum (Koltunow et al., 1990), as well as Ginkgo biloba (Sun et al., 2015) and family Lamiaceae (Firdous & Bhadula, 1988; Sanders et al., 1999; Zhang & Wilson, 2009), and it is not known whether there are differences in microspore development between tomato and the better studied model species.

The development of tomato microspores has previously been divided into eight stages based on bud length (Sawhney & Bhadula, 1988), and detailed studies of pollen development have revealed ultrastructural differences from the tetrad stage to pollen maturation (Polowick & Sawhney, 1992; 1993a; 1993b), as well as the developmental transition of microspores and flowers in wild type tomato genotypes and tomato mutants (Sawhney & Bhadula, 1988; Polowick & Sawhney, 1995; Brukhin et al., 2003). However, to our knowledge, microspore development has only been studied in domesticated tomato (S. lycopersicum) or S. lycopersicum var. cerasiforme, which has only a few flowers on each inflorescence (Polowick & Sawhney, 1993b; Brukhin et al., 2003). Consequently, within the genus Solanum, correlations between morphological features of the visible buds/flowers and microspore development over a defined time scale are not well defined, unlike the situation for rice and A. thaliana (Ma, 2005; Zhang et al., 2011; Chang et al., 2011; Yousuf et al., 2006).

In this regard, in order to effectively characterize the chemical, physiological and structural differences as well as the regulation and expression of stamen development associated genes, it is important to establish a precise correlation between the morphological features and key cytological events at each developmental stage (Yousuf et al., 2009). In the present study, we characterized microspore development at different developmental stages in wild tomato relative, S. pimpinellifolium (LA1585), since this species has ten or more buds/flowers in each inflorescence before anthesis, which allows the accurate staging of microspore development over a well-defined time scale. In addition to being the wild relative that is most closely related to cultivated tomato, S. pimpinellifolium can also...
cross freely with the domesticated species (Miller & Tanksley, 1990; Alvarez et al., 2001; Marshall et al., 2001), and it has the important characteristic that anthesis can occur over the entire year. In this study, changes in S. pimpinellifolium cytology during microspore development were observed in semi-thin sections and key features of S. pimpinellifolium microspore development were compared with those of the two experimental model plants, A. thaliana and rice, as well as with the equivalent features in S. lycopersicum.

Materials and Methods

Sample collection: Seeds of S. pimpinellifolium (accession LA1585) were kindly provided by the Tomato Genetic Resource Center (TGRC, http://tgrc.ucdavis.edu), and sown on petri dishes with 1/2 MS medium (Murashige and Skoog, MS basal medium 2.21 g/L, sucrose 20 g/L, phytogel 2.6 g/L, Sigma, pH 5.8). The plates were then incubated at 25°C, with a photoperiod of 16h light /8h dark to promote germination and seedlings were then transferred to sterile plastic pots (diameter 7.5 cm) containing seedling soil (20% vermiculite, 20% turf soil and 60% garden soil) and allowed to grow until four true leaves had fully spread. The plantlets were then transferred into 5 L plastic pots with nutrient soil (50% peat, 20% vermiculite and 30% soil) and grown in a greenhouse (25°C day/18°C night, 70% air humidity and natural light) in the School of Agriculture and Biology, Shanghai Jiao Tong University, China.

Inflorescences and flowers were sampled from healthy blossoming plants where only one open flower and small buds (generally more than ten) were present. Harvested inflorescences and flowers were brought back to the laboratory in an ice box. Photographs of the different developmental stages were taken with a digital camera (SONY Cyber-shot DSC-W350, Japan).

Histological analysis: The anthers or intact small buds at different development stages were harvested from the selected inflorescences using sterile forceps before being fixed in FAA (50% ethanol, 5.0% glacial acetic acid, 3.7% formaldehyde) overnight, and subsequently dehydrated in a graded ethanol series (2×50%, 60%, 70%, 85%, 95%, 3×100%) for 30 min. Sample were then embedded in Technovit 7100 resin (Emgrid Australia Pty Ltd, Australia), and sectioned (2 μm) using a microtome (Leica RM2265, Germany). Transverse sections of anthers derived from different development stages were stained with 1% toluidine blue for 20 s and observed under a microscope (Olympus BX51, Japan).

