Investigating the Mechanism of Self-Incompatibility in *Papaver rhoeas* and Functional Transfer of *Papaver* S-Determinants to *Arabidopsis thaliana*

by

SABINA VATOVEC

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ABSTRACT

Flowering plants have evolved complex genetic mechanisms of self-incompatibility (SI) to overcome the problem of self-fertilization. SI is a cell-cell recognition system where the interaction of genetically linked pollen and pistil S-determinants prevents self-fertilization. In *Papaver rhoeas*, the pistil S-determinant is PrsS, a secreted protein of around 15 kDa. The pollen determinant, PrpS, encodes a novel transmembrane protein of around 20 kDa. Upon the interaction of incompatible PrsS and PrpS variants, the SI response is triggered, activating a signalling network. Rapid increases in cytosolic free calcium ([Ca\(^{2+}\)]\(_i\)) are followed by changes to the actin cytoskeleton and activation of a DEVDases, resulting in programmed cell death (PCD).

Within this thesis, three inter-related studies are described. Initially, we investigated the role of the ubiquitin-proteasomal system during SI in *Papaver*, the second study focused on the PrpS protein. Thirdly, we also created transgenic *Arabidopsis thaliana* lines expressing PrpS and PrsS, in order to investigate if the *Papaver* SI system might be functionally transferable to other plant species. We have demonstrated that PrpS binds the PrsS in an S-specific manner, while the functional analysis “*in vitro*” revealed that PrpS expressed in *A.thaliana* is functional and that just PrpS and PrsS are sufficient for a fully functional SI response in *A.thaliana* pollen.
I dedicate this thesis to my dearest:

my parents Tatjana and Lojze Vatovec and to my partner Miha Zakotnik
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LIST OF ABBREVIATIONS

ABA – abscisic acid
ABP – actin binding protein
ACE-H – apical cap extension-hydrodynamics
ACR – Arabidopsis crinkly
ACRE – Avr9/Cf9 rapidly elicited
AGO7 – argonaute7
AP – alkaline phosphatase
ARC1 – Armadillo-repeat containing 1
ARF – auxin response factor
ARM – armadillo
as-ODN – antisense oligonucleotide
AtMC – Arabidopsis thaliana metacaspase
Avr – avirulence
BAK – BRI-associated receptor kinase
BCIP – 5-bromo-4-chloro-3-indolyl phosphatase
BR – brassinosteroid
BRI – brassinosteroid insensitive
BSA – bovine serum albumine
BTB – bric-a-brac-tramtrac-broad
CLE – clavata/ endosperm surrounding region
CLV – clavata
CP – 20S core protease
CRP – cysteine-rich proteins
DISC – death inducing signalling complex
DMSO – dimethyl sulfoxide
ECL – enhanced chemiluminiscence
ECM – extracellular matrix
EDTA – ethylenediamine tetraacetic acid
EGF – epidermal growth factor
ETI – effector-triggered immunity
ETS – effector-triggered susceptibility
FDA – fluorescein diacetate
Flg – flagellin
FLS – flagellin sensing
GABA – gamma-aminobutyric acid
GFP – green fluorescent protein
GM – germination medium
GPCR – G-protein coupled receptor
GSI – gametophytic self-incompatibility
HECT – homology to E6-associated protein C-terminus
HR – hypersensitive response
IDA – inflorescence deficient in abscission
IEF – isoelectric focusing
Kan – kanamycin
LRR – leucine rich repeats
MAMP – microbe associated molecular pattern
MAPK – mitogen activated protein kinase
MG132 – N-(benzyloxy carbonyl)-leucinyl-leucinyl-leucinal
MLPK – M-locus protein kinase
MOMP – mitochondrial outer membrane permeabilization
MS – Murashige and Skoog medium
MW – molecular weight
NBT – nitro blue tetrazolium
NO – nitric oxide
NS – not significant
PAGE – polyacrylamide gel electrophoresis
PAMP – pathogen-associated molecular pattern
PARP – poly-ADP-ribose polymerase
PBA – proteasome beta-subunit A
PBS – phosphate buffer saline
PCD – programmed cell death
PCR – polymerase chain reaction
PFA – paraformaldehyde
PiSBP – *Petunia inflata* S-RNase binding protein
PrpS – *Papaver rhoea* pollen S
PRR – pathogen-recognition receptors
PrsS – *Papaver rhoea* stima S
PSK – phytosulphokine
PSKR – phytosulphokine receptor
PTI – PAMP triggered immunity
PUB – plant U-box
PVDF – polyvinyl difluoride
RAR – require for Mla12 resistance1
RBX – RING-box
RLK – receptor-like kinase
ROP – Rho-related GTPase from plants
RP – 19S regulatory particle
RT – room temperature
RT-PCR – reverse transcription PCR
s-ODN – sense oligonucleotide
SA – salicylic acid
SAM – shoot apical meristem
SAR – systemic acquired resistance
SBP – S-protein binding protein
SC – self-compatibility
SCR – small cysteine-rich
SDS – sodium dodecyl sulphonate
SDW – sterile distilled water
SGT1 – suppressor of the G2 allele of skp1
SI – self-incompatibility
SKP – S-phase kinase-associated protein
SLF/SFB – S-locus F-box/ S-haplotype-specific F-box
SLG – S-locus glycoprotein
SP11 – S-protein 11
sPPase – soluble inorganic pyrophosphatases
SSI – sporophytic self-incompatibility
SSK – SLF-interacting SKP1-like
SRK – S-locus receptor kinase
SUB – strubbelig
RING – really interesting new gene
ROS – reactive oxygen species
ta-siRNA – trans-acting short interfering RNA
TBS – Tris buffer saline
TBST – TBS-Tween
TM – transmembrane
TMM – too many mouths
TNF – tumor necrosis factor
TNFR – tumor necrosis factor receptor
TPD – tapetum determinant
TRAIL – TNF related apoptosis inducing ligand
TTS – transmitting tract specific
UbP – ubiquitin-26S proteasome
UT – untreated
VPE – vacuolar processing enzyme
Y2H – yeast 2 hybrid
CHAPTER 1

INTRODUCTION
1.1 PLANT SEXUAL REPRODUCTION

Plant reproduction is widely studied due to its immense importance for biotechnology and crop science, medicine, conservation biology and evolution. An important feature in the subject of plant reproduction is also the pollen tube, which is used as a model system for cell biology studies. The accessibility, easy \textit{in vitro} germination and rapid growth makes it an excellent model to study mechanical, genetic and molecular principles of polarised tip growth, cytoskeleton organization, ion fluxes, periodic behaviour, endo- and exocytosis and cell-cell signalling (Cheung and Wu, 2008, Feijó et al., 2001, Hepler et al., 2001, Moscatelli and Idilli, 2009, Geitmann, 2010).

During plant sexual reproduction the parental male and female plant organs combine their genetic material. The resulting offspring forms from a diploid embryo created when two haploid gametes, generated by meiosis, fuse together to form a diploid zygote. Plant reproduction starts with pollination. When a compatible pollen grain, containing the male gamete, lands on the stigma, it adheres and hydrates. Following hydration, pollen germinates and produces a pollen tube, which, with its strictly apical cell growth, elongates through the tissue of pistil (see Figure 1.1) until it reaches the ovule. The pollen tube then releases its two sperm cells to the embryo sac where a double fertilization occurs (Boavida et al., 2005, Cheung et al., 2010, Lord and Russell, 2002).

After fertilization, embryogenesis takes place that ends with the production of seed. The seed then germinates and new plant starts growing and developing organs. When the plant is mature, flower development starts and within the flower, male and female gametophytes form in spatially distinct areas of this organ via the processes of microsporogenesis and
megasporogenesis respectively. When these spores are fully formed, a new cycle of pollination can occur.

![Figure 1.1: Schematic diagram of sexual reproduction in model plant Arabidopsis thaliana. Pollen adheres to stigma, germinates and produces a pollen tube, which grows through the stigmatic papillae towards the ovaries where double fertilization occurs. Main pistil organs are indicated in white boxes and biological processes are indicated in blue boxes, while red arrows indicate pollen-pistil interactions. Image adapted from Feijó, (2010).](image)

The majority of flowering plants have, however, evolved mechanisms with which to cope when pollinated with genetically too similar pollen, i.e. incompatible pollen. This work focuses primarily on the pollination and prevention of self-fertilization aspect of plant reproduction, more specifically, on the phenomena of self-incompatibility, which will be described in greater detail in later section.

### 1.1.1 Pollen-pistil interactions: Pollination and fertilization

Pollen-pistil interaction involves an exchange of chemical signals between the male and female cells in a clear physical connection. Six major pollen-pistil interactions events have
been identified during pollination (Heslop-Harrison, 1975): pollen capture and adhesion, pollen hydration, germination of the pollen to produce a pollen tube, penetration of the stigma by the pollen tube, growth of the pollen tube through the stigma/style and entry of the pollen tube into ovule, resulting in discharge of sperm cells (Hiscock and Allen, 2008) (Figure 1.1).

The first step of pollination is the adhesion of pollen to the stigma. Pollen is released from the anthers in a dehydrated state and dispersed. After the initial capture of the pollen on the stigma, the pollen and stigma proteins combine and the start the complex pollen-stigma interactions that are under tight genetic and cellular control (Swanson et al., 2004). Adhesion depends on the stigmatic surface (wet or dry stigmas, depending on the presence or absence of stigmatic secretion) and on the pollen adhesion components (Zinkl et al., 1999).

It was demonstrated that in the formation of the pollen-stigma interface, pollen coat and stigma lipids small cysteine-rich proteins (SCRs), reactive oxygen species (ROS), nitric oxide (NO) and gamma-aminobutyric acid (GABA) are engaged (reviewed in Hiscock and Allen, 2008, Higashiyama, 2010).

In the compatible interaction, the pollen grain hydrates and germinates. Major factors implicated in this process in tobacco are lipids, present on stigma and pollen coat (Wolters-Arts et al., 2002), while in Brassica aquaporin-like protein MIP-MOD regulates the water supply to pollen (Dixit et al., 2001). Additional proteins identified in the hydration are pollen coat protein GRP17 in A. thaliana (Mayfield and Preuss, 2000), extracellular lipase EXL4 (Updegraff et al., 2009), water channel protein aquaporin in Brassica (Ikeda et al., 1997) and in tomato, pollen specific secreted protein LAT52, was demonstrated to interact with
stigmatic LeSTIG1, via receptor kinase LePRK2, to promote pollen tube growth (Tang et al., 2004). Pollen hydration step is important when we take into account SI mechanisms. It is usually the step, which is severely inhibited in case of incompatible pollen-pistil reactions in Brassicaceae (Takayama et al., 2000). During pollen germination pollen tube penetrates from an aperture in the pollen cell wall through a process of secretion of digestive enzymes, and enters the stigmatic tissue (see Figure 1.1). Following germination, the pollen tube starts growing through the stigma, style and transmitting tract of the pistil towards the ovary (see Figure 1.1), where it is mediated mostly by the female guidance cues. Pistil influence on the pollen tube gene expression was recently demonstrated to be much higher than previously thought. The pollen tube transcriptome of pollen tubes growing in vivo involves ~700 genes more than pollen tubes grown in vitro (Qin et al., 2009). Pollen absorbs nutrients from the female tissue and interacts with the several components including pectins, stigma/stylar cysteine-rich adhesion protein, that function in adhesion mediated pollen tube guidance, GABA, that forms a gradient which controls pollen tube guidance to micropyle and arabinogalactan proteins among which are best characterised transmitting tissue specific (TTS) glycoproteins that form a glycosylation gradient that promotes pollen tube growth (Cheung and Wu, 1999, Park et al., 2000, Palanivelu et al., 2003). In the transmitting tract of Nicotiana a thioredoxin h was also identified and reported to interact with S-RNases and play a role in pollen-pistil interactions during the S-RNase based SI (Juárez-Díaz et al., 2006). During the penetration of pollen through the transmitting tract, extensin-like and expansin-like activities have been reported, as well as pectin methylesterases, that enhance pollen tube tip dynamics (Stratford et al., 2001, Grobe et al., 1999, Bosch et al., 2005).
In the final stages of pollen tube journey through the pistil, female guidance of the pollen tube to the embryo sac is important for successful fertilization, and synergid cells play crucial role in this step. Two synergid cells are flanking the entrance to the egg cell at the micropylar end of the female gametophyte and are the source of the chemoattractants for the pollen tubes. Their role was established in the past decade by studies in *Torenia fournieri* (Higashiyama, 2002, Okuda et al., 2009). Mathematical model describing the dynamics of pollen tube attraction towards the attractants released from the ovules is also established (Stewman et al., 2010). It was recently demonstrated that two synergid cells in *Torenia* secret two cysteine-rich proteins (CRPs) LURE1 and LURE2 (Okuda et al., 2009). They were identified by expressed sequence tag analysis of the synergid cell of *Torenia*, 29% of all the clones they have sequenced encode CRPs (Okuda et al., 2009). CRPs LUREs belong to subgroup of defensin-like proteins and are secreted toward the micropylar end of the synergid cell. Down-regulation of the LURE1 and LURE2 protein resulted in the decreased rates of pollen tube attraction and recombinant expression of LUREs exhibited strong attraction of pollen tubes *in vitro*, suggesting that LUREs are involved in synergid cell pollen tube attraction (Okuda et al., 2009). In *Zea mays* it was also reported that small predicted transmembrane protein ZmEA1 (*Zea mays* Egg Apparatus 1) plays a pivotal role in pollen tube attraction and guidance through the micropyle into the female gametophyte (Márton et al., 2005).

