Disorganization of F-actin cytoskeleton precedes vacuolar disruption in pollen tubes during the in vivo self-incompatibility response in Nicotiana alata

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• Background and Aims The integrity of actin filaments (F-actin) is essential for pollen-tube growth. In S-RNase-based self-incompatibility (SI), incompatible pollen tubes are inhibited in the style. Consequently, research efforts have focused on the alterations of pollen F-actin cytoskeleton during the SI response. However, so far, these studies were carried out in in vitro-grown pollen tubes. This study aimed to assess the timing of in vivo changes of pollen F-actin cytoskeleton taking place after compatible and incompatible pollinations in Nicotiana alata. To our knowledge, this is the first report of the in vivo F-actin alterations occurring during pollen rejection in the S-RNase-based SI system.

• Methods The F-actin cytoskeleton and the vacuolar endomembrane system were fluorescently labelled in compatibly and incompatibly pollinated pistils at different times after pollination. The alterations induced by the SI reaction in pollen tubes were visualized by confocal laser scanning microscopy.

• Key Results Early after pollination, about 70 % of both compatible and incompatible pollen tubes showed an organized pattern of F-actin cables along the main axis of the cell. While in compatible pollinations this percentage was unchanged until pollen tubes reached the ovary, pollen tubes of incompatible pollinations underwent gradual and progressive F-actin disorganization. Colocalization of the F-actin cytoskeleton and the vacuolar endomembrane system, where S-RNases are compartmentalized, revealed that by day 6 after incompatible pollination, when the pollen-tube growth was already arrested, about 80 % of pollen tubes showed disrupted F-actin but a similar percentage had intact vacuolar compartments.

• Conclusions The results indicate that during the SI response in Nicotiana, disruption of the F-actin cytoskeleton precedes vacuolar membrane breakdown. Thus, incompatible pollen tubes undergo a sequential disorganization process of major subcellular structures. Results also suggest that the large pool of S-RNases released from vacuoles acts late in pollen rejection, after significant subcellular changes in incompatible pollen tubes.

Key words: Confocal microscopy, F-actin cytoskeleton, Nicotiana alata, pollen tube, self-incompatibility, S-RNase, vacuolar system.

INTRODUCTION

The cytoskeleton of actin is an essential component of the polar cell growth machinery of pollen tubes (Cheung and Wu, 2008). As in few other examples of tip growth in nature, such as root hairs in plants, hyphae in fungi and neurites in animals, actin filaments (F-actin) in normally growing pollen tubes exhibit a characteristic organization. Long cables of polymerized F-actin are arranged in parallel to the major axis of the cell. These bundles of F-actin extend in the cytoplasm from the subapical region along the shank of the pollen tube. In the subapical region, shorter and randomly disposed F-actin fragments form the ‘collar zone’, a dense mesh of actin immediately behind the apex (Geitmann et al., 2000; Vidali et al., 2001). The apical zone possesses individual and fine filaments of actin, difficult to observe and mostly under the limit of resolution for light microscopy (Staiger et al., 2010). This highly structured organization of F-actin provides the appropriate frame for vesicle and organelle motility, including the transport to and from the apex of secretory vesicles, necessary to sustain the tip growth (Staiger et al., 1994; Cai and Cresti, 2009). Given that pollen tubes germinate and grow easily in culture medium, most studies of F-actin organization and dynamics have been made in in vitro systems. However, some in vivo studies performed in pollen tubes growing through the style have confirmed the F-actin general disposition (Geitmann et al., 2000; Chen et al., 2002).

