Proteomic analysis of *Brassica* stigmatic proteins following the self-incompatibility reaction reveals a role for microtubule dynamics during pollen responses

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Running Title: Microtubule depolymerization promotes pollen acceptance

Abbreviations

SI: Self-incompatibility; MT: Microtubules; SRK: S-locus receptor kinase; SCR: S-locus cysteine rich protein; ARC1: ARM-repeat containing protein
Summary

Mate selection and maintenance of genetic diversity is crucial to successful reproduction and species survival. Plants utilize self-incompatibility (SI) system as a genetic barrier to prevent self pollen from developing on the pistil, leading to hybrid vigor and diversity. In Brassica (canola, kale and broccoli), an allele-specific interaction between the pollen SCR/SP11 (S-locus cysteine rich protein/S locus protein 11) and the pistil S Receptor Kinase (SRK), results in the activation of SRK which recruits the Arm Repeat Containing 1 (ARC1) E3 ligase to the proteasome. The targets of ARC1 are proposed to be compatibility factors, which when targeted for degradation by ARC1 results in pollen rejection. Despite the fact that protein degradation is predicted to be important for successful self-pollen rejection, the identity of the various proteins whose abundance is altered by the SI pathway has remained unknown. To identify potential candidate proteins regulated by the SI response, we have used the two-dimensional difference gel electrophoresis (2-D DIGE) analysis, coupled with MALDI-TOF/MS. We identified 56 differential protein spots with 19 unique candidate proteins whose abundance is down-regulated following SI pollinations. The identified differentials are predicted to function in various pathways including biosynthetic pathways, signaling, cytoskeletal organization and exocytosis. From the 19 unique proteins identified, we investigated the role of tubulin and the microtubule network during both SI and compatible pollen responses. Moderate changes in the microtubule network were observed with SI pollinations; however, a more distinct localized break-down of the microtubule network was observed during compatible pollinations, that is likely mediated by EXO70A1, leading to the successful compatible pollination.
Introduction

Recognition of suitable pollen is a key aspect of reproductive success of angiosperms. Despite having little control over the type of pollen that lands on the stigma of the pistil, many plant species have developed elaborate recognition systems to allow the growth of only suitable pollen grains and to reject genetically similar (self-incompatible) and incompatible mates in order to increase the probability of successful reproduction and survival. Adherence of compatible pollen grains to the stigmatic papillae results in early signaling events leading to hydraulic connectivity with the stigma, which facilitates movement of water into the pollen grains. Following hydration, pollen grains germinate and produce tubes that penetrate the stigmatic cell walls and traverse the pistil down to the ovules where fertilization takes place (1, 2). In species of Brassicaceae (canola, broccoli, cabbage, Arabidopsis), which are characterized by dry stigmas, this recognition occurs at the earliest stages of pollen adhesion and hydration. Only compatible pollen is capable of inducing the stigma to release its resources such as water and other factors necessary for pollen growth, while incompatible pollen is blocked either prior to hydration or during attempts to penetrate the stigmatic barrier (3). Although multiple genetic screens have identified various compatible pollen factors necessary for the initial recognition mechanisms, the molecules or mechanisms behind delivery of stigma factors necessary to support pollen growth remain largely unknown (3,4). The lipids of the stigmatic cuticular layer are proposed to interact with the lipids from pollen coat to form a hydraulic conduit for passage of water from the stigma to pollen. For example, Arabidopsis fiddlehead mutants, which are altered in their cuticular properties, support pollen hydration on non-stigmatic tissues (5,6). Removal of the pellicle, the proteinaceous outermost layer of the papillary cell that functions as the initial site of the pollen-pistil interaction, also results in lack of pollen tube entry into the stigma (7). Two Brassica
stigma proteins, S-locus related 1 (SLR1) and S-locus glycoprotein, are implicated in pollen adhesion (8). Both of these bind to specific pollen coat proteins and are proposed to mediate pollen adhesion.

In contrast to the limited knowledge about the signaling mechanism behind compatible interactions, extensive studies have identified the stigmatic factors that regulate the self-incompatibility response (SI) in Brassica. Recognition of self-pollen induces a highly ordered signaling mechanism that rejects the self-pollen (i.e. blocks allocation of pistil resources for pollen hydration/germination). The SI response, which is initiated following attachment of the self-pollen to the stigma, involves binding of a secreted pollen ligand SCR/SP11 (9-10) to the S-Receptor Kinase (SRK) localized to the plasma membrane of the stigmatic cells. This binding event activates SRK, which, along with another cytoplasmic receptor kinase, M-locus protein kinase (MLPK) (11), can interact with and phosphorylate the E3 ubiquitin ligase, ARC1 (12), to promote rejection of ‘self-pollen’. ARC1 acts as a positive regulator of SI as activated SRK can target ARC1 to the proteasome (protein degradation machinery) (13,14) and suppression of ARC1 in self-incompatible Brassica napus W1 lines leads to breakdown of SI response (13). In addition to the genetic evidence, treatment of W1 stigmas with proteasomal inhibitors resulted in breakdown of SI response leading to pollen germination and growth, indicating the requirement of an intact ubiquitin-mediated, proteasomal degradation pathway in order to confer SI response (14). The present hypothesis is that, following initiation of SI, ARC1 rapidly targets specific substrate proteins for proteasome-mediated degradation. These substrates targeted for destruction are likely compatibility factors, and their degradation results in pollen rejection (3). These compatibility factors potentially play a role in promoting hydration of compatible pollen grains and/or allow loosening of the cell wall for the compatible pollen tubes to penetrate. Recently,
EXO70A1, a member of the exocyst complex involved in targeted vesicle trafficking was identified as one of the downstream targets of ARC1 (15). Suppression of EXO70A1 in both compatible Brassica (‘Westar’ canola line) and Arabidopsis resulted in pollen rejection at very early stages of hydration and germination. Consistent with this, overexpression of EXO70A1 in self-incompatible W1 line partially restored compatibility indicating the role of EXO70A1 as a compatibility factor and sufficiency of EXO70A1 to breakdown SI response (15). But the exact mechanism through which EXO70A1 imparts its function is still unknown. While EXO70A1 functions as one of the compatibility factors, whether there are other early signaling proteins involved in compatibility response remains unanswered. Given the fact that proteasome blockers could restore compatibility in an SI background (14), it may be likely that many compatibility proteins are targeted for degradation following SI response. Identification of these factors could provide us with significant insights into both how compatibility response operates and how SI abrogates this process by targeting these proteins for degradation. Thus, by quantifying changes in protein abundance through analysis of the proteome prior to, and after self-incompatible pollination, we can identify the subset of proteins whose abundance is modified following SI response. These proteins would be predicted to include critical compatibility factors that are targets of SI.