After pollen tube grows into the female gametophyte, it releases its two sperm cells: one fertilizes the haploid egg cell to form a diploid zygote and the other fuses with the diploid central cell nuclei giving rise to the triploid endosperm. RLK FERONIA/SIRENE and LORELEI were identified to be required for pollen tube arrest and burst in *A. thaliana*, and defensin-like protein ZmES4 in *Z. mays* (Amien et al., 2010, Escobar-Restrepo et al., 2007,
Capron et al., 2008). ZmES4 activates $K^+$ channel KZM1, localised in the plasma membrane at the pollen tube tip. Interaction between ZmES4 ligand and KZM1 channel triggers rapid influx of potassium ions and osmotic stress resulting in pollen tube tip burst (Amien et al., 2010). During fertilization, each ovule is penetrated by a single pollen cell and it was demonstrated using two-photon microscopy that late arriving pollen tubes are repelled from the fertilized ovules or do not approach it at all (Cheung et al., 2010).

1.1.2 Pollen tube

Pollen is an organ that must survive in variety of different environments. To penetrate from the stigma towards ovule, pollen tubes have a rapid growth rate. They are one of the fastest growing plant cells known, reaching speeds of 200-300 nm.s$^{-1}$ (Cheung and Wu, 2008). This is an important characteristic, as the ovules are located at distances of several thousand times the diameter of the pollen grain away from the stigma (Cheung and Wu, 2008).

There are four distinct zones or domains in highly polarised growing pollen tubes:

1. the tip domain outlines an inverted cone and is rich in secretory vesicles
2. the sub-apical domain, that contains metabolically active organelles, like mitochondria and endoplasmic reticulum (ER)
3. the nuclear zone, that contains large organelles and male germ cells
4. vacuole domain, that contains large vacuoles and callose plugs (Aström, 1997, Boavida et al., 2005).

The apical and sub-apical domains are jointly referred to as clear region, while nuclear and vacuolisation zones are jointly referred to as the shank.
Growth is restricted to the apical area. As the tip advances a periodic callose deposition occurs in the zone far behind the tip, restricting the pollen protoplast to the most proximal region of the elongating tube. Pollen tube growth requires the presence of a tip-focused calcium gradient, an intact actin and microtubule cytoskeleton and is supported by active vesicle trafficking. Actin forms long filaments that help with the transport and create one of the hallmark features of the growing pollen tube, reverse fountain cytoplasmic streaming (Figure 1.2).

**Figure 1.2.** Reverse fountain cytoplasmic streaming in pollen tube occurs opposite as the water in the fountain, hence name reverse fountain streaming. Reverse cytoplasmic streaming involves the movement of the cytoplasmic contents, with organelles and vesicles, toward the tip in the apical region of the tube where secretory vesicles will be discharged via exocytosis while excess membrane and recycled proteins will be retrieved via endocytosis (Zonia and Munnik, 2009).

Actin cytoskeleton, Ca$^{2+}$ signalling and vesicle trafficking are regulated by ROP that are related to animal RAC (Rop/Rac), which have a role at the apex in regulating membrane trafficking and polar expansion (Fu et al., 2001, Yang, 2008, Fu, 2010). F-actin is present in the form of long filamentous cable-like structures in the shank of the pollen tube, that are responsible for the organelle and vesicle movement and reverse cytoplasmic streaming (Cai and Cresti, 2009). F-actin in the subapex is organised in a mesh-like structure, named actin collar, and in the tip, actin forms a dynamic network (Geitmann et al., 2000, Vidali et al., 2009). Actin binding proteins (ABPs) play a pivotal role in the dynamic actin signalling (Hussey et al., 2006).
In the recent years, ROS and phosphoinositides were demonstrated as additional pivotal signalling factors during the pollen tube growth, both of them tightly connected with Ca$^{2+}$ signalling (Fu, 2010). It was demonstrated that ROS forms a gradient in the apical domain of the pollen tubes and by inhibiting ROS production, pollen tube growth was inhibited, while Ca$^{2+}$ stimulates ROS production in pollen tubes (Potocký et al., 2007).

Cytosolic free calcium is an essential secondary messenger in many signal transduction processes in plants (reviewed in Dodd et al., 2010) and a tip-focused Ca$^{2+}$ gradient is also required for growing pollen tubes. This apical gradient is maintained by Ca$^{2+}$ influx at the tip via the opening of calcium channels. It has been shown to oscillate and it is thought that these oscillations are associated with the pollen tube growth in vitro, however, it was recently demonstrated in A. thaliana and N. tabacum, that such regular oscillations are not essential for pollen tube growth (Iwano et al., 2009, Holdaway-Clarke et al., 1997). [Ca$^{2+}$]$_{cyt}$ oscillations are controlling a wide range of intracellular processes, therefore the range of the [Ca$^{2+}$]$_{cyt}$ must be kept low (~2x10$^4$ lower than extracellular [Ca$^{2+}$]), for the stable equilibrium of Ca$^{2+}$ in the pollen tube (Iwano et al., 2009). Malhó et al., (1994) and Malho and Trewavas, (1996) have shown, that calcium is involved in the reorientation of the growing pollen tube. They suggested that this element is part of a signal transduction system, which allows pollen tubes to respond to directional signals in the style that guide it towards the ovary. Mechanism of regulating Ca$^{2+}$ is very complex and is interconnected also with other signalling pathways, such as ROP, ROS and phosphoinositidide signalling and influences many processes in the growing pollen tube; fusion of vesicles within the cell wall; regulation of cytoplasmic streaming; controlling the direction of growth, influencing actin cytoskeleton and mediating the SI response and programmed cell death (Malhó, 2006).
1.2 PROGRAMMED CELL DEATH (PCD)

PCD is an essential conserved cell death process used to remove the unwanted cells in plants during development and in response to external stimuli. It has been defined as a sequence of (potentially interruptible) events that lead to the controlled and organized destruction of the cell (Lockshin and Zakeri, 2004).

In animals there are three main types of cell death distinguished: apoptosis, autophagic cell death and necrosis (Kroemer et al., 2009). The major morphological features of animal apoptosis are reduction of cellular volume, chromatin condensation, nuclear fragmentation, plasma membrane blebbing, formation of apoptotic bodies, engulfment by phagocytes and clearance of dead cell contents by lysosomal degradation. Apoptosis in animals can be triggered by intrinsic stimuli, that can result from stress, radiation and others, and extrinsic stimuli, that is normally caused by binding of extracellular ligands, such as TNF-α (tumor necrosis factor), Fas ligand or TRAIL ligand (TNF related apoptosis-inducing ligand) to TNF receptors and activates caspase-dependent pathway (see Figure 1.3) (Spencer and Sorger, 2011). Death processes are triggered by initiator caspases-8 and -10 at DISCs (death inducing signalling complexes), that cleave the effector caspase’s-3 and -7 pro-domain thus forming an activated protease and consequentially cell death occurs (Figure 1.3). However, in some cases mitochondrial outer membrane permeabilization (MOMP) leading to diffusion of the mitochondrial intramembrane space into the cytosol, is also required to activate effector caspases (type II apoptotic pathway). Mitochondrial permeabilization is a well studied mechanism and is controlled by Bcl-2 protein family (Figure 1.3): Bax and Bak (multidomain proteins causing mitochondrial pore formation); Bcl-2, Mcl-1 and Bcl-xL (inhibitors of Bax and
Bak); Bid and Bim (activators of Bax and Bak); and Bad, Bik and Noxa (sensitizers that act antagonistically to BCl-2-like proteins) (Spencer and Sorger, 2011, Oberst et al., 2008).

Figure 1.3.: Schematic representation of death receptor-mediated apoptosis signalling. Upon the receptor-ligand interaction, initiator caspase-8 (C8) is activated at DISC and cleaves Bid that will subsequently activate Bax and Bak to oligomerize. These oligomers will create a mitochondrial outer membrane pore (MOMP) and caused the release of apoptotic regulators (cytochrome c, Smac/Diablo). Cytochrome c binds with caspase-9 (C9) and Apaf-1 to form apoptosome. In type II pathway XIAP inhibits caspase-3 (C3) and -7 activity and promotes their UbP degradation. However, Smac, that is also released from mitochondria binds to XIAP and inhibits its action, therefore enabling C3 and C7 to confer cell death. Image adapted from Spencer and Sorger, (2011).

In contrast to animals, plants do not exhibit apoptotic bodies and also there are no phagocytes in plants. So, a recent classification emerged that set the ground for morphological classification and terminology of plant PCD (van Doorn et al., 2011). According to this new classification, two classes of PCD are distinguished: vacuolar cell death and necrosis. However, some examples of PCD in plants do not fit in either of these classes, so they are classified as separate categories. In contrast to necrosis, which is a form of cell-death that results from acute tissue injury and provokes an inflammatory response, PCD is carried out in a regulated fashion. PCD in plants is less well studied compared with animals, but at present there are vast array of plant cell culture models...
and developmental systems are being researched by different research groups (Korthout et al., 2000, Sundström et al., 2009, Hatsugai et al., 2009, Thomas and Franklin-Tong, 2004, Coll et al., 2010, Bozhkov et al., 2004, Chichkova et al., 2010, Coffeen and Wolpert, 2004, Danon et al., 2004, Vercammen et al., 2004, Watanabe and Lam, 2011).

Plant vacuolar cell death is for example found during aerenchyma formation, xylem differentiation in vascular plants, leaf remodelling in *Monstera* or during the formation of embryo-suspensor (Gunawardena et al., 2004, Ohashi-Ito et al., 2010, Filonova et al., 2000). It is morphologically noticeable due to the increase of the vacuole volume, which contains hydrolytic enzymes that swallow up cell cytoplasm and degrade its contents. The disassembly of nuclear envelope was also observed as well as the formation of actin cables. Vacuolar cell death ends with the rupture of vacuole membrane or tonoplast and release of hydrolytic enzymes destroying the protoplast, however until rupture the organelles are intact (van Doorn et al., 2011). Biochemically, vacuolar cell death can be detected due to its autophagic activity, acidification of vacuoles, reorganisation of cytokeletal elements and activation of vacuolar processing enzymes (VPEs).

The second type of plant PCD is necrotic PCD, which occurs in pathogen recognition during the hypersensitive response (HR) or in cells challenged by necrotrophic pathogens (van Doorn et al., 2011). It is distinguished from the vacuolar type by the absence of increasing size vacuole, mitochondrial swelling and shrinkage of proplasts caused by the early rupture of plasma membrane with spilled and unprocessed corpses of necrotic cells. Biochemically, the necrosis can be detected by changes in mitochondria membrane potential, decreased respiration, accumulation of reactive oxygen species (ROS) and reactive nitrogen species (NO) as well as decline in ATP level (van Doorn et al., 2011, Christofferson and Yuan, 2010).
However, despite these two classifications, some unique types of plant PCD exist that cannot be attributed to either of those two groups. Such examples are HR cell death and PCD during self-incompatibility (SI) response (Thomas and Franklin-Tong, 2004, Hatsugai et al., 2009, Hofius et al., 2009).

### 1.2.1 Plant caspase-like proteases

Caspases play a critical role in animal apoptosis and are a family of cysteine proteases with specificity for aspartic acid, hence their name (Shi, 2002). There are no homologies to the caspases in plants, however, the caspase-like proteolytic activities have been identified in plants. Caspase-like activities have been detected using synthetic tetrapeptide substrates designed using the preferred cleavage site consensus of the members of the mammalian caspase family. Therefore the synthetic substrates are not caspase specific but they represent the optimal cleavage site of specific caspases (Stennicke and Salvesen, 2000). Caspase activities are often referred to using the amino acid sequence of the substrate cleaved (e.g. an activity against the substrate DEVD will be referred to as a DEVDase activity; see Table 1.1). Good biochemical evidence exists for the activation of plant proteases that cleave the substrates of caspases and therefore exhibit caspase-like activity in plants during PCD (for recent reviews see (Bonneau et al., 2008, van Doorn and Woltering, 2005, Piszczek and Wojciech, 2007, Woltering, 2010). Table 1.1 presents caspase-like activities that have been observed in plants to date.
Table 1.1.: Plant caspase-like activities, plant species where they were identified and reference.