Self-incompatibility (SI) is a genetic mechanism controlled by the S-locus to prevent self-fertilization and the consequent inbreeding and fitness decay in plant populations (Rea and Nasrallah, 2008). SI has evolved several times in independent lineages of angiosperms and a variety of strategies are used by different plant families to stop germination or growth of incompatible pollen. At the molecular level, the best known SI mechanism is probably that of Papaver rhoeas which has been successfully reproduced in an in vitro system, mimicking in vivo conditions (Franklin-Tong, 2008; Poulet et al., 2011). This research has revealed the crucial role of F-actin cytoskeleton during the rejection of incompatible pollen in Papaver. Early after SI-challenge, a massive cytoplasmic influx of free Ca²⁺ at the shank of the pollen tube triggers dramatic depolymerization of F-actin cables and subsequent formation of...
punctate actin foci (Geitmann et al., 2000; Snowman et al., 2002; Poulter et al., 2010). Moreover, F-actin depolymerization induced by SI triggers a programmed cell death (PCD) process in incompatible pollen (Thomas et al., 2006; Bosch and Franklin-Tong, 2007; Wilkins et al., 2011). Thus, F-actin acts as a target and effector of SI-signalling network. Changes in F-actin organization linked to induction of PCD have also been reported in animals and yeast, in several biological processes (reviewed by Franklin-Tong and Gourlay, 2008; Smertenko and Franklin-Tong, 2011).

In the S-RNase-based SI system, studied mainly in Solanaceae, Plantaginaceae and Rosaceae (Hua et al., 2008; Chen et al., 2010; McClure et al., 2011), the role of F-actin has been considerably less studied than that of Papaveraceae. In Pyrus pyrifolia, in vitro germination and pollen-tube growth were selectively inhibited by an S-RNase of the same pollen haplotype added to the culture medium (Hiratsuka et al., 2001). S-RNase caused the disruption of tip-localized reactive oxygen species; this, in turn, induced the depolymerization of F-actin, which changed from the typical axial cables to punctate actin pattern (Liu et al., 2007; Wang et al., 2010). These changes occurred in parallel with mitochondrial alterations and nuclear DNA degradation (Wang et al., 2009) suggesting that configuration changes in F-actin would also trigger PCD during incompatible pollen rejection in Rosaceae (Wang and Zhang, 2011).

In Solanaceae, subcellular changes produced in incompatible pollen were studied early at the ultrastructural level. Morphological alterations in the rough endoplasmic reticulum, aggregation of cytoplasmic vesicles in the apex and an increase in callose deposition were reported as early manifestations of pollen rejection in Lycopersicum peruvianum (de Nettancourt et al., 1973) and Brugmansia suaveolens (Geitmann et al., 1995). More recently, it has been established that S-RNases were localized into vacuoles (Goldraij et al., 2006; Meng et al., 2009) which appear to selectively collapse after incompatible pollination, releasing the S-RNase to the cytoplasm of incompatible pollen tubes. However, no studies have been focused on the alterations of F-actin during pollen rejection in Solanaceae. On the other hand, to our knowledge, with the exception of a single experiment in Papaver (Geitmann et al., 2000), progress in understanding configurational changes in F-actin during the SI reaction has been exclusively made in in vitro-cultivated pollen rather than in in vivo-grown pollen.

In this work, we study the alterations in the F-actin cytoskeleton of pollen tubes during the in vivo SI reaction in pollinated styles of Nicotiana alata. As the incompatible pollen tubes progressed into the style, disorganization of F-actin cables into shorter fragments gradually increased, reaching almost 80 % of the pollen tubes when they were already fully arrested. However, at this point, in >80 % of pollen tubes the vacuolar system was still well-organized. This result indicates that F-actin disorganization precedes the vacuolar breakdown during the process of incompatible pollen rejection in Nicotiana. Thus, incompatible pollen tubes undergo a sequential disorganization of subcellular structures, rather than an uncontrolled cytoplasmic degeneration. Consequently, it appears that the large pool of S-RNase stored in vacuoles would act late in pollen rejection, once major subcellular changes have taken place in incompatible pollen tubes. A comparison of these results with the subcellular changes in incompatible pollen tubes of Papaver and Pyrus during the in vitro SI response is discussed.

MATERIALS AND METHODS

Plant material, conditions of growth and pollinations

Self-incompatible Nicotiana alata individuals Sc10Sc10 and S70S70, collected from a natural population previously described (Roldán et al., 2010), were self-pollinated at an immature stage of pistil development to generate the homozygous genotypes Sc10Sc10 and S70S70, used in this work. Plants were grown in a greenhouse at 28°C with a 16 h/8 h light/dark period.