In order to identify changes in stigma protein levels following SI, we utilized a two-dimensional difference gel electrophoresis (2-D DIGE technology). The 2-D DIGE approach has significant advantages over conventional 2-D electrophoresis where the analysis is biased due to gel to gel variations that hinder precise quantification (16). In 2-D DIGE, respective protein samples are labeled with different fluorescent dyes such as Cy2, Cy3 or Cy5 followed by mixing of these samples and electrophoresis through conventional 2-D. The scans for the various channels will
allow precise quantitation of the relative abundance of the perfectly matched spots. Subsequently, these spots can be isolated and protein identified through mass spectrometry. Previously 2D-DIGE technology has been effectively used to identify stress and hormone regulated proteins from various plant species including Brassica and Arabidopsis (16-18). Through using 2D-DIGE and mass spectrometry to identify protein changes following SI response, we have identified 19 unique SI-regulated proteins. Interestingly, all of these proteins exhibited reduced levels following the SI response suggesting that some of these proteins could be rapidly degraded following activation of SI. From the identified list of proteins, one candidate, tubulin, was chosen for further testing, and the role of microtubule (MT) network was explored during SI and compatible pollinations.

**Experimental Procedures**

*Plant growth conditions and treatment.* Brassica napus canola cultivars spp., compatible Westar and self-incompatible W1, were germinated on soil and maintained at standard growth conditions (22°C, 16 h/8 h day/night cycle). For proteomic analysis, W1 flowers were self-pollinated with W1 pollen for different times and pistils were harvested and frozen at -80°C until protein extraction. Arabidopsis Col-0 and mor1-1 were sterilized with 30% bleach for 10 min and germinated on half strength Murashige and Skoog basal salt mixture pH 5.7, grown for a week, transplanted on to soil and maintained at standard growth conditions. For restrictive temperature treatments (29°C), Col-0 and mor1-1 flowers were emasculated and moved to chambers pre-conditioned at 29°C followed by pollination after 12 hours.
**Extraction of Total Proteins for 2-D DIGE.** Total proteins were extracted using the phenol-methanol method (19). For each biological replicate, approximately 100 stigmas were ground in liquid nitrogen with a pestle and mortar, followed by addition of 1mL of protein extraction buffer (0.5M Tris (pH7.5), 10mM EDTA, 2% β-mercaptoethanol and 1x protease inhibitor) and centrifugation at 15,000g for 20 min at 4 °C and the supernatant was collected in 2 mL eppendorf tubes. The supernatant was mixed with equal volume of ice-cold phenol (Tris-buffered, pH 7.5–7.9) and spun down at 15,000g for 20min at 4°C to separate phenol. Following removal of the aqueous phase, the intact interface along with the phenol phase was extracted twice with the extraction buffer. To the interface and phenol phase, 5 volumes of cold 0.1M ammonium acetate in methanol was added and incubated at -20°C overnight, followed by centrifugation at 15,000g. The supernatant was discarded and the pellet was washed twice with 100µL of 80% acetone, allowed to dry and re-suspended in lysis buffer (7 M urea, 2 M thiourea, 4% CHAPS), and protein concentration determined using the Bio-Rad protein assay mixture using BSA as a standard.

**Protein CyDye Labeling**—Proteins were labeled with CyDye DIGE Fluor minimal dye (GE Healthcare) according to the manufacturer’s instructions with modifications. Briefly, 10 µl (around 3µg/µl) of proteins were mixed with 0.2µl 1.5 M Tris-HCl buffer (pH 8.8) to adjust the pH of the protein solution to around 8.8. To the protein solution,100 pmol CyDye (1 µl of 0.1mM CyDye stock solution) was added and incubated on ice in the dark for at least 2 h. The reaction was stopped by addition of 0.5 µl of 10 mM lysine and incubated on ice for 10 min.

**Two-dimensional Gel Electrophoresis**—Pairs of Cy3- and Cy5-labeled protein samples were mixed together. The DIGE sample buffer (7 M urea, 2 M thiourea, 4% CHAPS, 20 mM DTT, and 1% IPG buffer (GE Healthcare)) was added to bring the volume to 450 µl, and the samples
were then applied to 24-cm Immobiline Drystrips (GE Healthcare). IEF was carried out on an Ettan IPGphor II (GE Healthcare) at 20 °C with maximum 50 µA/strip and the following setting: rehydration for 2 h and then 50 V for 10 h, 500 and 1000 V each for 1 h, gradient increase to 8000 V in 3 h, and remaining at 8000 V until reaching the desired total V-h (80,000 for pH 4 –7 and 100,000 for pH 3.0-5.6/5.3-6.5/6.3-7.5/6-11 IPG strips). After IEF, IPG strips were equilibrated in equilibration buffer (6 M urea, 30% (w/v) glycerol, 2% SDS, 50 mM Tris-HCl, pH 8.8) first with 0.5% DTT and then with 2% iodoacetamide each for 15 min. The strips were then transferred to 10% SDS-PAGE gels for the second dimension electrophoresis using the Ettan Dalt Twelve gel system (GE Healthcare). SDS-PAGE was run at 50V for 2 h and then 120V for 12–16 h or until the bromophenol blue dye front reached the gel end. 