<table>
<thead>
<tr>
<th>Activity</th>
<th>Species and tissue</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>YVADase</td>
<td>Tobacco leaf tissue</td>
<td>(del Pozo and Lam, 1998)</td>
</tr>
<tr>
<td>(caspase-1-like)</td>
<td>Barley embryonic suspension cells</td>
<td>(Korthout et al., 2000)</td>
</tr>
<tr>
<td></td>
<td>Tobacco BY2 cells</td>
<td>(Mlejnek and Prochazka, 2002)</td>
</tr>
<tr>
<td></td>
<td>White spruce seeds</td>
<td>(He and Kermode, 2003)</td>
</tr>
<tr>
<td></td>
<td>Pisum sativum seedlings</td>
<td>(Belenghi et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>A. thaliana fumonisin B-induced leaf lesion</td>
<td>(Danon et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>Nicotiana TMV infected leaves</td>
<td>(Kuroyanagi et al., 2005)</td>
</tr>
<tr>
<td></td>
<td>Seed integumens</td>
<td>(Hatsugai et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>P. rhoeas pollen</td>
<td>(Nakaune et al., 2005)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Bosch et al., 2010)</td>
</tr>
<tr>
<td>DEVDase</td>
<td>Barley embryonic suspension cells</td>
<td>(Korthout et al., 2000)</td>
</tr>
<tr>
<td>(caspase-3-like)</td>
<td>Tobacco BY2 cells</td>
<td>(Mlejnek and Prochazka, 2002)</td>
</tr>
<tr>
<td></td>
<td>White spruce seeds</td>
<td>(He and Kermode, 2003)</td>
</tr>
<tr>
<td></td>
<td>Pisum sativum seedlings</td>
<td>(Belenghi et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>Picea abies embryogenic cell line</td>
<td>(Bozhkov et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>Avena sativa leaves</td>
<td>(Coffeen and Wolpert, 2004)</td>
</tr>
<tr>
<td></td>
<td>A. thaliana seedlings</td>
<td>(Danon et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>P. rhoeas pollen</td>
<td>(Thomas and Franklin-Tong, 2004)</td>
</tr>
<tr>
<td></td>
<td>A. thaliana bact. inf. leaves</td>
<td>(Hatsugai et al., 2009)</td>
</tr>
<tr>
<td>IETDase</td>
<td>Avena sativa leaves</td>
<td>(Coffeen and Wolpert, 2004)</td>
</tr>
<tr>
<td>(saspase)</td>
<td>P. rhoeas pollen</td>
<td>(Bosch and Franklin-Tong, 2007)</td>
</tr>
<tr>
<td>LEHDase</td>
<td>Nicotiana benthamiana leaves</td>
<td>(Kim et al., 2003)</td>
</tr>
<tr>
<td>LEVDase</td>
<td>P. rhoeas pollen</td>
<td>(Bosch and Franklin-Tong, 2007)</td>
</tr>
<tr>
<td>TATDase</td>
<td>Xanthi tobacco leaves</td>
<td>(Chichkova et al., 2004, Chichkova et al., 2008)</td>
</tr>
<tr>
<td>VEIDase</td>
<td>P. rhoeas pollen</td>
<td>(Bosch and Franklin-Tong, 2007)</td>
</tr>
<tr>
<td></td>
<td>Barley seeds</td>
<td>(Borén et al., 2006)</td>
</tr>
<tr>
<td></td>
<td>Norway spruce embryogenic cell line</td>
<td>(Bozhkov et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>A. thaliana seedlings</td>
<td>Rotari &amp; Gallois – unpublished</td>
</tr>
<tr>
<td>VEIDase</td>
<td>Tobacco and rice</td>
<td>(Chichkova et al., 2010)</td>
</tr>
<tr>
<td>(phytaspase)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VKMDase</td>
<td>Avena sativa leaves</td>
<td>(Coffeen and Wolpert, 2004)</td>
</tr>
<tr>
<td>(saspase)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>metacaspases</td>
<td>A. thaliana seedlings</td>
<td>(Coll et al., 2010)</td>
</tr>
<tr>
<td></td>
<td>Picea abies embryo</td>
<td>(Sundstrøm et al., 2009)</td>
</tr>
</tbody>
</table>
Metacaspases are a distant homologues of caspases that are found in plants, fungi and protozoa, and were demonstrated to play a role in plant PCD. They are arginine/lysine specific cysteine proteases. In \textit{A. thaliana}, two types of metacaspases exist, type I and type II metacaspases (Watanabe and Lam, 2011, Sundström et al., 2009, Vercammen et al., 2004, Coll et al., 2010). Type II metacaspase, Tudor staphylococcal nuclease was identified in pine that is cleaved during PCD (Sundström et al., 2009). Recently, (Coll et al., 2010) demonstrated in \textit{A. thaliana}, that type I metacaspases, AtMC1 and AtMC2 antagonistically control PCD. AtMC1 acts as a pro-death protein required for HR cell death, while AtMC2 antagonizes it.

Vacuolar processing enzyme (VPE) is a plant legumain and it exhibits YVADase activity. It was implicated to play its role during PCD in several different species in various pathways. It plays a vital role in the rupture of the vacuole membrane and HR cell death in response to plant virus infection (Hatsugai et al., 2004). VPE was also identified in \textit{Papaver} pollen, by binding to DEVD-biotin probe (Bosch et al., 2010). It also exhibits DEVDase and IETDase activities in addition to predominant YVADase. Although it does not play a role in the SI response it is suggested to play a role in processing mitochondrial proteins (Bosch et al., 2010).

Another group of plant caspase-like proteins are saspases, subtilisin-like serine-dependent aspartate-specific proteases. They are involved in victorin-induced degradation of Rubisco during PCD in oats and exhibit VKMDase, VNLDase and VEHDase activities (Coffeen and Wolpert, 2004).

Chichkova et al., (2010) recently reported identification of phytaspase, a plant aspartate-specific protease, another subtilisin-like protease from tobacco and rice. It plays an
important role in regulation of PCD response to TMV infection and abiotic stresses, like oxidative and osmotic stresses. It possesses VEIDase activity and is localised in the apoplast, however during PCD it is partly re-localized inside the cell and so it might play a role in both positions (Chichkova et al., 2010).

There are many examples of PCD in plant development and in response to external stimuli (reviewed in Bonneau et al., 2008), and presented in Table 1.2 on page 22). An example of PCD in response to an external stimulus comes from the hypersensitive response (HR) after pathogen attack (reviewed in Coll et al., 2011), that can be compared to the SI-induced events in *Papaver*. So these two examples of PCD are described in more detail in the next two sections.

### 1.2.2 Hypersensitive response (HR-) induced PCD

Hypersensitive response (HR) is a mechanism of cell death that exhibits features of necrotic cell death but in addition also growth of the vacuole and rupture of the vacuole membrane followed by the release of lytic vacuolar contents and also VPEs. HR is a genetically controlled mechanism, based on the specific interactions between the products of the complementary genes of plant and pathogen, resulting in shrinkage of the cytoplasm, chromatin condensation, vacuolization and disruption of chloroplasts, all of that leading to rapid and localized cell death at the site of infection. In this way plants protect themselves and prevent spread of pathogens into healthy tissues (Dangl and Jones, 2001, Coll et al., 2011). When a microbial pathogen invades the plant tissue, a defence mechanism is initiated. Pathogens are inhibited by a combination of a layer of
dead cells, the local production of antimicrobial compounds, and the induction of systemic acquired resistance in the host (Dickman et al., 2001, Lincoln et al., 2002).

In the recognition of ‘non-self’, plant immune responses are of two types: one against general microorganisms and one against specific pathogen races (Chisholm et al., 2006, Jones and Dangl, 2006). The general defense mechanism is known as a pathogen- or microbe- associated molecular pattern (PAMP/MAMP) triggered immunity (Schwessinger and Zipfel, 2008, He et al., 2007a) and is triggered by extracellular surface receptors, called pattern-recognition receptors (PRRs). PRR recognition PAMP/MAMP results in the activation of defense responses against the pathogenic and non-pathogenic microbes (He et al., 2007a). In plants there are numerous PRRs, majority belonging to the receptor-like kinase (RLK) transmembrane proteins (Shiu and Bleecker, 2001). They are very specific and can recognize general features of microorganisms, such as bacterial flagellin (Zipfel et al., 2004, Gómez-Gómez and Boller, 2000). As a result of the co-evolution, plant pathogens have developed various strategies to overcome PAMP triggered immunity (PTI). One of them is effector-triggered susceptibility (ETS), which deploys PTI-suppressing pathogen effectors. The second and more specific defense mechanism against pathogen ETS is known as effector-triggered immunity (ETI), which is stimulated by plant resistance proteins (R-proteins) recognizing pathogens effector proteins, avirulence (Avr) proteins. ETI is very rapid and overblown defense response compared to PTI (Jones and Dangl, 2006, Boller and Felix, 2009).

One of the early actions in ETI is the oxidative burst; a generation of reactive oxygen species (ROS), such as hydrogen peroxide (H₂O₂), hydroxyl radicals (•OH), superoxide anions (O₂•-) and nitric oxide (NO) and development of localized PCD of infected cells and this is
known as HR, the final defense mechanism in plants (Nimchuk et al., 2003). As a result of the ETI activation, an increased accumulation of salicylic acid (SA) is also observed. SA induces the induction of various pathogenesis-related (PR) genes and the activation of systemic acquired resistance (SAR) (Nimchuk et al., 2003).

Few components regulating R-protein interaction and responses have been characterized. ACRE (Avr9/Cf9 rapidly elicited) genes are such example and several of them encode Ubiquitin E3 ligases (Craig et al., 2009). Among them is ACIF1 (Avr9/Cf-0-induced F-box1), a positive regulator of HR against fungal, bacterial and viral pathogens (van den Burg et al., 2008). Plant U-box proteins (PUB) have also been reported to act as a positive or negative regulators of plant immunity. PUB17, for example, an A. thaliana homolog of tobacco ACRE276 (also closely related to Brassica ARC1) acts as a positive regulator of HR in a response to the infection of Cf9 expressing tobacco with Avr9 peptide (Yang et al., 2006). The functional involvement of ACRE276 for a resistance against Cladosporium fulvum was demonstrated by silencing experiments where the ACRE276 silencing caused a reduced HR and breakdown of the Cf9-mediated resistance against C. fulvum, which could complete a whole lifecycle in RNAi silenced tobacco expressing Cf9 and in VIGS silenced tomato plants. The expression of AtPUB17 in ACRE276 RNAi plants lead to the restoration of Cf9/Avr9 induced cell death, thus demonstrating the role of the E3 ubiquitin ligases (Yang et al., 2006). On the other hand, PUB22, PUB23 and PUB24 act cumulatively as negative regulators of resistance, as triple mutant pub22/pub23/pub24 displayed increased oxidative burst and cell death (Trujillo et al., 2008).

Hatsugai et al., (2009) recently demonstrated the role of the proteasome in HR induced cell death as a response to bacteria attack. They identified a novel plant defense strategy to bacteria attack by membrane fusion of vacuole membrane and plasma membrane. Such a
fusion forms a tunnel from the inside of cell to the outside and enables the discharge of vacuolar content and defense proteins, as they demonstrated by the use of fluorescent proteins (Hatsugai et al., 2009). Outside the cell, plant defense proteins attack the bacterial cells, while hydrolytic enzymes in plant cell cause HR cell death (Hatsugai et al., 2009). Moreover, with the use of inhibitors of caspase-3 activity and proteasome inhibitors, Hatsugai et al., (2009) also demonstrated, that such a cell death and membrane fusion are proteasome-dependent and shown that proteasome subunit PBA1 acts as a caspase-3-like enzyme.

1.2.3 Self-incompatibility (SI) induced PCD in incompatible Papaver, Pyrus and Olea pollen

SI is a genetic mechanism that prevents self-fertilization by recognition and rejection of self-pollen, thus promoting outbreeding (described in details in Section 1.4.). There are several different mechanisms of SI in different species, like gametophytic SI in Papaver, gametophytic S-RNase-based SI in Nicotiana, Petunia, and Pyrus, sporophytic SI in Brassica, and some other less studied SI types, for example SI in Olea.