Sc10Sc10 flowers were emasculated 2 d before floral maturity and pollinated 1 d after petal opening with pollen from S70S70 (compatible) or Sc10Sc10 (incompatible) genotypes.

Sample preparation for fluorescence microscopy

To determine the position of the pollen tube front in the style, pistils were collected at different times, stained with decolorized aniline blue and visualized by epifluorescence (Kho and Baer, 1968). For colocalization of F-actin with callose or vacuolar pyrophosphatase (vPPase), styles collected at different times were fixed for 45 min under slight vacuum with freshly prepared actin-stabilizing buffer (ASB) (Geitmann et al., 2000), pH 6.8, containing 250 μm m-maleimidobenzoyl N-hydroxysuccinimide ester. Subsequently, the styles were fixed for 1 h in freshly prepared ASB containing 4 % paraformaldehyde. After three washes in ASB, 15-mm segment of a style, including the growing front of pollen tubes, was selected. Style portions were embedded at −20°C in Cryoplast (Biopack, Argentina), cut into sections 10 μm thick in a cryostat (Leica microsystems, Germany) and put onto slides treated with 1-mm polylysine. Sections were incubated for 10 min with PBS containing 1 % each of Macerozyme R-10 (Yakult Honsha Co., Japan), Cellulase ‘Onozuka’ RS (Yakult Honsha Co.), and BSA. Samples were then washed three times with PBS, blocked for 30 min with PBS containing 5 % BSA and 0.1 % Triton X-100, washed again with PBS and incubated overnight with anti-callose (Biosupplies, Australia) or anti-vPPase (Goldraij et al., 2006) antibodies diluted in the blocking solution with 1 % BSA. After washing three times, goat anti-mouse Alexa 546-conjugated secondary antibody (Molecular Probes, Inc.) or donkey anti-rabbit Dy Light 549-conjugated secondary antibody (Jackson Immuno Research Laboratories, Inc.) were added for 4 h to label callose or vPPase, respectively. Following two washes with PBS and one with ASB, sections were incubated overnight with 700 nM Alexa 488-conjugated phalloidin diluted in ASB containing 5 mm DTT and 0-1 % Tween 20. Specimens were washed and mounted in Fluorsave (Calbiochem, Schwalbach, Germany). For single-labelling of F-actin, the sections were incubated directly with 500 nM Alexa 488-conjugated phalloidin in the conditions described above.
Fluorescence microscopy

Confocal laser scanning fluorescence microscopy images were recorded with Zeiss LSM 5 Pascal (Carl Zeiss, Oberkochen, Germany). To detect Alexa 488, samples were scanned by argon laser (excitation 488 nm; emission, filter BandPass of 505–530 nm). Alexa 546 and DyLight 549 were detected by using helium-neon laser (excitation 543 nm; emission, long pass 560 nm). Samples were visualized with a Plan-Apochromat ×100/1.4 NA Oil DIC objective using a ×2 digital zoom. Between 12 and 14 0.5-μm serial optical sections were acquired. The images were deconvoluted using the ‘advanced maximum likelihood estimation algorithm’ for Cell R software (Olympus Soft Imaging Solutions, Munchen, Germany), version 3.3, set with 20 iterations and an overlay sub-volume of 10 pixels. A theoretical point spread function was used.

The F-actin and the vacuolar membranes were visualized in the subapical region, including the first 50–100 μm of the pollen tube. For quantitation, F-actin and vacuolar membranes were scored as ‘organized’ or ‘disorganized’ in at least 75 pollen tubes for each replica.