**Image Scanning**—The gels were scanned using a Typhoon 8600 scanner (GE Healthcare). Cy3-labeled sample was scanned using a 532 nm laser and a 580/30-nm band pass emission filter, whereas Cy5-labeled sample was scanned with a 633 nm laser and 670/30-nm band pass emission filter. The photomultiplier tube was set to ensure that the maximum pixel intensity was between 60,000 and 90,000 pixels. Post-electrophoretic staining with deep purple (GE Healthcare) fluorescent dyes was performed essentially according to the manufacturer’s instructions, and subsequent scanning was performed using the same setting as that used for Cy3-labeled sample.

**2-D DIGE Data Analysis Using the DeCyder Software**—The DIGE images were analyzed using DeCyder version 6.5 software as described in the user manual (GE Healthcare). Briefly the differential in-gel analysis module was used for spot detection, spot volume quantification, and volume ratio normalization of different samples in the same gel. Then the biological variation analysis module was used to match protein spots among different gels and to identify protein
spots that exhibited significant difference. Manual editing was performed in the biological variation analysis module to ensure that spots were correctly matched between different gels and were not contaminated with artifacts, such as streaks or dust. Differentially expressed spots were selected based on the $p$ value of $t$ test ($p < 0.05$); presence on at least three of the five replicates, including presence on the reverse labeled one; and peak height of over 100 (absolute value prior to subtracting background for normalization) on at least three gels.

**Spot Picking**—Approximately 30 µg of protein were labeled with 50 pmol of Cy5 dye and mixed with 500 µg of unlabelled protein (250 µg each from pollinated and unpollinated samples) and separated with 2-DE. SDS-PAGE gels were stained with deep purple dye (GE Healthcare) or SYPRO Ruby dye (Bio-Rad). After scanning the gels with Typhoon 8600, spots of interests were selected with DeCyder software and picked by an Ettan spot picker (GE Healthcare) and stored in 1% (v/v) acetic acid until further processing.

**In-gel Tryptic Digestion, Mass Spectrometry, and Database searching and compilation**—Samples were received as excised spots from a 2D polyacrylamide gel and were reduced, alkylated, tryptically digested and extracted using the standard protocol outlined at [http://www.biol.yorku.ca/cm/proteomics/home.html](http://www.biol.yorku.ca/cm/proteomics/home.html). Extracted peptides were desalted and concentrated using Millipore C18 zip-tips®, then spotted in 1.5µl MALDI matrix (10mg/mL α-cyano-4-hydroxycinnamic acid (CHCA), in 60% acetonitrile, 0.3% trifluoroacetic acid) onto a 100-well MALDI target plate. The MALDI target plate was then loaded in to a QSTAR XL QqTOF mass spectrometer (Applied Biosystems/MDS Sciex, Foster City, CA) and MALDI MS spectra were collected for each sample over 20 – 50 seconds, with laser energy of 12uJ and frequency of 20Hz. Internal trypsin autolysis peptides were used to calibrate the instrument for each sample. MS spectra were then individually examined by hand and up to 17 monoisotopic
peaks were selected from each for MS/MS analysis, which was carried out over varied lengths of
time using the same laser settings and collision energy settings between 50 and 100 (arbitrary
units).

The mascot.dll data processing script from the Analyst QS 1.1 software package (Applied
Biosystems/MDS Sciex, Foster City, CA) was used to generate and concatenate peak lists for
each sample. Peak lists were searched against the entire NCBI non-redundant database (version
NCBInr_20100721 (11480388 sequences; 3917815449 residues)) using Mascot software version
2.3.02 (Matrix Science, Boston, MA). Peptide and MS/MS error tolerances were set to 100 ppm
and 0.2 Da respectively. Tryptic peptides having a charge of 1+ and up to 2 missed cleavages
were considered. Carbamidomethylation of cysteine and oxidation of methionine were set as
fixed and variable modifications respectively. All protein scores exceeding Mascot’s 95%
significance threshold (score > 56) were accepted (Supplemental data, S5). Identifications based
on single peptide identifications therefore required an individual ion score > 56 (Supplemental
data, S6).

Because the *B. napus* proteome is not exhaustively represented in the NCBI database, some
tryptic peptides interrogated by MS/MS will yield no protein match regardless of their quality.
In other cases, a protein from a source organism other than *B. napus* is identified because an
unannotated protein from *B. napus* shares some identical amino acid sequence and tryptic
peptides with an annotated protein from a close genetic relative (i.e. *Arabidopsis thaliana*). The
assumption here is that the protein being interrogated is similar in sequence, and therefore
similar in function to the identified protein. If the top MASCOT hit was not from Arabidopsis,
the identified protein was then subjected to a BLAST analysis against the Arabidopsis protein
database to find the closest homolog, thereby generating a homogenous list of Arabidopsis proteins. To help organize the list of proteins identified through MS/MS analysis, all proteins were then categorized according to GO ontology; ‘biological process’, ‘molecular function’, and ‘cellular compartment’ (The Gene Ontology Consortium, 2000). To assess whether these classifications are over-represented among Arabidopsis genes as a whole, all genes identified through MS/MS analysis were submitted to enrichment analysis using the DAVID Functional Annotation tool (20).

*Treatment of of Brassica Stigmas with MT manipulating Drugs.* Taxol and oryzalin were used as MT manipulating drugs. The stigmas of self-incompatible W1, stage 12 flowers from *Brassica* were used for the study. These flowers actively reject genetically similar W1 pollen (self-pollen) while they allow genetically dissimilar compatible pollen from Westar cultivar to develop on them. Thus the same female can be used to analyze both self-incompatible and compatible traits. The pedicel (flower stalk) of W1 flowers were immersed for 3 hours with either the drugs at 1µM and 10µM concentration or with DMS0 (1:1000 dilution), followed by pollination for four hours with self-incompatible W1 or compatible Westar pollen. The stigmas were then fixed in 3:1 ethanol and glacial acetic acid, followed by aniline blue assay.