PCD was initially identified as a final and key downstream mechanism during SI in Papaver rhoeas pollen tube (Thomas and Franklin-Tong, 2004). PCD in the incompatible Papaver pollen is characterized by loss of pollen viability, depolymerization of actin cytoskeleton, activation of ROS and NO, SI-mediated DNA fragmentation and increased caspase-3-like activity (Bosch and Franklin-Tong, 2007, Wilkins et al., 2011). Recently, PCD was also implicated in response to SI in Pyrus pyrifolia (pear) pollen and Olea europea (olive) stigmatic cells (Serrano et al., 2010, Wang et al., 2009a, Wang et al., 2010). P. pyrifolia
from family Rosaceae is the first example of gametophytic S-RNAse based SI where nuclear DNA degradation was demonstrated in incompatible pollen tube in vivo and in vitro, prior to the inhibition of incompatible pollen tube (Wang et al., 2009a, Wang et al., 2010). In addition to DNA degradation, collapse of membrane potential and cytochrome c leakage in the cytosol were also demonstrated, indicating that SI in pear pollen could result in PCD of incompatible pollen (Wang et al., 2009a, Wang et al., 2010).

The O. europea SI system is less well understood and studied, however it is implicated with the stylar gametophytic SI (Serrano et al., 2010). Morphological changes with vacuolization of the cells, chromatin condensation, plasma membrane blebbing and loss of cell integrity were observed in the Olea stigmas. Trypan blue staining confirmed loss of papillar and pollen cell viability in incompatible conditions, DNA degradation was demonstrated by TUNEL and DEVDase activity were observed in both pollen and pistils of Olea under free pollination (Serrano et al., 2010).

1.3 RECEPTORS & CELL SIGNALLING

Cells coordinate their intracellular status with the external environment by means of a large number of clearly defined signalling pathways. The interaction of a typical cell with the extracellular matrix and with neighbouring cells influences a variety of signalling events. Signalling pathways can be extracellular, activated by an external stimuli or generated within the cell. There are various signals received by the cells: chemical signals (i.e. specific ligands, such as PrsS in case of Papaver rhoeas SI signalling, or hormones), electromagnetic radiation (such as light) or mechanical inputs (such as touch). The purpose of signalling is to
encode information and convey a message about the internal or external environment into chemical signals, to which the cell can respond. This is done through the reception of the signal, which is then transduced intracellularly until a cellular response is activated. Information is communicated either through direct protein-protein interactions or by diffusible elements, usually referred to as second messengers.

Plant signalling is mediated, as in many animal systems, by hormones, steroids, sterols and lipids, reactive oxygen species (ROS) and nitric oxide (NO), signalling peptides, and various small proteins (Matsubayashi, 2003, Mittler et al., 2004, Wang, 2004, Vert et al., 2005). Only some of these signalling molecules have their receptor components identified and the next section 1.3.1 gives an overview.

### 1.3.1 Plant receptors

Plant receptors have been classified according to their structural characteristics. The largest group are receptor-like kinases (RLKs), comprising a family of over 600 genes in *A. thaliana* (Shiu et al., 2004). Other plant receptors include G-protein coupled receptors (GPCRs) and other receptors for hormone signalling, some of which are components of the ubiquitin-26s proteasome (UbP) pathway (Chow and McCourt, 2006, Spartz and Gray, 2008). Wheeler et al., (2009) reported identification of a pollen transmembrane receptor PrpS, which was identified as the male determinant of SI in *Papaver*. This type of receptor has no homology to any other known transmembrane protein and it represents a completely new group of plant receptors. Table 1.2. summarizes the plant receptors and their corresponding ligands in cases where they have been identified.
Table 1.2: Receptor-ligand pairs identified in plants

Different types of receptors are highlighted by different colours for easier overview: RLKs are in blue, GPCR is in brown, PrpS is in red and hormone receptors are in green.

<table>
<thead>
<tr>
<th>Ligand</th>
<th>Receptor</th>
<th>Type of receptor</th>
<th>System</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brassinosteroids (BR)</td>
<td>BR11/BAK1</td>
<td>LRR-RLK</td>
<td>Plant Growth control: cell expansion and division, senescence, male fertility, pollen development fruit ripening, response to environmental factors</td>
<td>(Ye et al., 2011, Clouse, 2011, Kinoshita et al., 2005)</td>
</tr>
<tr>
<td>CLAVATA3 (CLV3)</td>
<td>CLV1/CLV2</td>
<td>LRR-RLK</td>
<td>Growth regulation of apical shoot meristem</td>
<td>(Clark et al., 1997, Brand et al., 2000, Ogawa et al., 2008)</td>
</tr>
<tr>
<td>TRACHEARY ELEMENT DIFFERENTIATION INHIBITORY FACTOR (TDIF)/CLE41 &amp; CLE44</td>
<td>PHLOEM INTERCALATED WITH XYLEM/TDF RECEPTOR (PXY/TDR)</td>
<td>LRR-RLK</td>
<td>Vascular cell division</td>
<td>(Fisher and Turner, 2007, Etchells and Turner, 2010, Lito et al., 2006, Hirakawa et al., 2008)</td>
</tr>
<tr>
<td>IINFLORESCENCE DEFFICIENT IN ABSCISSION (IDA)</td>
<td>HAESA/HAESA-LIKE2 (HAE/HSL2)</td>
<td>LRR-RLK</td>
<td>Control of floral abscission in Arabidopsis</td>
<td>(Cho et al., 2008, Stenvik et al., 2008)</td>
</tr>
<tr>
<td>systemin</td>
<td>SR160</td>
<td>LRR-RLK</td>
<td>Defense signalling</td>
<td>(Scheer and Ryan, 2002)</td>
</tr>
<tr>
<td>Phytosulphokine (PSK)</td>
<td>PSK receptor (PSKR)</td>
<td>LRR-RLK</td>
<td>Cellular de-differentiation and re-differentiation</td>
<td>(Matsubayashi and Sakagami, 2006)</td>
</tr>
<tr>
<td>Bacterial flagellin (flg22)</td>
<td>FLAGELLIN SENSING2 (FLS2)</td>
<td>LRR-RLK</td>
<td>Plant innate immunity; bacterial disease resistance; mediating stomatal response</td>
<td>(Gómez-Gómez and Boller, 2000, Zeng and He, 2010, Zipfel et al., 2004)</td>
</tr>
<tr>
<td>EPIDERMAL PATTERNING FACTOR1 (EPF1) &amp; EPF2</td>
<td>ERECTA (ER) &amp; TOO MANY MOUTHS (TMM)</td>
<td>LRR-RLK</td>
<td>Stomatal initiation and development</td>
<td>(Hara et al., 2007, Hara et al., 2009); (Hara et al., 2009, Hunt and Gray, 2009, Shpak et al., 2005)</td>
</tr>
<tr>
<td>Not identified</td>
<td>STRUBBELIG (SUB)</td>
<td>LRR-RLK</td>
<td>Epidermal maintenance and cell specification</td>
<td>(Chevalier et al., 2005, Yadav et al., 2008)</td>
</tr>
<tr>
<td><strong>TAPETUM DETERMINANT1</strong> (TPD1)</td>
<td><strong>EXCESS MICROSPOROCYTE ES1/EXTRASPOROGENOUS CELLS (EMS1/EXS)</strong></td>
<td><strong>LRR-RLK</strong></td>
<td>Anther tissue development and microsporocyte numbers</td>
<td>(Canales et al., 2002, Zhao et al., 2002, Yang et al., 2005, Jia et al., 2008)</td>
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<tr>
<td><strong>CLAVATA3/ENDOSPERM SURROUNDING REGION (CLE40)</strong></td>
<td><strong>ARABIDOPSIS CRINKLY 4 (ACR4)</strong></td>
<td><strong>TNFR-RLK</strong></td>
<td>Root meristem development</td>
<td>(Becraft et al., 1996, De Smet et al., 2008, Gifford et al., 2005, Stahl et al., 2009)</td>
</tr>
<tr>
<td><strong>S-locus cysteine rich (SCR) also known as S-protein11 (SP11)</strong></td>
<td><strong>S-locus –RLK (SRK)</strong></td>
<td><strong>SRK-RLK</strong></td>
<td>Self-Incompatibility in <em>Brassica</em></td>
<td>(Kachroo et al., 2001, Takayama et al., 2001)</td>
</tr>
<tr>
<td><strong>Abscisic acid (ABA)</strong></td>
<td><strong>G-PROTEIN COUPLED RECEPTOR (GPCR)</strong></td>
<td><strong>GPCR</strong></td>
<td>ABA signalling in guard cells</td>
<td>(Liu et al., 2007b)</td>
</tr>
<tr>
<td><em>Papaver rhoes stigma</em> S (PrsS)</td>
<td><em>Papaver rhoes pollen</em> S (PrpS)</td>
<td><strong>PrpS</strong></td>
<td>Self-Incompatibility in <em>Papaver</em></td>
<td>(Wheeler et al., 2009)</td>
</tr>
<tr>
<td><strong>Ethylene &amp; Cytokinins</strong></td>
<td><strong>ETR1, ERS1, ETR2, EIN4 &amp; ERS2 for ethylene CRE1, AHR2 &amp; AHR3 for cytokin</strong></td>
<td><strong>Two component Histidine kinase sensors</strong></td>
<td>Regulation of cell division and differentiation, promotion of fruit ripening.</td>
<td>(Chow and McCourt, 2006)</td>
</tr>
<tr>
<td><strong>Auxin, Jasmonic acid (JA)</strong></td>
<td><strong>Ubiquitin protein ligase SCF^{ETR1}, SCF^{CRE1}, F-box proteins</strong></td>
<td><strong>Plant growth and development</strong></td>
<td></td>
<td>(Katsir et al., 2008, Kepinski and Leyser, 2005, Dharmasiri et al., 2005)</td>
</tr>
<tr>
<td><strong>Gibberellins (GA)</strong></td>
<td><strong>GID1</strong></td>
<td><strong>Hormone-sensitive lipases</strong></td>
<td>Seed germination, stem elongation, leaf expansion, pollen maturation and induction of flowering</td>
<td>(Ueguchi-Tanaka et al., 2005)</td>
</tr>
</tbody>
</table>

### 1.3.1.1 Plant receptor-like kinases

RLKs represent nearly 2.5 % of the total number of proteins encoded in the genome of *A. thaliana* and have diverged to perform different roles (Shiu et al., 2004). The first plant RLK was identified in *Zea mays* (Walker and Zhang, 1990). Its role was determined because of
similarity to the *Brassica* S-locus glycoprotein, implicating RLKs in mediating self-incompatibility (Walker and Zhang, 1990).

RLKs encode a typical single-pass, transmembrane-spanning domain, with an extracellular N-terminal domain and an intracellular C-terminal serine/threonine (Ser/Thr) kinase domain. Recently it was reported that LRR RLKs, such as brassinosteroid insensitive 1 (BRI1) receptor and BRI1-associated receptor kinase (BAK1), can autophosphorylate on tyrosine residues in addition to Ser/Thr and are thus dual specificity kinases (Oh et al., 2009, Oh et al., 2010, Jaillais et al., 2011).

Plant RLKs consist of several subfamilies, recognised because of their distinct extracellular domains that display variable structural features that bind different types of ligands, like S-domain, leucine-rich repeats (LRR), epidermal growth factor repeats (EGF) and lectins (Figure 1.4.). These will be briefly explained in the next section.

![Figure 1.4: Subfamilies of plant RLKs, based on the structure of the extracellular domain: S-domain class RLKs based on S-receptor kinase (SRK) RLKs; leucine rich repeats (LRR) RLKs are the largest subfamily of RLKs; tumor-necrosis factor receptor (TNFR)-like class RLKs; epidermal growth factor (EGF) class RLKs; PR5 like receptor kinase (PR) class RLKs; lectin class RLKs (Adapted from the website of Torii Laboratory)](image-url)
1.3.1.1 LRR-RLKs

The largest subfamily of RLKs are the leucine rich repeats (LRR-RLKs) with over 200 members in *A. thaliana*, all containing tandem domains of approximately 24 amino acids featuring conserved leucines (Figure 1.4). Despite the similarity of the extracellular domain of the LRR-RLKs they bind a very diverse set of ligands, including: brassinosteroids, peptides, such as phytosulfokine and systemin and even secreted proteins (e.g. CLV3). Accordingly, LRR RLKs play critical roles in various stages of plant growth and development (see Table 1.2). For example, LRR RLKs are involved in the maintenance of the shoot apical meristem (SAM) where the CLAVATA (CLV) signalling pathway plays a central role in the regulation of SAM by controlling the size of the stem cell population (Fletcher and Meyerowitz, 2000, Clark et al., 1997). LRR RLKs are also involved in the brassinosteroid signal transduction pathway, which is regulated by the BRs. BRs are growth promoting steroid hormones (Grove et al., 1979). BR signal transduction and its mechanism is one of the best understood among the plant RLK family. Their role is associated with wide range of processes, such as light signalling, PCD, innate immunity response, male fertility, pollen development, fruit ripening, regulated senescence, cell expansion and division (for recent review see Clouse, 2011).