Results

Flower features: S. pimpinellifolium (accession LA1585) is a small perennial bush with a slim stem and a uniparous scorpoid cyme, with more than ten flowers per inflorescence (Fig. 1A). Generally, a perfect flower consists of five sepals, alternating with five petals and a style surrounded by stamens, where five anthers are connection laterally to form a flask-shaped cone with an elongated sterile tip at the apex (Fig. 1A and B). The long style, which projects from the anther cone in S. pimpinellifolium at anthesis, is distinct from that of the cultivated tomato which is embedded within the anther cone (Fig. 1A and C). Ten buds/flower, including one open flower and nine buds, which harvested from same inflorescence and ranged from 3.5 mm to 11.5 mm in length, and these were divided into ten stages of anther development (Fig. 1C). Landmark cytological events for each S. pimpinellifolium stamen developmental stage were noted over a time course starting from the emergence of the U-shaped sporogenous cell region to the release of mature pollen from the anther locules (Fig. 2 and Fig. 3).

Fig. 1. Buds/flowers and inflorescence of Solanum pimpinellifolium.
A: a scorpoid cyme inflorescence; B: a perfect flower with sepal, petal, anther and carpel; C: developing buds/flower from stages 1 to 10. (Bars=10mm).
Stage 1: the early sporogenesis stage: At stage 1, the bud length was approximately 3.5 mm, and a yellow colored corolla showed the first signs of emergence from the enwrapping sepals (Fig. 1C). The fully developed anther was divided into a bilateral structure (two half anthers) with an anther wall, connective tissue, vasculature and four locule regions. Each anther half included two separate locules (the outer being larger than the inner), and each locule region was filled by two layers of sporogenous cells (Sp) and the clevis, or U-shaped locule region was surrounded by the tapetum (Fig. 2A). The anther wall differentiated into four distinct cellular layers: an epidermis layer (E), an endothecium layer (En), a middle layer (ML) and a tapetum layer (T), each comprising a single layer of cells (Fig. 2, stage 1; Fig. 3, stage 1).

Stage 2: the late sporogenesis stage: At stage 2, bud lengths reached 4.5 mm, the corollaceous top was distinctly exposed from the calyx, and sepals initiated opening (Fig. 1C). The distinct E, En, ML and T cell layers in the anther showed continued development and cell number gradually increased in both the E and En layers due to periclinal division and the cell number of the ML cell layer also increased, giving rise to three or four layers due to anticlinal division. The ‘U’-shaped Sp cells initiated differentiation into microspore mother cells (MMC), and those MMCs were stained with toluidine blue to reveal a distinct blue-black region. At this stage, the tapetum cell layer was thicker and intact (Fig. 2, stage 2; Fig. 3, stage 2).
Stage 3; the MMC formation stage: At stage 3, sepals were only half the length of the corolla (Fig. 1C). The MMCs showed evidence of vacuolation, with an enlarged locular space and they also entered meiosis. A concave stomium region (Str) arose at both sides of the anther, the outer tapetum layer became gradually thicker and the emergence of binucleate tapetum cells was the most notable characteristic of this stage. The outer E cells increased in size and were observed to be larger than the inner E cells (Fig. 2, stage 3; Fig. 3, stage 3).

Stage 4; the tetrad stage: At stage 4, all flower organs continued developing and some sepals expanded further (Fig. 1C). The MMCs underwent one round of meiosis and mitosis and differentiated into tetrads that were coated by callose. Concomitantly, the cells of the tapetum layer, which supply nutrients to the developing microspores, began to burst and form an amoeboid shape, in contrast to the previous uniform appearance of the cells in the tapetum layer (Fig. 2, stage 4; Fig. 3, stage 4).

Stage 5; the early uninucleate microspore stage: At stage 5, all sepals were fully open (Fig. 1C). Free haploid microspores (Msp) were released from the irregularly shaped callose enwrapped tetrads in each of the four locules, the nucleolus was densely stained by toluidine blue and most of the Msps had a larger vacuole. The tapetal layer showed evidence of continued degeneration (Fig. 2, stage 5; Fig. 3, stage 5).

Stage 6; the late uninucleate microspore stage: At stage 6, buds reached approximately 8.5 mm in length, and the lower part of the corolla began to swell as the anther surrounded by petals developed quickly (Fig. 1C). Microspores were converting from their irregular shape into angular structures, and the blue-black toluidine blue staining associated with the nucleus was localized in the limbic regions of the microspores. The cells of the tapetum layer degenerated further and only some patches remained (Fig. 2, stage 6; Fig. 3, stage 6).