*Sample Preparation for Anti-tubulin Staining.* Whole mount immunostaining was performed as described earlier (21). *In-planta* pollination was done on self-incompatible *Brassica* W1 stigmas with either compatible Westar or incompatible W1 pollen for 30 minutes (n=10). The unpollinated and pollinated stigmas were fixed by infiltration with 4% paraformaldehyde for 5 min followed by incubation at RT (room temperature) for 1 hour. After fixation, stigmas were washed three times with 1 X MTSB (MT stabilizing buffer; 100mM PIPES, 2mM EGTA, 1mM
MgSO₄) and subjected to freeze fraction. The stigmas were then permeabilized overnight in 1 X PBS with DTT and 0.005% Tween 20. The next morning, blocking was performed with 1% BSA for 30 minutes, followed by staining with 1:150 dilution of anti-alpha tubulin antibodies (Sigma). After two hours of incubation with the primary antibodies, the samples were incubated with the secondary antibodies conjugated to Alexa Fluor 594 (Invitrogen) (1:150 in 1X PBS) for 2 hours. The stigmas were then mounted in 50% glycerol and visualized under the Perkin Elmer UltraVIEW VoX spinning disc confocal microscope. The samples were imaged using a 63X objective lens and Z-stacks collected were subjected to restorative deconvolution to improve signal to noise ratio and visualized following extend focus view of a volume.

**Genetic analysis using mor1-1 mutant.** A temperature sensitive *Arabidopsis* mutant of MOR1 (microtubule organization 1) gene was used to study the impact of MT depolymerization on pollen acceptance. At restrictive temperature of 29°C the microtubule in *mor1-1* becomes short, sparse and misaligned (22). *Arabidopsis mor1-1* and Col-0 plants were grown under the permissive (22°C) growth conditions. Stage 11/12 flowers from Col-0 and *mor1-1* were emasculated and maintained at either at permissive temperature 22°C or restrictive temperature 29°C for 12 hours. The next morning, these stigmas (permissive and restrictive) were pollinated using wild type Col-0 pollen collected from plants maintained at 22°C. Pollination was allowed to continue for 15, 30 and 60 minutes respectively, followed by aniline blue assay to analyze pollen attachment and germination.

**Transient expression of RFP-Exo70A in GFP-tubulin BY2 Cells.** Biolistic-mediated transformation of log-phase GFP-TUA3 BY2 suspension cultured cells was performed using RFP-EXO70A1 and RFP constructs. Transformations were performed with a total of 10 µg of purified plasmid DNA. Following transformations, BY2 cells were left in the dark and then
harvested at 20 h post bombardment and visualized directly using a Leica DMR epifluorescence microscope.

Aniline Blue Assay. Aniline blue assays were performed as described previously (12). After pollination, Brassica and Arabidopsis pistils were fixed in 3:1 ethanol, glacial acetic acid for 30 minutes, followed by incubation in 1N NaOH at 60°C for 1 hour. After 1 hour, the pistils were washed with distilled water and stained with basic aniline blue (0.1 M K₃PO₄ 0.1% aniline blue). The stained samples were mounted in 50% glycerol to visualize pollen germination and pollen tubes under Leica DMR epifluorescence microscope.

Results

Identification of differentially expressed proteins following self-incompatible pollinations.

In Brassica, following incompatible pollination, increased ubiquitination has been reported to occur and the proteasome’s proteolytic function was shown to be essential for manifestation of successful SI response (14). In order to re-confirm the increased ubiquitination of proteins following the SI response, extracted proteins from unpollinated and SI pollinated stigmas were resolved through either SDS-PAGE or 2D gel electrophoresis followed by Western blotting using anti-UBQ antibodies. The blots revealed basal level of protein ubiquitination in the unpollinated stigma sample and an increased ubiquitination at 30min following SI pollination, consistent with previous observations (Fig 1a, S1a, 14). We hypothesized that the increasing ubiquitinated pools of potential SI targets will likely be degraded at 60min and thus can be identified as differential spots on a 2-dimensional gel. To test this, we adopted the 2-D DIGE
technique and resolved differentially expressed proteins from unpollinated stigmas and SI pollinated stigmas, followed by scanning and BVA analysis. Five biological replicates of stigmas harvested from different batches of W1 plants were subjected to DIGE analysis. Initially these samples were processed using various narrow range IPG strips (pH 3.0-5.6/5.3-6.5/6.3-7.5/4-7/6-11) to provide highest resolution. We found that most of the protein spots identified from narrow range 2-DE can also be seen in 2-DE gel using pH 4-7 IPG strips. The five biological replicates were then processed using pH 4-7 strips (Fig 1b). Interestingly, following DeCyder and BVA analyses (p value of t test (p < 0.05) and presence on at least three of the five replicates, including presence on the reverse labeled gel), we did not observe any proteins that were reproducibly and significantly up-regulated, while a total of 56 proteins were consistently and significantly down-regulated displaying 1.24 to 2.13 fold change in protein levels (Fig 1, Table 1).

These 56 spots were then isolated with robotic spot picker using preparative gels which were stained by deep purple following electrophoresis. These spots were subjected to MALDI/MS analysis that identified proteins from 28 spots with high confidence, of which 19 were unique proteins (Table 1, Supplement S5, S6). These 19 proteins can be classified into several functional categories based on GO ontology annotation (Fig 2). The three most frequently identified biological processes were various metabolic processes, response to stimulus and response to stress at 45%, 14%, and 13% respectively (Fig 2). The three most frequently identified molecular functions were catalytic activity, nucleotide and nucleoside binding, and hydrolase and lyase activity at 23%, 21%, and 17% respectively (Fig 2). To assess whether these classifications are over-represented among Arabidopsis genes as a whole, all genes identified through MS/MS analysis were submitted to enrichment analysis using the DAVID Functional
Annotation tool (Table S1) (20). As observed in Table S1, the identified candidates were strongly enriched relative to the gene content in the Arabidopsis genome.