RESULTS

Pollen-tube growth in compatible and incompatible pollinations

We determined the growth rate for compatible and incompatible pollen tubes in pollinated pistils of plants derived from a natural population of Nicotiana alata (Fig. 1). No differences were seen within the first 12 h after pollination, when pollen-tube tips had penetrated about 5 mm into the style. Then, in the compatible cross Sc10Sc10 × S70S70, pollen tubes grew through the style at a linear rate and reached the ovary within 72 h post-pollination. Incompatible pollen tubes from self-pollinated Sc10Sc10 plants also grew linearly on the first 3 d after pollination and subsequently continued growing at a slightly slower rate, until growth stopped below mid-style by day 5 after pollination. Since the styles were about 69 mm in length, the average growth rate was 1.0 and 0.4 mm h⁻¹ for compatible and incompatible pollen tubes, respectively.

F-actin pattern in compatible and incompatible pollinations

We checked first whether the fixation method used in this study was efficient to preserve F-actin disposition in in vivo-grown pollen tubes. Fixed sections of cross-pollinated pistils were stained with Alexa 488-phalloidin to label F-actin. The typical bundle of F-actin cables parallel to the long axis of the cell was clearly distinguished from the subapical zone to the shank (Fig. 2). A shorter filament array was seen in the ring zone, in the transition between apical and subapical regions, while the dome of the tip had fainter and disperse actin filaments. This pattern of F-actin disposition was similar to that previously reported in pollen tubes stained by different methods (Geitmann et al., 2000; Wilsen et al., 2006; Cheung et al., 2008), confirming that the fixative procedure was appropriate to preserve F-actin organization of in vivo-grown pollen tubes.

Sections of cross- and self-pollinated styles collected at different times after pollination were double-labelled with phalloidin and with an antibody against callose, to stain F-actin and to demarcate the pollen tube boundaries, respectively. Representative images of compatible and incompatible pollen tubes of similar lengths are shown (Fig. 3). A similar pattern of well-organized F-actin cables was visualized 12 h after pollination (Fig. 3A, B), in consistency with the similar growth rates of compatible and incompatible pollen tubes during early pollination. In contrast, a remarkable difference was seen at a late stage of pollination. Two days after pollination, when compatible pollen tubes were in the lower style, the F-actin organization was similar to that observed in early pollination (Fig. 3C). Conversely, 5 d after incompatible pollination, when pollen growth was already arrested (see Fig. 1), F-actin had lost its normal appearance. Instead, smaller and discontinuous F-actin fragments of variable size were detected by using helium-neon laser (excitation 543 nm; emission, long pass 560 nm). Samples were visualized with a Plan-Apochromat ×100/1.4 NA Oil DIC objective using a ×2 digital zoom. Between 12 and 14 0.5-μm serial optical sections were acquired. The images were deconvoluted using the ‘advanced maximum likelihood estimation algorithm’ for Cell R software (Olympus Soft Imaging Solutions, Munchen, Germany), version 3.3, set with 20 iterations and an overlay sub-volume of 10 pixels. A theoretical point spread function was used.

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lengths were extended along the shank of incompatible pollen tubes (Fig. 3D). Additional images of F-actin in compatible and incompatible pollen tubes are shown in Supplementary Data Fig. S1. The two F-actin configurations shown in Fig. 3 were quantified over time for compatible and incompatible pollinations (Fig. 3E). In both types of pollinations, 30% of pollen tubes exhibited disorganized F-actin 12 h post-pollination. While in compatible pollinations this percentage was kept constant throughout the trajectory of pollen tubes, the number of incompatible pollen tubes with disorganized F-actin exhibited a progressive increase, reaching 70% at the time of growth arrest and almost 80% 8 d after pollination. The pattern of F-actin disorganization shown in Fig. 3D was largely predominant at all time points after incompatible pollination. Although punctate foci of actin characteristic of in vitro-grown incompatible pollen tubes (Poulter et al., 2010; Wang et al., 2010) were occasionally observed, by day 8 after pollination they represented <13% of pollen tubes with altered F-actin organization (Supplementary Data Fig. S2).