Stage 7; the vacuolated microspore stage: At stage 7, the sepals only reached one-third of the length of the corolla, and the lower part of the corolla expanded further (Fig. 1C). The angular microspores adopted a more spherical shape and both more extensive vacuolation and a high density of nuclear substances were observed in the microspores. Although remnants of a very thin tapetum layer were still detected, the septum layer became considerably thinner and only a remnant wall of broken tapetum cells remained. Septum (Sp) tissues, which were located between the two locules in each half of the anther, began to disappear due to cell degradation (Fig. 2, stage 7; Fig. 3, stage 7).

Stage 8; the binucleate microspore stage: At stage 8, buds were 10.5 mm in length and the corolla size increased to become equal to that of an open flower, while the top of the corolla was still closed (Fig. 1C). The large vacuole within the microspores disappeared, or was converted into many smaller vacuoles, and the irregularly shaped microspores were present as spherical binucleate pollen grains (PGs), which stained blue-black with toluidine blue. The tapetum layer had entirely disappeared from the locules and the cells of the Sp were completely absorbed or degraded, resulting in an opening between the locules. The two locules started to form one larger locule in each half of the anther, and the PGs were able to freely move within this larger locule. Eventually, anthers with four locules converted into anthers with two larger locules. Meanwhile, the cells of the stomium (St) also started to degenerate and form a thin layer of only one or two cell layers (Fig. 2, stage 8; Fig. 3 stage 8).

Stage 9; the early mature pollen stage: At stage 9, the buds were approximately 11.0 mm in length, and the flat top of the yellow corolla had opened appreciably (Fig. 1C). The smaller vacuoles within the pollen grains disappeared, and mature pollen grains, which stained very strongly blue-black with toluidine blue, were present. Three distinct colporates/germinal apertures were observed on each of the spherical pollen grains. Stomium cells were gradually absorbed and only one or two layers cells remained, but the anther sporangium was intact (Fig. 2, stage 9 and Fig. 3, stage 9).

Stage 10; the late mature pollen stage: At stage 10, the corolla had opened fully and the bright-yellow petals curled down in the direction of the peduncle, exposing the bright-yellow anther cone and the style projecting from the anther cone (Fig. 1C). Finally, the anther sporangium dehisced due to degeneration of the stomium cells, and mature pollen grains were released from the locules, where only a few pollen grains remained (Fig. 2, stage 10; Fig. 3, stage 10).