Of the 19 unique proteins identified, many of the proteins were predicted to be in metabolic pathways. This may be a reflection of the SI system in the stigma reducing metabolic activities to stop the self-pollen (as the pollen requires resources from the stigma to develop). Another candidate is the S-locus related-1 (SLR1) protein which has previously been found to be involved in the adhesion of compatible pollen grains (23); thus, the down-regulation of SLR1 could be related to an SI response to reduce the adhesion of the SI pollen. Annexin is an interesting candidate as annexins have been implicated in Golgi-mediated secretion (24). Vesicle-mediated secretion is proposed as a mechanism to provide resources to the pollen during compatible pollinations, and we have previously uncovered evidence for this process to be blocked in SI pollinations (15). Both actin filaments and MTs make up the plant cytoskeleton and are required for various developmental processes including, cell division, intracellular trafficking and controlling cell shape. Previous studies have established a functional role for actin polymerization during both compatible and incompatible interactions in *Brassica* (21). The reduced abundance of actin in this analysis is consistent with a previous study demonstrating the depolymerization of actin in the stigmatic papilla following SI pollination (21). Actin-based vesicle targeting is a common theme in eukaryotic systems, and it has been previously shown that blocking actin polymerization inside the stigmatic papilla could abrogate compatible pollen acceptance (21). We confirmed that both preventing actin polymerization through cytochalasin D treatment and blocking vesicle trafficking with brefeldin A treatment in the stigma resulted in reduced compatible pollen attachment, mimicking the SI response (Fig. S2). Thus, the reduced abundance of both annexin and actin would be consistent with the SI response inhibiting vesicle
delivery to the pollen attachment site. Finally, alpha 2-4 tubulin, a subunit of the MT network, was another interesting candidate that was identified from this study and was selected for further analyses.

In plants, apart from their function during cell division in the nucleus, there is an extensive network of acentrosomal microtubule arrays in the plant cell cortex that is required for controlling cell expansion, cell morphogenesis, and is proposed to control anisotropic expansion in plant cells. In the Arabidopsis genome, the alpha tubulins are represented by at least six genes that code for four isoforms with TUA2 and TUA4 coding for the same isoform (alpha 2-4 tubulin) (25, 26). Mutation in \textit{TUA4} has been shown to result in left handed helical growth of Arabidopsis hypocotyls (27, 28). Although both actin filaments and MTs make up the cytoskeletal network, information is lacking on the precise role of MT network during pollen responses. Using a multi-faceted approach of pharmacological, genetic and cell biological techniques, we were able to further elucidate the significance of altered alpha 2-4 tubulin levels and the microtubule network following pollen responses.

\textit{Localized de-stabilization of MT network following compatible pollinations in Brassica}

The reduced abundance of alpha 2-4 tubulin in this DIGE analysis indicated that MT dynamics could be altered following the induction of an SI reaction. To visualize \textit{in vivo} changes in MT dynamics following both SI and compatible pollination, W1 stigmas were pollinated with either W1 pollen (SI) or with Westar pollen (compatible) and subjected to whole mount immuno-staining procedure using anti-tubulin antibodies and observed through a spinning disc confocal microscope. When the extended focus from the Z-stacks obtained from unpollinated W1
stigmatic papilla was examined, an intricate dense cortical MT network could be observed throughout the cell cortex (Fig. 3a, left panel). When samples were examined 30 min following SI pollinations, this intricate cortical network exhibited changes, where the dense network became re-organized into longer bundles, although no MT shortening or depolymerization could be observed (Fig. 3a, right panel). When stigmatic papillae were examined following compatible pollinations, consistently, the dense MT network could not be visualized. In particular, close to the pollen attachment site, the dense cortical network of MTs was largely eliminated or reduced to shorter segments, indicating a depolymerized state of MT (Fig. 3a middle panel, 3b). These observations suggest that while SI pollination resulted in moderate changes in MT organization (resulting in tubulin being identified in the DIGE analysis); compatible pollination appeared to trigger a more dramatic localized depolymerization of MTs.

Pharmacological modulators of MT dynamics reveal a role for MTs during compatible pollinations

In order to investigate the role of MTs during pollen-pistil interactions, we used pharmacological modifiers of MT stability and tested the ability of these drugs to interfere with both self-incompatible and compatible responses. For both interactions, the W1 pistil was used as the female parent, and following drug treatment, was either pollinated with compatible Westar or self-incompatible W1 pollen. The pistils were treated with taxol, a microtubule stabilizing drug, or oryzalin, a microtubule depolymerizing agent. Interestingly, neither taxol nor oryzalin affected the self-incompatible phenotype at the level of either pollen attachment or germination (Fig 4a). In contrast, when the effect of these inhibitors on compatible pollinations was
analyzed, treatment with taxol significantly reduced pollen attachment and pollen germination, while oryzalin treatment had the opposite effect of promoting pollen attachment and germination (Fig 4a,b,c). Overall, these observations indicated that MT depolymerization in the stigmatic papilla is a requirement for compatible pollen acceptance, which is consistent with the observations in Fig 3. That is, pre-treatment with oryzalin would increase the degree of MT depolymerization leading to increased compatible pollen acceptance; while taxol would prevent MT depolymerization by MT stabilization and decrease compatible pollen acceptance. In the case of SI pollinations, if the observations in Fig 3a of longer MT bundles reflect a stabilization of MTs, then taxol’s effect would not be detected, as the pollen are already being rejected by the endogenous SI pathway. However, this is unlikely to be a primary response in the rejection of SI pollen as MT depolymerization by oryzalin also had no effect, indicating that the SI outcome is unaffected by alteration in MT dynamics.