Disruption of F-actin cables in incompatible pollen tubes precedes vacuolar compartment disorganization

In the S-RNase-based SI system, the progression of morphological changes during incompatible pollen rejection is not fully understood. After the occurrence of early subcellular alterations, such as the excess of callose deposition, the disposition of the endoplasmic reticulum in concentric circles, and
vesicle accumulation in the apex, the degeneration of the pollen tube cytoplasm has been proposed (Geitmann, 1999; de Graaf et al., 2006). On the other hand, the disorganization of vacuolar compartments, where S-RNases are confined (Goldraij et al., 2006; Meng et al., 2009), has been reported as a late event during self-pollen rejection (Goldraij et al., 2006). We assessed whether there was a simultaneous disorganization of F-actin bundles and the vacuolar system or, conversely, if disorganization of these major subcellular structures was carried out in a progressive and sequential manner. Colocalization experiments for F-actin cytoskeleton and the vacuolar system of pollen tubes were performed after incompatible pollinations. An antibody against vPPase was used to visualize the tonoplast of pollen vacuolar compartments. In the pollen subapical zone, these compartments were clearly defined and considerably diverse in size and morphology (Hicks et al., 2004; Supplementary Data Fig. S3). Representative images of different patterns of F-actin and vacuolar membranes in incompatible pollen tubes at different days after pollination are shown in Fig. 4. Both F-actin and

![Image](https://example.com/image.png)

**Fig. 4.** Disorganization of F-actin cytoskeleton precedes vacuolar breakdown in incompatible pollen tubes during the in vivo SI response. Colocalization of F-actin (green) and vPPase (red) in incompatibly pollinated pistils. The images are representative of the three patterns observed: (A) organized F-actin and vacuolar endomembrane system; (B) disorganized F-actin and organized vacuolar endomembrane system; (C) disorganized F-actin and vacuolar endomembrane system. Merged optical sections (top panels), full projections of phalloidin-stained pollen tubes (central panels) and the corresponding bright fields (bottom panels) are shown. The days after pollination are indicated below the images. Scale bar = 10 μm. (D) Quantification of morphology of F-actin and vacuolar endomembrane system. White, organized F-actin and vacuolar system; dark grey, disorganized F-actin and organized vacuolar system; light grey, disorganized F-actin and vacuolar system. Seventy-five pollen tubes were examined in each experiment. The values are the average ± s.d. for two or more independent experiments.
vacuolar compartments were predominantly well-organized 1 d after pollination (Fig. 4A). Instead, by day 3 after pollination, the majority of pollen tubes showed that F-actin bundles mostly disappeared and were replaced by much smaller fragments, while the integrity of the vacuolar system appeared to be normal (Fig. 4B). By day 8 post-pollination, the loss of integrity of F-actin in individual cells was similar to that observed 3 d after pollination. Most pollen tubes still exhibited organized vacuoles; however, there was an increase in the frequency of pollen tubes in which no signal of vacuolar compartments was detected, while the vacuoles of the adjacent transmitting tissue were normally labelled (Fig. 4C). The simultaneous quantification of F-actin organization and the integrity of the vacuolar compartment in incompatible pollen tubes is shown in Fig. 4D. From day 1 to day 3 after incompatible pollination, pollen tubes with disorganized F-actin increased from 48% to 67%, but <8% of pollen tubes showed both disorganized F-actin and vacuolar compartments by 3 d after pollination. By 8 d after pollination almost 70% of pollen tubes showed intact vacuoles but >84% of pollen tubes had disorganized F-actin. No pollen tubes with organized F-actin and disrupted vacuolar compartments were seen on any day. These results suggest strongly that F-actin disorganization occurs earlier than vacuolar breakdown during the process of incompatible pollen rejection in *Nicotiana*.

**DISCUSSION**

In this work we studied the *in vivo* subcellular changes of *Nicotiana alata* pollen tubes after compatible and incompatible pollinations, focusing on the organization of F-actin cytoskeleton and the vacuolar endomembrane system. We analysed the temporal sequence of alterations in incompatible pollen tubes establishing that the disorganization of F-actin occurs progressively over time and precedes vacuolar disruption, which takes place mainly after pollen-tube growth has been completely inhibited.