Discussion

Microspore development in angiosperms starts with the formation a stamen primordium and ends with the maturation of the pollen grains and this process has been characterized and staged in a number of model species (Sanders et al., 1999; Brukhin et al., 2003; Zhang & Wilson, 2009). Tomato has emerged as a model plant for researchers in a wide range of research disciplines (Li & Chetelat, 2010; Yeats et al., 2010; Bedinger et al., 2011; Matas et al., 2011), and the precise definition of microspore developmental stages is therefore essential for studies of tomato reproductive and developmental biology. In this current study, the wild tomato species, S. pimpinellifolium (LA1585), was chosen for a detailed study of microspore development stages, since it is taxonomically placed between domesticated tomato and other wild relatives, with the closest relationship being to domesticated tomato (Zuriaga et al., 2009). To our knowledge, domesticated tomato has to date served as the primary experimental material for the analysis of tomato microspore development (Mazzucato et al., 1998; Sawhney & Bhadula 1998; Brukhin et al., 2003); however, domesticated tomato differs from S. pimpinellifolium in the number of flowers on each inflorescence, where ten or more buds/flower are present in S. pimpinellifolium, compared to only a few in S. lycopersicum. This characteristic of S. pimpinellifolium facilitates a fine dissection of stamen development over a time course.
<table>
<thead>
<tr>
<th>Tomato anther stage</th>
<th>Tomato anther length (mm)</th>
<th>A. thaliana anther stage</th>
<th>Rice anther stage</th>
<th>Major events and morphological markers</th>
<th>Tissues present</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.63</td>
<td>3 – 4</td>
<td>3–4</td>
<td>Emergence of the U-shaped sporogenous cell region and formation of the four-lobed anther pattern. The vascular region was clearly visible.</td>
<td>E, En, M, V, T, C, Sp</td>
</tr>
<tr>
<td>2</td>
<td>2.64–3.33</td>
<td>4–5</td>
<td>5–6</td>
<td>The number of MMCs increased due to anticlinal division; the middle layer also increased but due to periclinal division.</td>
<td>E, En, M, V, T, C, MMC</td>
</tr>
<tr>
<td>3</td>
<td>3.34–3.81</td>
<td>6</td>
<td>6–8a</td>
<td>MMCs initiated genomic DNA replication, and entered meiosis and mitosis. The tapetum layer became thicker, and the size and number of tapetum cells increased due to anticlinal division and growth, and generated binucleate tapetum cells. Stomium regions emerged between two locules in each anther half and the middle layer initiated degradation.</td>
<td>E, En, M, V, T, C, MMC, SrR</td>
</tr>
<tr>
<td>4</td>
<td>3.82–4.19</td>
<td>7</td>
<td>8b</td>
<td>MMCs completed meiosis and generated free microspore tetrads within each locule. Only one thin layer of the middle layer was present. The tapetum layer started to degrade.</td>
<td>E, En, M, V, T, C, SrR, Tds</td>
</tr>
<tr>
<td>5</td>
<td>4.20–4.58</td>
<td>8</td>
<td>9–10</td>
<td>Individual microspores, surrounded by a callose cell wall, were released from the tetrads, and vacuolated into irregular shapes, and underwent one round of mitosis. The tapetum layer became thinner due to the degradation of cells and only one of layer cells remained as the middle layer.</td>
<td>E, En, V, T, C, SrR, Msp</td>
</tr>
<tr>
<td>6</td>
<td>4.59–5.32</td>
<td>8</td>
<td>10</td>
<td>Microspores were further vacuolated and generated three-celled microspores. The tapetum degraded further. The shapes were similar to those seen in A. thaliana but not rice.</td>
<td>E, En, V, T, C, SrR, Msp</td>
</tr>
<tr>
<td>7</td>
<td>5.33–6.66</td>
<td>9–10</td>
<td>11–12</td>
<td>Microspores generated an exine wall, and changed from an irregular to a spherical shape. Septum tissue became thin. Stomium tissue differentiation began and only a few remnants of the tapetum remained.</td>
<td>E, En, V, T, C, SrR, Msp</td>
</tr>
<tr>
<td>8</td>
<td>6.67–7.93</td>
<td>12</td>
<td>13</td>
<td>The microspore exine walls generated numerous granules that resulted in a rough surface on the pollen grains. Septum cells fully degraded resulting in an opening between the two locules in each anther half. Stomium cells continued to degrade.</td>
<td>E, En, V, C, SrR, Mp</td>
</tr>
<tr>
<td>9</td>
<td>7.94–8.72</td>
<td>12</td>
<td>13</td>
<td>Three colporates/germinal apertures were presented on the surface of the pollen grains. Only one layer of cells remained in the stomium.</td>
<td>E, En, V, C, SrR, Mp</td>
</tr>
<tr>
<td>10</td>
<td>8.73–9.06</td>
<td>13–14</td>
<td>13–14</td>
<td>Stomium cells were fully degraded and a cleft emerged on the outside of each anther half. Mature pollen grains were released from the anthers.</td>
<td>E, En, V, C, Mp</td>
</tr>
</tbody>
</table>

*a BM, binucleate microspore; E, epidermis; En, endothecium; ML, middle layer; MMC, microspore mother cells; MSp, microspores; PG, pollen grains; Sm, septum; Sp, sporogenous cells; Sr, stomium; SrR, stomium region; T, tapetum; TdR, tetrads; V, vascular bundle; Va, vacuoles.