**mor1-1 mutants maintained at the restrictive temperature exhibit rapid pollen acceptance**

To further evaluate the role of MTs during compatible pollinations, we used a genetic approach with the Arabidopsis *mor1-1* temperature sensitive mutant. MOR1 (At2g35630) is highly expressed in Arabidopsis stigmas as observed from public microarray databases (29,30) and thus, *mor1-1* mutant plants serve as a useful genetic tool to understand role of MT organization during pollen responses. In *mor1-1*, the cortical MTs become shortened, mimicking a de-polymerized state after exposing the plants to the restrictive temperature (29°C) for shorter periods (2 hours) of time (22). Stage 12 *mor1-1* flowers were emasculated in-planta and maintained either at permissive (22°C) or restrictive (29°C) temperatures overnight and then were pollinated with pollen from wild-type Col-0 flowers maintained at permissive conditions. When the pollinated
pistils were left for 24h at the restrictive temperature, no differences could be observed for the germination of pollen grains/pollen tube growth, in comparison to wild-type Col-0 pistils (Fig S3). Based on our immune-staining and pharmacological evidence, if MT breakdown in the stigma is necessary for compatible pollen interactions, then in the mor1-1 mutant (which exhibits constitutive MT depolymerization at the restrictive temperature) both pollen germination and growth will be expected to occur more rapidly than the controls maintained at permissive conditions. In order to test this, we performed a shorter temporal pollination profile (15min - 1hr) and examined changes in pollen responses using the aniline blue staining assay. At 15min after pollination, mor1-1 stigmas at 29°C allowed pollen germination to occur, while very minimal germination could be observed in both wild-type Col-0 (restrictive and permissive) and mor1-1 (permissive) controls (Fig 5a). By 30 minutes, the difference was greater with stigmas from mor1-1 maintained at 29°C allowing much more rapid growth of pollen tubes through them (Fig 5b). By 1 hour after pollination, the average number of pollen germinated was insignificant between wild-type Col-0 and mor1-1 stigmas suggesting that within one hour after pollination, the compatible pollen grains on the control stigmas had caught up producing the same number of pollen tubes (Fig 5a). Thus, the results indicate that the switching of MTs from an elaborate network to a shortened depolymerized state following treatment at 29°C in mor1-1 stigmas, allowed a more rapid compatible pollen response to occur.

**EXO70A1 expression leads to a breakdown of the MT network in tobacco BY2 cells**

We have recently identified one of the exocyst complex members, EXO 70A1, as a compatibility factor in *Brassica* and *Arabidopsis*, but the exact mechanism by which it regulates pollination is still unknown (15). There is precedence for a role of exocyst complex in regulating MT
dynamics in mammalian systems. Previous studies have reported that an intact MT network is required for appropriate localization of EXO70 and the exocyst complex, while EXO70 and exocyst complex function to cause end instability of MT network (31). Based on these previous findings and our observations that both EXO70 and MT depolymerization are essential for compatible pollinations, we hypothesized *Brassica* EXO70A1 could cause depolymerization of the MT network. To test this, we used a transgenic tobacco BY2 cell system expressing GFP-TUA3 protein that illuminated the MT network (Fig 6, S4). Through biolistics, we transformed these cells either with an RFP construct or an RFP-EXO70A1 construct and examined the cells for presence of intact MT network. In cells expressing RFP alone, ~90% of the cells also displayed an intact MT network (Fig 6a,b), while in cells that expressed RFP-EXO70A1, the MT network was completely disrupted or absent more than 70% of the time (Fig 6a,b), indicating that EXO70A1 over-expression can de-stabilize the MT network.

**Discussion**

Unlike non-specific, unrelated pollen that are incapable of inducing the compatible response on the dry stigmas of Brassicaceae, SI pollen are fully capable of triggering the compatible pollen response (3,32,33). However, the presence of the self-pollen rejection response in the stigmas allows the active inhibition of the compatibility signaling, thus preventing self-pollen germination. During this active rejection process, binding of the pollen ligand SCR/SP11 induces the SRK-mediated SI pathway in the stigmas that stops any compatible signaling events. Thus, instead of being a linear module to self-rejection, the SI system is now proposed to negatively regulate the compatibility responses (3, 33). The identification of ARC1, an E3 ubiquitin ligase
that likely targets compatibility factors such as EXO70A1 following activation of SI response, is further evidence for such a theory (14, 15). The fact that blocking ubiquitination-mediated protein turnover in the stigma results in the breakdown of the SI response leading to incompatible pollen acceptance (14) indicates that post-translational targeting through ubiquitin tagging is a predominant mechanism during SI response. This proteasome-mediated regulation could be reflected in changes in protein abundance following SI pollinations. To identify potential targets of this pathway, we analyzed the stigma proteome through the 2D-DIGE approach, for differential regulation of protein abundance following the activation of the SI response.