1.3.1.2 SRK RLKs

The *S*-locus receptor-like kinases (SRK) are described in greater detail in the section 1.4. Highly polymorphic SRK has been identified as a female determinant of *Brassica* SI and is specifically expressed in the plasma membrane of the stigmatic surface (Stein et al., 1996). It consists of an SLG–like extracellular domain, a single transmembrane domain and a
cytoplasmic Ser/Thr kinase domain (Figure 1.4) (Stein et al., 1991). The ligand for SRK is a pollen coat protein SCR/SP-11. The receptor-ligand interaction, activated in an allele-specific manner, triggers intracellular signalling cascade resulting in the pollen tube inhibition (Kachroo et al., 2001, Takayama et al., 2001) also reviewed in (Nasrallah, 2005, Takayama and Isogai, 2005). SRK RLKs also have another function. It was demonstrated that some RLKs could interact with Calmodulin (CaM), a small protein containing two globular domains with EF hands domain that bind Ca$^{2+}$ (Vanoosthuyse et al., 2003, Charpenteau et al., 2004, Gifford et al., 2007). Recently, a protein from the SRK family, a calmodulin-binding RLK (CBRLK1) was identified in A. thaliana and Glycine soya (Kim et al., 2009, Yang et al., 2010). CBRLK binds specifically to a Ca$^{2+}$-dependent calmodulin binding domain (CaMBD) in the C-terminus (kinase domain), and there is a possibility that the interaction of CaM with SRK CBRLK1 plays a role in recycling of RLKs in plants (Kim et al., 2009).

1.3.1.3 Other RLKs

Among other classes of plant RLKs, illustrated in Figure 1.4. are TNFR, EGF, PR and lecitin class of RLKs. TNFR class is best represented by the ACR4 (ARABIDOPSIS CRINKLY4). ACR4 is involved in the maintenance of the root meristem and is expressed in differentiating root cells. ACR4 acts together with CLE40 peptide, a CLV3 homologue, which is expressed in fully differentiated columella cells (cell layer at the root tip, which forms the root cap together with the lateral root cap cells), to control the expression of transcription factor WOX5 (WUSCHEL-RELATED HOMEOBBOX5) in the quiescent centre in order to maintain distal root stem cells (De Smet et al., 2008, Stahl et al., 2009). The root meristem
signalling works in opposite direction to the shoot signalling. In root signalling, signals originate from differentiated cells and determine the size and position of the root stem cell niche. Another role implicated for ACR4 is promoting initiation of lateral roots (De Smet et al., 2008).

1.3.1.2 G-protein coupled receptor

The GPCR family of receptors is also known as a seven transmembrane domain (7TM) receptor and it acts as a guanine nucleotide exchange factor. G-protein signalling cascade is composed of heterotrimeric protein complexes, which are composed of α-, β- and γ-subunits and GPCR. These complexes link perception by GPCRs with downstream signalling cascade (Jones and Assmann, 2004). A change in GPCR conformation upon the perception of the signal leads to the conformational change in the α- subunit and an exchange of GDP for GTP. This reorientation disrupts the interaction between α and β subunits and promotes dissociation of the complex into Gα and Gβγ dimers, which can interact with an array of downstream signalling molecules (Jones and Assmann, 2004).

The genome of A. thaliana encodes around 25 GPCR candidate proteins (Grill and Christmann, 2007). In plants this family of receptors mediates various responses to adjust cell growth and differentiation, metabolism, embryogenesis, abscisic acid (ABA) and pathogen-associated molecular pattern (PAMP) regulation of guard cell ion channels and stomatal apertures and ROS signalling (Zhang et al., 2011).
1.3.1.3 PrpS receptor in *Papaver rhoeas*

*Papaver* pollen *S*-determinant (PrpS) will be described in greater detail in the next section. PrpS is a novel transmembrane protein, localized in pollen plasma membrane. Its structure has not yet been resolved, but it most likely contains four transmembrane domains with an extracellular loop of 35 amino acids, which is the binding side for its cognate ligand (Wheeler et al., 2009). The ligand for PrpS is globular stigmatic protein PrsS. The *S*-specific interaction between PrpS and PrsS triggers downstream signalling cascade resulting in the PCD of pollen tube and thus prevention of self-fertilization (Thomas and Franklin-Tong, 2004). The identification of PrpS, a predicted novel transmembrane protein is of considerable interest, since very few plant ligand-receptor partners have been identified to date. Given that most of the transmembrane plant receptors identified to date are plant receptor kinases, the discovery of the PrpS could represent a novel class of receptor.

1.4 SELF-INCOMPATIBILITY

The majority of plants develop both male and the female organs on the same plant in close proximity. For most plants, like for mammals, the problem of self-fertilization is a serious one. In mammals inbreeding can result in genetic disease and poor health, and inbred plants also experience decreased fitness as the benefits of out-crossing are lost and inbreeding is encouraged with a resulting loss of genetic variability. In order to avoid self-pollination and ensure that their offspring benefits from hybrid vigour, plants have developed a number of different mechanisms. The most widespread mechanism for prevention of inbreeding is SI; a
system where self-pollen is “rejected” at the molecular level. It is estimated that some form of SI is present in ~60 % of all flowering plants (Hiscock and Kües, 1999).

SI is genetically controlled cell-cell recognition system where self-pollen is recognised by pistil and inhibited in an S-specific manner. There are two major mechanisms of SI: sporophytic (SSI) and gametophytic (GSI) (see Figure 1.5.).

Figure 1.5: An illustration of the genetic basis of gametophytic and sporophytic SI. For the sporophytic type, the SI phenotype of pollen is determined by the S-genotype of its diploid parent, thus each pollen grain carries the products of two S-genotypes. If an S₁S₃ pollen pollinates S₁S₁ or S₂S₃ stigma the outcome is incompatible interaction, while pollination of S₂S₄ stigma results in compatible pollination and fertilization. For the gametophytic type, the SI phenotype of pollen is determined by the S-genotype of its haploid genome, thus each pollen grain carries the product of one S-genotype. If S₁ and S₂ pollen lands on S₁S₂ stigma it results in self-incompatible reaction, while pollination of S₁S₃ stigma results in half-compatible situation where S₁ pollen is inhibited and S₂ allowed to grow and fertilize, and pollination of S₂S₄ stigma results in completely compatible situation. The spots in the pistils represent S-allelic component of the female S-determinants (i.e. in the sporophytic SI model, Brassica SRK₁ is represented by blue spots, SRK₃ by red spots, SRK₂ by yellow spots and SRK₄ by purple spots, while in gametophytic SI model spots represent S-RNases or PrsS proteins).

The pollen SI phenotype in SSI is determined by diploid genome of its parents (sporophyte), whereas in GSI the pollen SI phenotype is determined by its own haploid (gametophytic) genome. GSI is the most widespread SI system and has been extensively studied in the Solanaceae (Lycopersicon (tomato), Nicotiana (tobacco), Petunia (Petunia) and Solanum (potato)), Rosaceae (Malus (apples), Prunus (apricots, almonds, cherries) and Pyrus (pears)),

29
Plantaginaceae (*Antirrhinum* (snapdragons)), Poaceae (*Lolium*) and Papaveraceae (*Papaver* (poppy)). The SSI system has been extensively studied in Brassicaceae but also occurs and is studied in Asteraceae (Oxford ragwort), Convolvulaceae (sweet potato relative *Ipomea*) and Betulaceae (hazelnut). Interestingly, not only SSI and GSI have different mechanisms, but also there are also differences in GSI within different plant families, almost certainly as the result of SI evolving independently many times (Takayama and Isogai, 2005, McClure and Franklin-Tong, 2006, Allen and Hiscock, 2008, Klaas et al., 2011).

SI is usually controlled by a single, polymorphic S-locus with multiple alleles and can be compared with major histocompatibility complex (MHC) loci in mammals and mating type loci in fungi and algae. The exception is GSI in grasses, where SI is controlled by two loci, S and Z (Klaas et al., 2011). The S-locus in Brassicaceae, Solanaceae and Papaveraceae consists of at least two polymorphic genes, one encoding the male component and the second encoding the female component (Figure 1.6.) (Takayama and Isogai, 2005). This multi-gene complex is known to be inherited as one segregating unit. The variants of this gene complex are known as S-haplotypes rather than alleles. The recognition of self or non-self operates at the level of protein-protein interactions between male and female components. Incompatible interactions occur when both S-determinants are derived from the same S-haplotype.
Figure 1.6: (a) Male and female determinants of SI: in Brassicaceae the SSI type female component is an $S$-receptor kinase (SRK) and the male component is an $S$-cysteine rich (SCR)/$S$-locus protein 11 (SP11); in the GSI type found in Solanaceae, Rosaceae and Plantaginaceae the female determinants are $S$-RNases and male determinant are $S$-locus F-box (SLF)/$S$-haplotype-specific F-box (SFB); in GSI in Papaveraceae the female component are PrsS proteins while the male determinant was recently identified as PrpS protein (Takayama and Isogai, 2005, Wheeler et al., 2009). (b) A schematic diagram of the $S$-locus, which containing pollen (blue bars) and pistil (pink bars) determinants and basis of SI. The interaction of pollen and pistil determinant with the same $S$-haplotype (i.e. encoded by the same allele) results in incompatible response (green line with arrow ends), while in case of different $S$-haplotype interaction a compatible response is observed (green line without ends).

### 1.4.1 Sporophytic SI in the Brassicaceae

The $S$-locus encoded highly polymorphic receptor-ligand pair that determines SI specificity in *Brassica*, are the $S$-receptor kinase (SRK), that is expressed in the epidermal cells of stigma, and $S$-cysteine rich (SCR) protein, also identified as $S$-protein 11 (SP11), expressed in the pollen coat (Stein et al., 1991, Suzuki et al., 1999, Schopfer et al., 1999, Takayama et
The proposed mechanism of SI is presented in Figure 1.7. Incompatible pollen tube growth is inhibited very rapidly at the stigma surface.

**Figure 1.7**: Schematic model for recognition of self pollen and mechanism of sporophytic SI in *Brassica*. Male determinant of SI are SCR molecules expressed in pollen coat and female determinant are SRKs, expressed in stigmatic epidermal cells. Upon landing of self pollen on the stigma, SCR peptide is delivered to the stigma epidermal surface where it binds to the extracellular domain of cognate SRK. This causes autophosphorylation of the receptor and triggers signalling cascade, the cartoon proposes three different SI cascades. MLPK acts as a signalling intermediate. The cartoon in the middle illustrates SI model that involves ubiquitination of EXO70A1 by ARC1, while the cartoons on the side illustrate the existence of unknown compounds in the ARC1/Exo70A1-independent signalling pathways (Image adopted from Tantikanjana et al., 2010).

SRK, encodes a single pass transmembrane Ser/Thr kinase that is anchored to the stigma epidermis, presenting its highly polymorphic extracellular S domain to the ligand molecules on the pollen surface (Nasrallah, 2002). The extracellular, S-locus glycoprotein (SLG)–like extracellular domain is the site of interaction with pollen SCR ligand (Stein et al., 1991). There are several splice variants of SRK, such as eSRK (glycosylated extracellular SRK) and tSRK (membrane bound form of eSRK) (Shimosato et al., 2007). SRK has a plasma membrane localization, however, it is distributed in the smaller patches or SI domains, where it is kept in the ready-to-be-activated state (Ivanov and Gaude, 2009). SRK is also localised in the endosomes, below the plasma membrane surface, where the thioredoxin THL1 keeps it in the unactivated state (Ivanov and Gaude, 2009).
The S-specific interaction of SP11/SCR with SRK is localised on the interacting area of the stigmatic papillae, while the rest are ready to react on their own independent interactions. The S-specific interaction induces the autophosphorylation of SRK, triggering a signalling cascade that results in the rejection of self-pollen (Kachroo et al., 2001, Takayama et al., 2001, Ivanov et al., 2010). Inhibition occurs within minutes of pollination and the papillae cell wall may not even be penetrated. Downstream of SRK signalling were found proteins ARC1 (Armadillo-repeat containing 1) and MLPK (M-locus protein kinase) (see Figure 1.7) (Murase et al., 2004, Stone et al., 1999).

Membrane anchored cytoplasmic serine/threonine kinase was identified by positional cloning of the $M$ (modifier) locus (Murase et al., 2004) and is thought to act with phosphorylated SRK to phosphorylate ARC1. ARC1 was identified with Y2H screening for interactors of SRK kinase domain and it encodes a U-box E3 ligase with an arm repeat region (Stone et al., 2003, Stone et al., 1999). ARC1 is a positive regulator of the SI because the downregulation of ARC1 results in the partial breakdown of the SI (Stone et al., 1999). As an ARC1 binding protein, EXO70A1, component of an exocyst complex, was recently identified (Samuel et al., 2009). It normally promotes compatible pollinations as its reduced expression disturbed non-self pollen tube growth, therefore ARC1 mediated ubiquitination and proteasomal degradation of EXO70A1 is required for normal SI to occur (Samuel et al., 2009).