The highly organized disposition of F-actin cytoskeleton in pollen tubes is essential to support polar tip growth (Cheung and Wu, 2008). Longitudinal cables along the shank of the pollen tube provide the appropriate track for organelle movement and cytoplasmic streaming, while finer and shorter actin filaments in the subapical and apical zone are presumably involved in vesicle transport to the apex (Staiger et al., 2010). Disruption of F-actin cytoskeleton organization causes a severe reduction or inhibition of pollen-tube growth (Gibbon et al., 1999; Cheung et al., 2000; Thomas et al., 2006). Thus, F-actin disorganization in *Nicotiana* could be an early manifestation at the cellular level of a more complex process leading to pollen rejection, as was reported on *Papaver* (reviewed in Franklin-Tong, 2008). To place the F-actin disruption within the time course of SI-induced alterations, actin structures were colocalized with vPPase, a tonoplast marker of the vacuolar endomembrane system. Vacuolar disruption was characterized as a late event in the SI reaction in *Nicotiana*. This is particularly relevant, since vacuoles demarcated by vPPase store the S-RNase (Goldraij et al., 2006), the cytotoxic agent ultimately responsible for pollen rejection in the S-RNase-based SI systems (McClure et al., 1990; Huang et al., 1994). Simultaneous visualization of F-actin cytoskeleton and vacuolar membranes allowed us to establish a sequence of cellular events during pollen rejection (Fig. 4). Six days after pollination, when the incompatible pollen-tube growth was fully inhibited, about 80% of pollen tubes exhibited disorganized F-actin structures; however, the vacuolar membrane system was still intact in 80% of pollen tubes. Thus, during incompatible pollen rejection in *Nicotiana*, vacuole disruption occurs almost entirely after an extensive F-actin disorganization.
Ultrastructural studies in the S-RNase-based SI system revealed that incompatible and compatible pollen tubes cannot be distinguished early after pollination. The first morphological changes in the Brugmansia and Lycopersicum populations studied were visualized between 4 h and 8 h post-pollination and consisted of concentric structures in ER, accumulation of vesicles in the tube apex and thickened cell wall by the increase of callose deposition (de Nettancourt et al., 1973; Geitmann et al., 1995). Subsequently, a general cytoplasmic degeneration was reported to occur (Geitmann, 1999). Based on these morphological changes and the slow timing of pollen rejection, it has been suggested that a necrotic process in incompatible pollen in the S-RNase-based SI system occurs (Geitmann, 1999; Geitmann et al., 2004; de Graaf et al., 2006). Interestingly, both F-actin disorganization and tonoplast disruption have been described as typical alterations in many processes of plant PCD, including SI (for recent reviews, see Hara-Nishimura and Hatsugai, 2011; Smertenko and Franklin-Tong, 2011). F-actin depolymerization induced PCD during pollen rejection of incompatible pollen tubes in *Papaver* (Thomas et al., 2006) through a well-studied signal-cascade (Franklin-Tong, 2008) that resulted in caspase activation and a pronounced acidification of cytosol (Bosch and Franklin-Tong, 2007). This suggested that tonoplast rupture could be involved in *Papaver* SI-reaction. Even more significant, in *Pyrus*, where pollen is rejected by the S-RNase-based SI system, extensive F-actin depolymerization was induced within 30 min after the addition of self-S-RNase to *in vitro*-grown pollen tubes (Liu et al., 2007; Wang et al., 2010). Alterations in mitochondrial permeability, DNA degradation and tip-localized reactive-oxygen species disruption were also triggered in *Pyrus* pollen tubes by S-RNase added to the growth medium (Wang et al., 2009, 2010). Since all these changes took place before pollen-growth arrest, the authors suggested that they were the cause but not the consequence of pollen inhibition (Liu et al., 2007; Wang and Zhang, 2011) and proposed a cascade-signal pathway that would produce PCD in *Pyrus* pollen. In this model, RNA degradation by S-RNase could be only the beginning of the SI response (Wang and Zhang, 2011). On the other hand, the results reported here from *in vivo* experiments, show that pollen growth inhibition and F-actin disorganization occurred in parallel and preceded tonoplast disruption. Since a large pool of S-RNases is compartmentalized into pollen vacuoles (Goldraij et al., 2006; Meng et al., 2009), these results suggest that, in *Nicotiana*, massive RNA degradation would be a late event in the SI reaction, occurring after major cellular changes take place in rejected pollen tubes. It could be possible that this late and massive release of S-RNase was necessary to warrant the fulfilment and the irreversibility of the rejection process. However, we cannot rule out the existence of a different pool of S-RNases exerting some cytotoxic action in an early stage of pollen rejection. Recently, questions have been raised about possible differences between *in vivo* and *in vitro* mechanisms of pollen rejection in the S-RNase-based system (Zhang et al., 2009; Chen et al., 2010). In any case, since the nature of pollen rejection in S-RNase-based SI is complex, the exact role of S-RNase is still a matter of debate (Zhang et al., 2009). Early grafting experiments conducted by Lush and Clarke (1997) showed that growth inhibition caused by incompatible styles could be reverted in some pollen tubes, suggesting that the cytotoxic effect of S-RNase was not permanent and not exclusively related to RNA degradation. In this regard, it is noteworthy that, in several T2 ribonucleases, some biological roles, including cytotoxicity, are independent of their catalytic activities (Thompson and Parker, 2009; for a review, see Luhtala and Parker, 2010). Interestingly, a fungal T2-RNase, termed ACT1BIND, exerts its cytotoxic activity by binding actin and disrupting the actin network in colon cancer cells (Roiz et al., 2006). It will be crucial to know if additional roles to RNA degradation of incompatible pollen can be attributed to S-RNases.