From the DIGE analysis, 19 unique proteins were identified with reduced abundance following SI pollinations, and a large component of these proteins was identified as belonging to metabolic pathways. During compatibility responses, there is a requirement of increased energy for supporting the hydration, germination, penetration and growth of the compatible pollen tube. Thus, to maintain the stigmatic papillary cells in an energy efficient state, the SI system actively works toward stopping any SI pollen from developing on the surface conserving its resources for the compatible pollen. Since most of this energy is usually derived from increased glycolysis, a block in glycolytic pathway enzymes (enolase, fructose bisphosphate aldolase) could directly result in ATP depletion, blocking various energy-dependent processes. AtCIMS is the cobalamine-independent methionine synthase which is involved in the biosynthesis of methionine, providing methyl groups for various metabolic processes including the production of ethylene and biotin (34, 35). Interestingly, Biotin carboxylase is also down-regulated, which is a subunit of Acetyl-Co A carboxylase involved in the production of the precursor malonyl CoA for fatty acid synthesis. Fatty acids are important for successful pollen-pistil interactions as cer
mutants defective in long chain fatty acids produce pollen that are unable to interact with the stigmas (1,36). *Arabidopsis* fiddlehead mutants possess altered cuticular properties that support pollen hydration on non-stigmatic tissues (6). In Papaver, SI is known to target pollen soluble inorganic pyrophosphatases (sPPases) which are required to provide the energy for various biochemical pathways (37). Activation of SI in Papaver causes rapid phosphorylation of sPPases resulting in their reduced activity and increased inorganic pyrophosphate levels in the system (37).

Down-regulation of proteins from various compartments such as the cytosol, endoplasmic reticulum, mitochondria and chloroplast indicates a system under stress, leading to a general shut-down of various processes required for pollen germination on the stigma. If the SI response causes a general shut down of the various energy centers, then how does the stigmatic papillary cell keep itself receptive for the compatible pollen? This is likely reflected in the moderate levels of protein changes observed (1.24 fold to 2.13 fold) following SI pollinations, which suggests that the response is quite localized so that stigmatic papillary sites remaining unoccupied by incompatible pollen can still accommodate compatible pollen. This is quite consistent with the model of polarized secretion and molecular signaling observed exclusively in the foot region of the pollen attachment site (2, 38).

Two other candidate proteins with reduced abundance following SI pollinations are the S-locus related-1 (SLR1) and annexin. SLR1 is a stigma-specific secreted glycoprotein, and its suppression has been shown to result in reduced pollen-stigma adhesion. Treatment of stigmas to anti-SLR1 antibodies resulted in reduced pollen adhesion, confirming the role of SLR1 in mediating compatible responses (23). The observed down-regulation of SLR1 following SI
response indicates that suppression of SLR1 protein levels could be one of the modes through which SI operates. Annexins are multifunctional proteins that have been observed to accumulate in growing tips of secretory cells where they have been implicated in Golgi-mediated secretion (24). In Brassicaeae, the stigmatic papillae provide resources to the quiescent pollen to initiate metabolic activity in pollen grains following attachment to compatible stigmas. In this context, a regulatory circuit that controls targeted secretion of cargo necessary for pollen germination at the site of pollen attachment should be functioning on the stigmatic papillary side. In mammalian systems, annexins have been found to be associated with actin, exhibiting a calcium-dependent binding to F-actins, and are involved in actin bundling (39). Absence of annexin A2 resulted in the inhibition of actin-dependent apical transport of exocytotic vesicles in polarized epithelial cells (39). This suggests that annexins play an important role in vesicle trafficking along with calcium and actin. A similar role for calcium and actin is also observed in stigmatic papillae following compatible pollinations. A triphasic increase in cytosolic calcium is observed immediately after pollen attachment, and actin polymerization has also been shown to be required for compatible pollen acceptance (40, 21). In contrast, induction of SI response leads to a reduced, single calcium spike and complete de-polymerization of actin at the site of pollen attachment (40, 21).

In Brassica, prior to pollination, a large vacuole that is interconnected and branching into a globular and tubular vacuolar network can be observed at the apical region of stigmatic papillary cells (21). Following compatible pollination, this vacuolar network extended towards the site of pollen attachment, while following SI pollination, the vacuolar network was altered into a more fragmented pattern with few elongated vacuoles near the pollen attachment site (21). Given the fact that the vacuole contains the majority of ions, water and nutrients required for pollen
germination, it is likely that a complex involving calcium-induced exocytosis, mediated by both annexin and actin could result in proper delivery of the vacuolar cargo necessary for pollen germination. We recently showed that EXO70A1 functions as a compatibility factor, and at least one mode through which SI response could operate is by targeting EXO70A1 for degradation (15). EXO70A1 is part of the exocyst complex that is involved in polarized secretion and thus, could perform a compatible function along with actin and annexin in delivering vesicles to the site of pollen attachment. Interestingly, during cytokinesis in plants, a tubular-vesicular network is formed at the equatorial plane, and exocyst-like tethering complexes are proposed to initiate the fusion of multiple vesicles to give rise to the tubular-vesicular network (41). The proteomic data presented here with both annexin and actin showing reduced abundance following SI pollination suggests that the SI response could potentially target to downregulate annexin and the actin cytoskeleton, to reject pollen. Our proteomics approach failed to identify EXO70A1 as one of the proteins that is targeted by SI response; this could be due to low abundance of EXO70A1 in the system or due to the basic nature of EXO70A1 protein which would prevent it from resolving properly on a 2D gel.

Our pharmacological, cell biological and genetic approaches provide evidence that MT depolymerization is a pre-requisite for a successful compatible pollen acceptance to occur. The Arabidopsis mor1-1 mutant was used to examine the effects of MT depolymerization on the acceptance of wild-type compatible pollen. MOR1 (microtubule organization 1) belongs to the MAP215 family of microtubule-associated proteins (42). Point mutations that result in single amino acid substitutions in the N-terminal HEAT repeat of MOR1, result in, sparse, misaligned and shortened microtubules following exposure of these mor1-1 mutants to restrictive temperatures of 29° C (42). When maintained at 22° C, the MT organization in these mutants is
similar to wild-type. The MT shortening induced by the 29°C treatment of mor1-1 plants allowed for a much more rapid pollination to occur compared to the respective controls (Fig 5). This indicates that priming of the stigmatic tissue by prior MT depolymerization results in rapid pollen germination, likely due to a much faster rate of delivery of resources from stigmatic papillary cells to the site of pollen attachment. This is also consistent with the pollination data from the Brassica W1 pistils pre-treated with the microtubule depolymerizing drug, oryzalin (Fig 4).