However, recent re-analysis of the Brassica SI signalling network revealed that it is much more complex then previously thought and that downstream signalling components identified so far (MLPK, ARC1, EXO70A1) are not required for SI, at least not for a Brassica SI in A. thaliana (see Figure 1.7) (Tantikanjana et al., 2010).
Tantikanjana et al., (2009) reported, that SRK also plays a role in pistil elongation. This additional role of SRK, whose expression was detected throughout the pistil, was identified in rdr6 mutants of transgenic Arabidopsis thaliana, expressing Arabidopsis lyrata SI genes. RDR6 functions in production of trans-acting short interfering RNA (ta-siRNA), and was found to enhance SI and causes stigma exsertion. In this mutant background they demonstrated that SRK gene further enhances pistil elongation and stigma exertion, which indicates that positive regulators of SI are regulated by ta-siRNAs and that SI trait and physiological distance between stigma and anthers have coevolved (Tantikanjana et al., 2009). They also identified targets of ta-siRNA as AGO7 (ARGONAUTE 7), that is an ARF (auxin response factor) and ARF3 and ARF4 are already demonstrated regulators of pistil development (Tantikanjana et al., 2009, Tantikanjana et al., 2010).

### 1.4.2 S-RNase based gametophytic SI

GSI in the Solanaceae (petunia, potato, tobacco, tomato, etc.), Rosaceae (apple, cherry, almond, pear, etc) and Plantaginaceae (Antirrhinum) share a common pistil S-determinant, an S-RNase. In these families inhibition of pollen tubes occurs in the transmitting tract of the style. Incompatible pollen grains lands on the stigma, hydrates and germinates as normal. The pollen tube grows through the stigmatic papillae and the transmitting tract of the style. It is only once pollen tubes reach about one-third of the way through the style that growth is arrested (Kao and Tsukamoto, 2004).

The S-RNases were identified by searching for pistil specific proteins that exhibited S-allele-specific differences in molecular weight and isoelectric point, as the allelic variants of protein involved in SI must be divergent in sequence in order to serve as a recognition
molecule. Using gain of function and loss of function experiments the S-RNase was demonstrated as the S-determinant in *Petunia* (Lee et al., 1994). S-RNases are produced in the transmitting cells and secreted in the transmitting tract where the pollen is growing and they contain five conserved domains and two hypervariable regions (HVa and HVb) (Matton et al., 1997). The second gene at the highly polymorphic S-locus that controls SI is the *SLF* (S-Locus F-Box)/*SFB* (*S*-haplotype-specific F-box) gene, which encodes the pollen S-determinant (Sijacic et al., 2004, Qiao et al., 2004a). SLF/SFB are members of F-box family of proteins, which generally function as a component of an E3-ubiquitin ligase complex and are involved in ubiquitin mediated protein degradation of non-self S-RNases. Recently new discoveries were made in *Petunia* where it was demonstrated that several related SLF factors encode pollen S-determinant and each SLF-type recognizes a range of S-RNases (Kubo et al., 2010).

Two proposed models, degradation and compartmentalization, can explain the self-compatibility and SI, and are presented in Figure 1.8. Degradation model of S-RNase-based SI (Figure1.7.a) is based on the destruction of S-RNases upon the interaction of SLF and S-RNase (Hua et al., 2008). It proposes that SLF determinants act collaboratively to prevent RNA degradation by non-self S-RNases and it also enables a wider variety of mating partners (Kubo et al., 2010). SLF determinants collaboratively mediate ubiquitination and degradation of the non-self S-RNases through the F-box domain, so that only self S-RNases function inside incompatible pollen tube (Meng et al., 2011). For example, during incompatible pollination of $S_1S_2$ style by $S_2$ pollen, the $S_1$- and $S_2$-RNases are taken up by $S_2$ pollen tube. The strong interaction between SLF$_2$ and $S_1$-RNAse would result in degradation of $S_1$-RNase as a consequence of ubiquitin-mediated proteasomal degradation, while the self $S_2$-RNase would remain intact. However, in the case of pollination of the $S_1S_2$ style by
heteroallelic $S_2S_2$ pollen, SLF$_1$ would preferentially interact with non-self $S_2$-RNases and SLF$_2$ with $S_1$-RNase, thus resulting in heteroallelic pollen compatible with $S_1S_2$ pistil (Meng et al., 2011). Using both allelic products of SLF, produced in heteroallelic pollen together mean that heteroallelic pollen is accepted by pistils, regardless of the $S$-genotype (Meng et al., 2011). However during incompatible pollination none of the SLF varieties bind and degrade self-$S$-RNases, which are therefore free to degrade the RNA of pollen tube (Kubo et al., 2010).

**Figure 1.8.** Two models for S-RNase based SI. (a) Degradation model: multiple SLF proteins are linked to certain $S$-RNase and act collaboratively. In the compatible situation (top) the array of SLF proteins (for example SLF$_1$ and SLF$_1$) bind to $S_x$-RNase thus preventing degradation of pollen RNA. During incompatible interaction (bottom) pollen SLF proteins, for example SLF$_1$, SLF$_2$, and SLF$_3$, do not bind self-$S$-RNase, which degrades the pollen RNA and thus inhibits pollen tube growth. (b) Compartmentalization model: during compatible pollination (top), the non-self RNase enters the pollen tube and are most likely compartmentalized into vacuoles and HT-B proteins are degraded. It is proposed that SLF promotes degradation of the non-self RNAses that are not compartmentalized by 26S proteasome. The RNA of the pollen remains intact. In incompatible pollination (bottom), the $S$-RNase and HT-B enter the endomembrane system by endocytosis. HT-B is required for $S$-RNase to exert its cytotoxic activity. When self $S$-RNase enters pollen tube, the conformational change occurs in SCF$^{SLF}$. The self-$S$-RNase-SLFI interaction inhibits the pollen protein responsible for HT-B degradation. Self-interaction therefore suppresses HT-B degradation. $S$-RNase is not degraded and its cytotoxicity results in RNA degradation and consequent pollen tube growth inhibition (Goldraij et al., 2006, Kubo et al., 2010, McClure, 2006). Image from McClure et al., (2011).

The S-RNase compartmentalisation model involves S-RNase compartmentalisation by the pollen tube endomembrane system (Figure 1.8.b) (McClure et al., 2011, McClure, 2009, Goldraij et al., 2006). Compartmentalisation is a common mechanism for controlling potential cytotoxins and probably protects both transmitting tract cells and pollen tubes.
S-RNases form complexes with other proteins: 120K, which is a 120 kDa glycoprotein abundant in the stylar ECM, and HT-B, a small asparagine-rich protein (McClure et al., 1999, Cruz-Garcia et al., 2005). S-RNase, 120K and HT-B are taken up by endocytosis and sorted to the vacuole (Goldraij et al., 2006). In a compatible reaction, S-RNase remains compartmentalised and HT-B levels are downregulated, so although S-RNase is present, it is not cytotoxic because it is sequestered. In an incompatible reaction, HT-B facilitates the S-RNase transport from endomembrane compartment to the cytoplasm where S-RNases, bound by 120K acts cytotoxic and degrade self-RNA thus leading to the inhibition of pollen tube growth (Goldraij et al., 2006, McClure, 2006).

1.4.3 Gametophytic SI in Papaveraceae

In *Papaver rhoeas*, the SI system is also under the control of a single, highly polymorphic, multiallelic, *S*-locus with gametophytic control of pollen *S*-gene expression (McClure et al., 1989). It is distinctly different to the S-RNase based SI system. In *Papaver*, the inhibition of the pollen tube is extremely quick and occurs soon after polarity is established during germination on the surface of the stigma. It has been estimated that around 66 *S*-alleles exist in poppy, several of which have been cloned (Foote et al., 1994, Walker et al., 1996, Kurup et al., 1998). The female component are small and secreted, ~15 kDa proteins, named PrsS (*Papaver rhoeas* stigma *S*) that act as a signalling ligand (Foote et al., 1994, Wheeler et al., 2009). *S*-stigma extracts and also recombinant PrsS have been shown to have *S*-specific pollen inhibitory activity when tested in an *in vitro* SI bioassay (Kakeda et al., 1998, Jordan et al., 1999).
1.4.3.1 Pistil S-determinant

The stigmatic S-alleles PrsS₁, PrsS₃, PrsS₈ and PrsS₇ from *Papaver rhoeas* and one from *Papaver nudicaule* have been cloned and sequenced nearly 2 decades ago (Foote et al., 1994, Walker et al., 1996, Kurup et al., 1998). The first one analysed was PrsS₁ and the protein was scraped directly from the stigmatic papillae of the flowers onto the isoelectric focusing (IEF) gel surface and separated by polyacrylamide gel electrophoresis (PAGE)-IEF (Foote et al., 1994). Silver staining revealed two proteins PrsS₁a and PrsS₁b with isoelectric points 7.55 and 6.90, while the sodium dodecyl sulphonate (SDS)-PAGE analysis showed that both separated into two proteins with size ~16.7 and ~14.7 kDa (Foote et al., 1994). Northern analysis showed that the gene was expressed specifically in stigmas with the expected temporal expression, while Southern analysis indicated that it was a single copy gene (Foote et al., 1994). PrsS₁ was produced as recombinant protein in *E.coli* (Foote et al., 1994).

PrsS proteins are easily extracted from *Papaver* stigmatic surface and have predicted signal peptide, suggesting they are secreted proteins. Secondary structure predictions suggest that they are comprised of 6 β-strands and 2 α-helixes, joined by 7 hydrophilic loops. They are cysteine-rich proteins with 4 identically positioned cysteine residues involved in disulphide bridge formation (Kakeda et al., 1998). It was demonstrated using a mutagenesis approach that residues in the hydrophilic loop number 6 are most likely to interact with the pollen S-determinant as the mutation of Asp79 of this loop resulting in complete loss of biological function (Kakeda et al., 1998).
1.4.3.2 Pollen S-determinant

Identification of pollen S-determinant has been long and difficult. A potential candidate was identified as integral membrane glycoprotein, S-protein binding protein (SBP) (Hearn et al., 1996). The problem was that SBP did not bind to PrS in an S-haplotype specific manner. The exact role and identity of SBP is yet to be found although it is thought to play an important role in SI.

So what are exactly criteria that a putative pollen S-determinant must meet? Firstly it must be linked to the pistil gene at the S-locus and it must have had co-evolved together with the pistil S-alleles. It must also be sufficiently polymorphic, exhibit correct temporal and spatial expression and it must play a functional role in the SI response.

Pollen S-determinant, PrpS, meets all of the above criteria (Wheeler et al., 2009). ORF of PrpS1 was identified very close (475 bp) to the pistil S gene when genomic DNA library, from the S1 containing Papaver plants, was screened using the S1 pistil gene as a probe. With two more alleles identified (PrpS8 and PrpS3), high level of polymorphism was confirmed. PrpS1 and PrpS8 sequences are only 59% identical at the amino acid level (for comparison, PrsS1 and PrsS8 have 64% sequence identity) (Wheeler et al., 2009, Kurup et al., 1998). PrpS1 and PrpS3 are 50% identical and PrpS3 and PrpS8 53% identical on amino acid level.

PrpS is a 20.5 kDa hydrophobic protein that is predicted to contain three to five membrane spanning domains. It was determined by expression analysis using RT-PCR that it is specifically transcribed in pollen with the highest levels of expression at the anthesis and the western analysis showed that it was only detected in the pollen membrane-enriched fraction. Immunolocalization analysis, using a specific PrpS1 antisera, visualized PrpS at the plasma membrane of S1 pollen tube (Wheeler et al., 2009).
Functionality of PrpS involvement in SI was demonstrated by the *in vitro* SI bioassays where PrpS peptide representing extracellular domain alleviated SI response, while randomised peptide did not. Antisense oligonucleotides (as-ODNs) and sense oligonucleotides (s-ODNs) were used in the *in vitro* assays to check that PrpS could function in an *S*-specific manner. Plants with the *S₁S₃* alleles were used and maximum inhibition was observed with *S₁* and *S₃* s-ODNs. *S₁* and *S₃* as-ODNs did not stop pollen tubes from growing while as-ODNs from *PrPS₈* had no effect on pollen inhibition (Wheeler et al., 2009). *S*-specific binding between PrpS and PrsS was detected with slot-blot peptide binding assay (Wheeler et al., 2009). PrpS has no homology to any protein in the existing databases and it is predicted to be a novel transmembrane protein (Wheeler et al., 2009). Due to the nature of the early poppy SI response, which includes an instantaneous influx of external Ca²⁺ via channels and rise in cytosolic Ca²⁺, the finding of pollen specific gene adjacent to stigma *S* that encodes a putative transmembrane protein could be of great potential significance.