*b The A. thaliana and rice data in the table are based on Sanders et al. (1999) and Zhang and Wilson (2009), respectively.
Based on the bud/flower length, which ranged from 3.5 mm to 11.5 mm, or the flower number, the stamen development was divided into ten stages (Fig. 1C). The landmark morphological and cytological events associated with each developmental stage were determined and compared to those of *A. thaliana* and rice at the corresponding stages (Table 1). We propose that the emergence at stages 1 and 2 in *S. pimpinellifolium* of specific sporogenous cells (Sp) and their differentiation into MMCs is equivalent to stages 4 and 5 in both *A. thaliana* and rice (Table 1). The U-shaped region of the Sps and MMCs was only found in tomato since connective tissue projected inward to the locules in the anther, which was not the case in *A. thaliana* and rice (stages 1 and 2 Fig. 2; Sanders *et al.*, 1999; Zhang & Wilson, 2009). We also observed two locules in each half of the anther in *S. pimpinellifolium*, which were different in size, with the outer locule (close to the petals) being larger than inner (close to pistil) locule (Fig. 2). This is similar to what is seen in *A. thaliana*, but not in rice (Sanders *et al.*, 1999; Zhang & Wilson, 2009). This feature might be related to the specific flower structure in tomato and Arabidopsis that sepals and petals closely pack the anther cone, but it maintains unknown whether extensively existing in dicotyledonous plants. Three colporates/germinal apertures were detected in *S. pimpinellifolium*, but not in *A. thaliana* and rice (stages 8 and 9 in Fig. 2; stages 8 and 9 in Fig. S1; Sanders *et al.*, 1999; Zhang & Wilson, 2009). In addition, we also observed that stage of the tapetum layer degradation was different in *S. pimpinellifolium* from what has been reported for *A. thaliana* and rice (Sanders *et al.*, 1999; Zhang & Wilson, 2009). Indeed, numerous differences in stamen development were seen between tomato and the two other model species, which presumably lend some evolutionary advantage; however, the adaptive significance of these features remains unknown.

*S. lycopersicum* flower development, and specifically that of the stamen and carpel, has been divided into twenty stages (Brukhin *et al.*, 2003), ranging from the sepal primordial stage (stage 1) to the anther dehiscence stage (stage 19), as well as the stage where the corolla is fully expanded (stage 20) and the pollen and ovules are mature (stage 20). The emergence of MMCs that had differentiated from sporogenous cells (Sp) was observed at stage 1 in *S. pimpinellifolium* and stage 7 in *S. lycopersicum*, while mature pollen release from each of the anther locules occurred at stage 10 in *S. pimpinellifolium* and at stage 20 in *S. lycopersicum* (Fig. 2 and Brukhin *et al.*, 2003). We observed that some morphological characteristics of the *S. pimpinellifolium* flower were somewhat different from those of *S. lycopersicum* during stamen development. In *S. lycopersicum*, the calyx emerged at the top of the bud at stage 12, sepals (5.0 mm) were longer than the petals (4.5 mm in length), and free irregularly shaped microspores were released from the tetrads. The corolla protruded from the calyx at stage 13, and the sepals (5.4 mm) were somewhat shorter than the petals (5.8 mm) (Brukhin *et al.*, 2003). However, in *S. pimpinellifolium* the corolla emerged from the calyx at stage 1 and, at the same time, stamen development was associated with the differentiation of sporogenous cells (Sp), making this stage equivalent to stage 7 in *S. lycopersicum*. Free, irregularly shaped microspores released from the tetrads were observed at stage 5 in *S. pimpinellifolium* (Fig. 2 stage 5 and Fig. 3 stage 5), where the sepals reached half the length of the petals, while the corolla was still surrounded by sepals in *S. lycopersicum* (Brukhin *et al.*, 2003). Initiation of the tapetum layer degeneration occurred at stage 13 in *S. lycopersicum*, a stage where the corolla first emerged from the calyx; however the cells of the *S. pimpinellifolium* tapetum layer began to degenerate at stage 4 (Fig. 2 stage 4), at which time the corolla was already projecting beyond the calyx (Fig. 1C). These results suggest differences between *S. pimpinellifolium* and *S. lycopersicum* in both morphological features and microspore developmental processes during stamen developmental, and that data derived from studies of *S. lycopersicum* are not fully analogous to *S. pimpinellifolium* or other wild tomato species. Consequently, the results derived from *S. pimpinellifolium* might be more applicable for research into the morphology, histology and cytology at particular stamen developmental stages, and will provide a useful platform for dissecting the molecular mechanisms of developmental and reproductive biology in tomato and its wild relatives.

Conclusions

Based on distinct flower developmental stages, *S. pimpinellifolium* anther/stamen development was divided into ten stages and a correlation between the morphological characteristics and cytological landmark events at each developmental stage was established.

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References


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Sato et al., 2011. missing in ref. list.


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