We also observed that the overexpression of EXO70A1 in tobacco BY-2 cells resulted in the break-down of the MT network (Fig 6) and this corroborates well with previously published work in mammalian systems. In rat kidney cells, overexpression of EXO70 led to a breakdown of MT network and resulted in increased membrane additions (43). In addition, localized MT depolymerization through MT disrupting drugs was also shown to induce plasma membrane additions (44). This study proposed that EXO70 and the exocyst complex could mediate transfer of Golgi-derived vesicles from MT ends to actin and to the plasma membrane through regulating MT dynamics (43). In Arabidopsis, delivery of cellulose synthase (CESA) complexes were unaffected by depolymerization of microtubules, but MTs were required for guiding these complexes to particular sites on the plasma membrane (45). It was also shown that the motility of the small CESA compartment was mediated by MT depolymerization (45). Thus, a localized depolymerization of MTs following a compatible pollination could be an essential event for targeted secretions, which are required for stimulating the metabolic activity of the pollen.

While a reduced abundance of tubulin (alpha 2-4) was observed following SI pollinations, this apparently did not result in any drastic change in MT dynamics, although the dense network of
MTs was altered into a more helical network, suggesting that the reduction in the relative abundance of tubulin could be related to altered MT morphology (Fig 3a). Consistent with this, neither taxol nor oryzalin was able to alter the Brassica SI response suggesting that the regulation of SI is independent of the MT organization status. Nevertheless, the down-regulation of actin along with a previous study (21) documenting the de-polymerization of actin at the site of SI pollen attachment, indicate that, a general alteration of cytoskeleton could be partly responsible for manifestation of the SI response. In conclusion, through DIGE analysis of the stigma proteome for SI regulated proteins, we identified a change in the levels of a tubulin isoform which allowed us to uncover a novel and essential role for MT depolymerization in the stigma during the compatible pollen acceptance response. This analysis has also revealed a number of previously unknown candidate proteins which are being further investigated for their respective role during pollen-pistil interactions.

Acknowledgements

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References


Tables

Table 1. *Brassica* SI regulated proteins. Changes in protein abundance following SI pollination. Average ratio of protein levels, SI-pollinated (p)/unpollinated (U). The $t$ test $p$ values were calculated from three independent biological replicates of the samples.

Figure Legends

Fig 1. SI response leads to increased ubiquitination and down-regulation of a subset of proteins. a) Increased ubiquitination following SI (30min) was observed through Western blotting using anti-ubiquitin antibodies. b) 2-D DIGE analysis of SI regulated proteins. Proteins from W1 stigmas either unpollinated (U) or pollinated (P) for 60 min with self-incompatible W1 pollen were compared through 2-D DIGE using 24 cm pH 4-7 IPG strips followed by 12.5% SDS-PAGE. Proteins that are down-regulated by SI appear red and that are up-regulated following SI appear green and those that are unaffected are yellow. Interesting spots identified are marked with their respective identifications, the differentials predicted are shown in Fig S1b and the fold changes are shown in table 1.

Fig 2. Distribution of SI regulated proteins based on proposed molecular function and biological processes classification. The identified SI regulated proteins were classified according to Gene Ontology (GO) terms relating to biological processes and molecular function. The subcellular localization and biological functions are listed in table S1.

Fig 3. Compatible pollination induces MT break-down. Status of the MT network was observed through whole mounting of unpollinated (W1), 30min compatible pollinated (Westar
pollen) and 30 min SI pollinated (W1 pollen) stigmas, followed by immunostaining using anti-tubulin antibodies (a). Stacked images from a single papillary cell, pollinated for 30 min with compatible pollen showing a general breakdown of MT network close to the site of pollen attachment (b). (n=10 stigmas/treatment) P: pollen grain S: Stigmatic papillary cell ; Scale: 5μM

**Fig 4. Pharmacological modulators of MT organization affect compatible pollination.**

Average number of adhered pollen grains (a) and germinated pollen tubes (b) on W1 stigmas treated with either taxol or oryzalin, followed by compatible or self-incompatible pollinations with Westar and W1 pollen respectively. n=5; error bars are ± standard error. The representative DIC and aniline blue fluorescence images (to visualize pollen tubes) of stigmas from the compatible treatments are shown in 4c. HAP: Hours after pollination

**Fig 5. mor1-1 mutant allows rapid pollination at restrictive temperatures.**

Average number of pollen tubes on stigmas of Col-0 and mor1-1 plants maintained at either permissive or restrictive temperatures, followed by pollination for various times with Col-0 pollen maintained at permissive temperature (a). n=4; error bars are ± standard error. The representative aniline blue fluorescence images of mor1-1 pollinated stigmas (30min) from the different temperatures are shown in 5b.

**Fig 6. EXO70 expression leads to break-down of microtubule network.** Transgenic BY2 cells expressing GFP-TUA3 were either bombarded with RFP alone (n>50) or with RFP-EXO70A1 (n>50) and observed for intact MT network, 20h after bombardment. Number of RFP positive
cells with MT network (present/absent) was expressed in percentage (a). b. Representative cells expressing either RFP or RFP-EXO70A1 and the status of MT network are shown.
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Table 1: The pI and molecular mass values shown are theoretical. Score, the Mascot score. Cov., the protein sequence coverage (%). (No. peptide) Number of observable peptides per protein detected through Mass spec.
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**Fig 5.** *mor1-1* mutant allows rapid pollination at restrictive temperatures. Average number of pollen tubes/stigma following pollination for various times of Col-0 and *mor1-1* plants maintained at either permissive or restrictive temperatures (a). In all cases, pollen was from Col-0 maintained at permissive temperature. n=4; error bars are ± standard error. The representative aniline blue fluorescence images of *mor1-1* pollinated stigmas (30min) from the different temperatures are shown in 5b.
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