### 1.4.3.3 Mechanism of SI in *Papaver*

When incompatible pollen lands on a receptive stigma, secreted PrsS proteins interact with pollen transmembrane PrpS and thus trigger a Ca²⁺-dependent signalling cascade, affecting many downstream cellular components, such as pyrophosphatase activity, MAPK, actin and tubulin cytoskeleton, ROS and NO, caspase-like activities, and ultimately leading to the PCD of the pollen tube (for recent review see Poulter et al., 2010b). Next sections discuss these events, which are also schematically presented in Figure 1.9.
Figure 1.9: Schematic model of the SI response in *Papaver*. Upon interaction of secreted stigmatic PrsS proteins and pollen transmembrane protein PrpS, a downstream signalling network is triggered, starting with an increase in K\(^+\) and [Ca\(^{2+}\)], who is signalling to the inhibition of pyrophosphatase activity and pollen tube tip growth, activation of MAPK, depolymerization of actin and microtubule cytoskeleton and appearance of punctate actin foci, activation of ROS and NO and activation of caspase-like activities, leading to the PCD of the pollen tube.

### 1.4.3.3.1 Role of Ca\(^{2+}\) in the SI response

Calcium plays an important role in the SI-specific response in *P. rhoeas* pollen tubes. Ca\(^{2+}\) imaging studies revealed a rapid, transient increase in pollen [Ca\(^{2+}\)], when pollen tubes were treated with incompatible but not compatible PrsS (Franklin-Tong et al., 1993, Franklin-Tong et al., 1995, Franklin-Tong et al., 1997). In the growing pollen tubes, a tip-focussed gradient of the intracellular Ca\(^{2+}\) concentration [Ca\(^{2+}\)], can reach 1-2 µM, while in the shank region basal levels of [Ca\(^{2+}\)]\(_i\) are low, with a mean value of ~200 nM. The SI challenge resulted in instantaneous and dramatic increase of [Ca\(^{2+}\)]\(_i\) in the shank and sub-apical region of pollen tubes, while tip-focused oscillating [Ca\(^{2+}\)]\(_i\) gradient rapidly disappeared (Franklin-Tong et al., 1995, Franklin-Tong et al., 2002). The SI induced increase in [Ca\(^{2+}\)]\(_i\) was blocked with calcium blocker La\(^{3+}\) and Gd\(^{3+}\), providing confirmation that Ca\(^{2+}\) influx is required for the increase in [Ca\(^{2+}\)]\(_i\) during the incompatible reaction (Franklin-Tong et al.,
Influx of extracellular Ca$^{2+}$ in the shank of pollen tube acts as a secondary messenger that triggers signalling cascades which results in irreversible inhibition of incompatible pollen tubes (Figure 1.9). Recently, a nonspecific cation channel activities were identified as being activated in incompatible *Papaver* pollen by using whole-cell patch clamping that are permeable for monovalent (K$^+$ and NH$_4^+$) and divalent (Ba$^{2+} \geq$Ca$^{2+} \geq$Mg$^{2+}$) ions upon the S-specific incubation with PrsS proteins (Wu et al., 2011). While Ca$^{2+}$ ions are well demonstrated to play an important role in pollen tube germination and growth, the role of K$^+$ is less well documented. So far in growing pollen tubes hyperpolarization-activated Ca$^{2+}$ channels, stretch activated Ca$^{2+}$-permeable cation channel and Ca$^{2+}$ pump ACA9 have been identified (reviewed in Michard et al., 2009). Role of K$^+$ was demonstrated recently in maize pollen, where small CRP ZmES4 protein was identified to activate the potassium channel KZM1 in the final step of fertilization and causes the K$^+$ influx and subsequent burst of the pollen tube that enables sperm cells to be released (Amien et al., 2010). There is a parallel with the PrsS-activated channel activity in *Papaver* pollen; both pollen systems respond to a clearly defined biological stimuli in a form of small CRP protein, that stimulates channel activity, leading to influx of K$^+$ ions, and in case of poppy pollen also to Ca$^{2+}$ ions (Amien et al., 2010, Wu et al., 2011).

### 1.4.3.3.2 Role of soluble inorganic pyrophosphatases in poppy SI

Soluble inorganic pyrophosphatases (sPPases) are enzymes that catalyse the hydrolysis of inorganic pyrophosphate to inorganic phosphate and are Mg$^{2+}$ dependent. sPPases are regulating biosynthesis and biochemical pathways in prokaryotes and eukaryotes. They have essential role in plant cells, driving biosynthetic reactions and generating ATP and
biopolymers such as pectin, hemicellulose and cellulose (Cooperman et al., 1992). Examples of cytosolic sPPases in plants are rare because plant sPPases are localized primarily to plastids rather than the cytosol (Gross and Ap Rees, 1986). Possible explanations for this could be that sPPases are abundant in metabolically active cells and thus, pollen tubes may express cytosolic sPPases because they require high metabolic activity to generate new membrane and cell wall for pollen tube extension (de Graaf et al., 2006).

In *Papaver* pollen the sPPases play a crucial role, as they provide the driving force for biosynthesis of pollen tube germination and growth. Ca\(^{2+}\)-dependent hyperphosphorylation of Pr-p26.1a/b (two soluble inorganic pyrophosphatases) leads to the reduced sPPase activity and occurs at around 90 s after SI (de Graaf et al., 2006, Rudd et al., 1996). sPPases can be inhibited by Ca\(^{2+}\) and by phosphorylation and this causes arrest of pollen tube growth, highlighting the importance of this regulatory mechanism during the SI response in *Papaver* pollen (Figure 1.9) (de Graaf et al., 2006).

### 1.4.3.3.3 Role of MAPK in poppy SI

Another target downstream of the Ca\(^{2+}\) signals, triggered by the SI response in poppy, is protein p56, named after its 56 kDa size, and is identified as MAPK (Li et al., 2007, Rudd et al., 2003). Its activation is peaking 10 min post SI implicated its role in signaling in incompatible pollen tube and regulation of PCD (Figure 1.8) (Li et al., 2007, Rudd et al., 2003).

MAPKs are the key elements of a of protein kinase cascade that can trigger responses to a wide variety of signals and stimuli. Collectively, different MAPK cascades regulate important cellular processes including gene expression, cell proliferation, cell survival and
death in eukaryotic cells (Chang and Karin, 2001). MAPKs were known to be functionally involved in regulating PCD in plants (Ligterink et al., 1997, Kroj et al., 2003) and the same was demonstrated for p56 in incompatible Papaver pollen (Li et al., 2007). The crucial role for p56 during SI in Papaver was demonstrated using specific MAPK inhibitor U0126, by whom the DNA fragmentation, PARP cleavage and SI-induced DEVDase activity were inhibited and loss of pollen viability rescued. Therefore, this represented the first evidence for MAPK signalling in pollen tubes during SI (Li et al., 2007).

1.4.3.3.4 Role of actin in poppy SI

One of the most rapid and dramatic physiological changes observed during Papaver SI response is a dynamic rearrangement of the actin cytoskeleton in the pollen tube, within 1-2 min after a challenge with incompatible PrsS proteins (Figure 1.9) (Geitmann et al., 2000, Snowman et al., 2002). F-actin filaments depolymerize and stable punctate actin foci are formed, whose size increases with time and are most characteristic 3 h post SI (Poulter et al., 2010a). Those punctate actin foci are the large aggregates of the F-actin that can reach up to 1 μm in diameter and are very resistant to high concentrations of latrunculin B, actin depolymerising drug that prevents polymerization of F-actin (Poulter et al., 2010a). Associated with the punctate actin foci are several ABPs, such as CAP and ADF (Poulter et al., 2010a).

Isolation of F-actin from untreated and SI pollen protein extracts and subsequent analysis by mass spectrometry FT-ICR-MS revealed a large amount of proteins implicated in binding the F-actin, however, the peptides were screened against Arabidopsis database. Among most prominent candidates that were more abundant in SI sample compared to the untreated
sample were: 14-3-3 proteins, Rab-like proteins, heat-shock proteins and chaperonins, and further investigations of them would provide important information about the complex actin signalling network in incompatible poppy pollen tube (Poulter et al., 2011, Poulter et al., 2010a).

An additional target of SI in incompatible pollen tube is also the microtubule cytoskeleton (Figure 1.9); cortical microtubules rapidly depolymerize upon the addition of PrsS, while the microtubules around the generative cell are more stable, still evident 1 h post-SI, as identified by immunolocalization studies (Poulter et al., 2008). Microtubule and actin cytoskeleton show different and distinct temporal alterations upon SI. There is one-way cross-talk between the two cytoskeletal elements since the actin depolymerization induced by latrunculin B can trigger depolymerization of microtubules but not vice versa, demonstrating that actin alterations act upstream of microtubule alterations (Poulter et al., 2008).

F-actin depolymerization or stabilization can push the cell into PCD and a first indication of this in *Papaver* pollen tubes was provided when DNA fragmentation was observed by using latrunculin B, and by jasplakinolide that stabilizes actin filaments, but was reduced when pollen was pretreated with the inhibitor of DEVDase activity, Ac-DEVD-CHO (Thomas et al., 2006). It was also established that stabilization of actin filaments by jasplakinolide prior to depolymerization of actin cytoskeleton by SI induction or latrunculin challenge could rescue pollen DNA degradation, thus preventing PCD (Thomas et al., 2006). Despite the evidence for the crosstalk between actin and microtubule cytoskeleton and the crucial role of actin cytoskeleton during the PCD, microtubule depolymerization or stabilization does not trigger PCD (Poulter et al., 2008).
1.4.3.3.5 Programmed cell death triggered by poppy SI

PCD was first indicated in incompatible *Papaver* pollen by Jordan et al., (2000) with the identification of nuclear DNA fragmentation and inhibition of pollen viability, which was observed in an S-specific manner. PCD was demonstrated with the rapid increase in cytochrome c release, detected upon pollen challenge with incompatible PrsS proteins, cleavage of bovine Poly-ADP-Ribose Polymerase (PARP), classic substrate for caspases, that generated a 24 kDa PARP cleavage product and activation of caspase-3-like activity (Franklin-Tong et al., 1996, Thomas and Franklin-Tong, 2004). Pretreatment of pollen tubes with the inhibitor of caspase-3-like (or DEVDase) activity, Ac-DEVD-CHO, prior to the SI induction resulted in a significant reduction in the amount of DNA fragmentation measured by TUNEL, thus demonstrating DEVDase activity in the SI-mediated DNA fragmentation (Thomas and Franklin-Tong, 2004). DEVDase activity and several other caspase-like activities were demonstrated by caspase-like cleavage assay using tetrapeptide substrates with fluorescent probe attached (e.g. Ac-DEVD-AMC). Live cell imaging revealed DEVDase activity between 1-2 h post SI in the cytosol of incompatible pollen tube, which moved to the nucleus with the progression of SI, peaking at 5 h post SI (Bosch and Franklin-Tong, 2007). Major caspase-like activity identified was DEVDase, however, a significant increase in VEIDase (caspase-6-like) activity was also detected and significantly lower YVADase (caspase-1-like) and LEVDase (caspase-4-like) activities (Bosch and Franklin-Tong, 2007).
1.4.3.3.6 Role of ROS in *Papaver* SI

ROS and NO are well established to regulate rate and orientation of pollen tube growth and are involved in the polarized growth of pollen tubes (Prado et al., 2008, Potocký et al., 2007). They were recently demonstrated to play an important role during S-RNase based GSI in incompatible pear pollen, where S-RNases disrupt tip-localized ROS and arrest ROS production in mitochondria, which subsequently decreased Ca$^{2+}$ currents and induced depolymerization of actin cytoskeleton and DNA degradation (Wang et al., 2010).