Importantly, our results fit well the two models that currently explain pollen rejection in the S-RNase-based SI system. The S-RNase degradation and S-RNase compartmentalization models can be conciliated, since both assume that recognition and cytotoxicity take place in pollen cytoplasm, where pollen recognition factors – the SLF proteins – are localized (Wang and Xue, 2005; Meng et al., 2010). A minor portion of S-RNases can gain access to the pollen cytoplasm by retrograde transport for recognizing the SLF products (Goldraij et al., 2006). In incompatible pollinations, self-S-RNases and/or other still unknown factors could initiate a cytotoxic effect targeting gradually the organization of the F-actin cytoskeleton (Wang and Zhang, 2011). This in turn might trigger a cascade of signals, affecting subsequently vacuolar membrane integrity and releasing the large pool of compartmentalized S-RNases in the final steps of pollen rejection. Conversely, in compatible pollinations, degradation of ubiquitinated non-self-S-RNases by proteosome machinery (Hua et al., 2008) would preserve the integrity of F-actin and vacuolar compartments, allowing the pollen tube to progress to the ovary. It is expected that non-S-specific pistil factors also contribute to regulate both compatible and incompatible scenarios, although their precise role is still unknown.

In conclusion, during the SI response in *Nicotiana*, significant subcellular changes took place in incompatible pollen tube through a sequence of time-ordered events. The normal bundles of pollen F-actin cytoskeleton were gradually disrupted, in parallel with a reduced growth rate. The breakdown of the vacuolar endomembrane system occurred subsequently, only after an extensive F-actin disorganization. These results indicated that during rejection of incompatible pollen there was a progressive rather than a rapid and uncontrolled collapse of cellular structures. This also suggested that the catalytic activity of S-RNases, which was shown to be essential for the SI response (Huang et al., 1994), would manifest itself in a late stage of pollen rejection, after major cellular dismantling of incompatible pollen tubes.

**SUPPLEMENTARY DATA**

Supplementary data are available online at www.aob.oxfordjournals.org and consist of the following. Figure S1: changes in the F-actin cytoskeleton of pollen tubes during the *in vivo* SI response. Figure S2: punctate foci of actin in incompatible pollen tubes. Figure S3: morphological diversity of pollen tube vacuoles.
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LITERATURE CITED