The role of ROS and NO signaling was also demonstrated *in vitro* in the incompatible *Papaver* pollen (Wilkins et al., 2011). Upon the S-specific PrsS challenge of pollen tubes, the increases in ROS were visualised using specific probes at around 5 min post SI, followed by an increase in NO at around 10-20 min (Wilkins et al., 2011). The crosstalk between the two signalling molecules was also established, using H$_2$O$_2$ that stimulates ROS production and DPI, that inhibit ROS, and examined the NO production using fluorescence probes (Wilkins et al., 2011). In incompatible poppy pollen tubes, the increases in [Ca$^{2+}$]$_{cyt}$ stimulated increase in ROS and NO. The connection with depolymerization of the actin cytoskeleton and subsequent appearance of punctate actin foci and an increase in DEVDase activity was also made, using ROS scavengers DPI (diphenyleneiodonium chloride) or TEMPO and NO scavenger cPTIO, placing ROS and NO, demonstrating that alleviated ROS and NO production reduced the formation of punctate actin foci and DEVDase activity. Thus, ROS and NO production is placed to act downstream of Ca$^{2+}$ signalling, but upstream of the actin cytoskeleton alterations and DEVDase activity (Wilkins et al., 2011).
1.5 Arabidopsis thaliana AS A MODEL FOR SELF-INCOMPATIBILITY

*Arabidopsis thaliana*, also known as thale cress or mouse-ear cress is a small flowering plant from family Brassicaceae. Its importance in plant science is not due to agronomic significance but due to its use as a model plant organism. It has a small genome of 125 Mb that was sequenced, which makes it extremely useful for genetic manipulations. *A. thaliana* is amenable and susceptible for genetic transformation by *Agrobacterium tumefaciens* and large collections of mutants exist in various ecotypes. *A. thaliana*’s lifecycle is very rapid, lasting only 6 weeks and it produces large amount of self-progeny due to its self-compatible state. That has made it an ideal plant for research as it can give results to test hypotheses quickly and there is a large worldwide research community working with it.

*A. thaliana* is already well-established model for expression of *Brassica*-type of SI (Rea et al., 2010, Nasrallah et al., 2002, Nasrallah et al., 2004). *A. thaliana* lost SI and became self-compatible ~1 million years ago by a mutation in the *SCR* gene, a 213 bp inversion (Tang et al., 2007, Tsuchimatsu et al., 2010). Studies on transgenic *A. thaliana* expressing *A.lyrata* and *C.grandiflora* S-determinants revealed that transition from SI to self-fertility occurred multiple times by independent mutations in different accessions of *A. thaliana* as they identified polymorphisms at the S-locus and at the SI modifier loci (Sherman-Broyles et al., 2007, Boggs et al., 2009c). These polymorphisms were explained by different strength of SI, from developmentally stable SI to very transient, weak SI or no SI, in different accessions of *A. thaliana*, when it was transformed with a functional SRK/SCR gene pair from *A.lyrata* (Boggs et al., 2009c). The approach of transgenic *A. thaliana* expressing *A.lyrata* and *C.grandiflora* SRK/SCR not only provided information on evolution of selfing but also
enabled new detailed studies of the basis of the SI S-specificity. Over 100 residues were
tested by mutagenesis in different regions of extracellular domain of SRK, but only six
amino acid residues clustered at the same positions within two variants tested were identified
to be responsible for SCR-specific activation of SI (Boggs et al., 2009a). The Colombia
ectype of A. thaliana was expressing the transient SI and when it was used in mutational
analysis a mutation was revealed in RNA-dependent RNA polymerase (RDR6) functioning
as a negative regulator of SI in trans-acting siRNA (ta-siRNA) production. This mutation
caused an enhancement of SI, pistil elongation and stigma exsertion, indicating dual role for
SRK in SI and pistil development (Tantikanjana et al., 2009).
New information and questions appeared regarding the role of ARC1 and MLPK1 proteins
as well as links between SI and pathogen resistance pathway (Rea et al., 2010). They crossed
SRK/SCR expressing A. thaliana plants with plants carrying the mutations in various plant-
defense implicated loci. If either of numerous plant resistance pathways examined (such as
etr1-1, ein2-1, npr1-1, pad4-1, rar1-21, sgt1b or eds1-1) would be important for SI pathway
then the breakdown of the SI was expected, however, in neither of the analysed crosses or its
progeny the breakdown of SI was observed, indicating that these pathways are not required
for SI and also illustrating the importance of such a transgenic A. thaliana model.

1.6 THE UBIQUITIN-26S PROTEASOME SYSTEM

So far I have given the background on the plant reproduction, signaling, PCD and SI, and I
have also made comparisons between SI and HR. Ubiquitin-dependent proteasomal
degradation (UbP) plays an important part of HR as well as some SI models. Part of my
studies, presented later in this thesis were involved with investigation of involvement of UbP during SI in *Papaver*, this section gives background on UbP pathway.

All aspects of plant physiology and development are controlled by regulated synthesis of new polypeptides and degradation of existing proteins. Posttranslational addition of the ubiquitin (Ub) to the proteins serves as a tag for the proteasomal degradation by the 26S proteasome, using ATP as an energy source (see Figure 1.10). Substrates for UbP degradation are long-lived proteins or damaged, misfolded or mutated proteins which can cause damage in the cell if they accumulate and so UbP degradation plays a vital role in regulation of various cell signaling processes in plant physiology and development (Vierstra, 2009, Dreher and Callis, 2007). The UbP complex is extremely important for plants since ~6% of proteome (> 1300 genes) in *Arabidopsis* genome is involved in the UbP pathway (Vierstra, 2009). Molecular genetic analyses have also connected individual components of the proteasome to almost all aspects of plant biology, including the cell-cycle, embryogenesis, photomorphogenesis, circadian rhythms, hormone signaling, disease resistance, senescence and self incompatibility (Vierstra, 2003, Vierstra, 2009).

![Figure 1.10.](image-url) The mechanism of UbP system. Ub activating enzyme E1 activates Ub, which is then transferred to an Ub conjugating enzyme E2 and attached to the target protein by Ub ligase E3. Once the polyUb tail is long enough, protein is targeted to the 26S proteasome and degraded, while Ub is recycled. Image adapted from Dielen et al., 2010.
Ubiquitin (Ub) is a 76 amino acid globular protein that is highly conserved in all eukaryotes (yeasts, animals, and plants). Proteins are marked for degradation when a covalent inter-Ub linkages are made from C-terminal glycine to lysine of the previous Ub to form a poly-Ub chain (polyubiquitination), although some proteins could also be ubiquitinated with a single molecule (monoubiquitination) or on multiple lysine residues by single Ub molecules (multimonoubiquitination) (Vierstra, 2009, Komander, 2009).

Ubiquitination is a multi-step process. Ub becomes activated by attachment to the Ub-activating enzyme, E1, in an ATP-dependent step. The Ub is then transferred to the second enzyme, called Ub-conjugating enzyme, E2. The final transfer of Ub to the target protein is mediated by a third enzyme, called ubiquitin-ligase, or E3, which is responsible for the selective recognition of appropriate substrate proteins. In some cases, the Ub is first transferred from E2 to E3 and then to the target protein. In other cases, the Ub may be transferred directly from E2 to the target protein in a complex with E3. Most cells contain a single E1, but have many E2s and multiple families of E3 enzymes. Different members of the E2 and E3 families recognize different substrate proteins, and the specificity of these enzymes is what selectively targets cellular proteins for degradation by the UbP pathway (Vierstra, 2009).

In *A. thailand* genes coding for different E3 components fall into different families, based on their domains. HECT (homology to E6-associated protein C-terminus), U-box and RING (really interesting new gene) (Figure 1.11.a,b) are composed from a single polypeptide which has characteristic recognition site for binding of E2 with Ub and different protein binding sites with motifs such as Armadillo (ARM), LRR, Serine/Threonine kinases and others (Jackson et al., 2000).
The largest group of E3 ligases are cullin-RING ligases (CRL), which contain subunits cullin, RBX1 (RING-box1) and other variable target and are further divided into subtypes (Figure 1.11.c).

**Figure 1.11.**: Structures of different types of E3 ligases: (a) HECT; (b) U-box/RING; (c) CRLs are divided in subtypes: SCF (SKP1 (S-phase kinase-associated protein1)-CUL1 (cullin1)-F-box) use F-box proteins for their target specificity, BTBs (bric-a-brac-tramtrac-broad complex) proteins and the third are DDB (DNA damage-binding) using WD40 domain for target recognition; (d) APC (anaphase promoting complex) E3 ligase. Image from Vierstra, 2009.

SCF ligases consist of 4 different polypeptides (SKP1 (S-phase kinase-associated protein1)-CUL1 (cullin1)-F-box (SCF) and RBX1). Specificity of the SCF complex is conferred by the F-Box domain, which contain on its C-terminus one of many protein interaction motifs (ARM, LRR, WD-40, Kelch, DEAD-Box,...) and signature F-Box motif on N-terminus for binding the SKP1, which bind CUL1 through SKP1 bridging protein (ASK1 in A. thaliana). Fourth component is RBX1 and it contains RING domain where E2-Ub binds. This unique organization of SCF E3s provides an effective mechanism for recognizing many substrates simply by exchanging F-Box subunits.

Second subtypes are E3s that use BTB (bric-a-brac-tramtrac-broad) complex proteins for target recognition and the third are DNA damage-binding E3s who use WD40 domain for target recognition (Craig et al., 2009, Dielen et al., 2010, Vierstra, 2009).
1.6.1 The 26S Proteasome

The 26S proteasome is a 2 MDa complex that degrades Ub-tagged substrates (see Figure 1.12). It is divided into two particles, the 20S core protease (CP) and the 19S regulatory particle (RP). The CP is created by the assembly of stacked heptameric rings of related α and β subunits (see Figure 1.12). Three of the seven β subunits are responsible for the proteolytic activity: β1 (PBA1), that has caspase-like activity, β2 (PBB1/2) that has trypsin-like activity and β5 (PBE1/2), that has chymotrypsin-like activity (Gu et al., 2010).

![Figure 1.12: Organization and structure of 26S proteasome.](image)

Recently, a direct connection between proteasome activation and plant PCD was established by Hatsugai et al., (2009), where they demonstrated, using genetical, biochemical and biological approaches, that proteasome subunit PBA1 had a DEVDase activity. It acts as a
long sought plant caspase-3-like enzyme and is responsible for the vacuolar membrane and plasma membrane fusion-induced HR cell death in response to bacteria attack. By using a proteasome inhibitors β-lactone and more specific Ac-APnLD-CHO, inhibitor of the β1 subunit, and caspase-3 inhibitor they demonstrated that the DEVDase activity and the PBA1 activity were abolished, and the HR was prevented (Hatsugai et al., 2009). In animals and yeast, β1 subunits of the proteasomes are long known to have caspase-like sites (Kisselev et al., 2003).

Identification of selective proteasome inhibitors has allowed a definition of the roles of the UbP pathway in various cellular processes (Lee and Goldberg, 1998, Kisselev and Goldberg, 2001). Proteasome inhibitors represent valuable tool to enhance cell content in the protein studies since proteins destined for rapid degradation are often hard to isolate or express. The most widely used are the peptide aldehydes, such as Cbz-leu-leu-leucinal (MG132), acting on a chymotrypsin-like activity, or lactacystin and β-lactone.

Recently a development of small membrane permeable fluorescent probe was presented that interact with the plant proteasome catalytic subunit and can allow the quantification of proteasome activities for in vitro and in vivo studies (Gu et al., 2010, Kolodziejek et al., 2011). It was also demonstrated that plant proteasome activity was inhibited using caspase-3 inhibitor. This activity-based probe will certainly expand studies on plant proteasomes in the future.
1.6.2 Ubiquitination and self-incompatibility

SI mechanisms are described in details in the section 1.4. The link between the S-RNase-based GSI system and ubiquitination is the male S-determinant, an F-box protein (SLF/SFB) which is part of an SCF ubiquitin ligase complex (SCF$^{SLF/SFB}$) (Meng et al., 2011, McClure et al., 2011). The compartmentalization models of SI presumes, that following pollination, self and non-self S-RNases are localized in the pollen tube. In the compatible interaction S-RNases are ubiquitinated by SCF$^{SLF/SFB}$ and afterwards degraded by the 26S proteasome, while S-RNases in the incompatible interaction are left intact and due to their cytotoxic activity they degrade RNA of the pollen tube and inhibit the pollen growth (Figure 1.13.b) (Goldraij et al., 2006, Meng et al., 2011). However, the issue emerging is, that SCF E3 ligases interact with the F-box protein with Skp1 and also with the specific substrate for degradation though the specific protein-protein interaction domain (ARM, WD40, leucine-rich repeats, etc) and none of that domains have been identified in SLF. Additionally, in the Antirrhinum, the SCF complex was termed a conventional SCF complex that contains novel SSK1 protein (SLF-interacting SKP1-like1), but is not encoded by polymorphic gene, but monomorphomorphic and might be important for the interaction with AhSLF2 (Zhao et al., 2010). In Petunia inflata, the SLF complex is also not as simple as initially thought but might be a component of new E3 ligase complex, containing PiCUL1-g and PiSBP1 ($P.inflata$ S-RNase Binding Protein1, a RING-HC protein), and it does not contain SKP1 or RBX1 (Hua and Kao, 2006). So in order to organise the complex, instead of Skp1, the PiSBP1 forms complex with PiCUL1-G and PiSLF, and also interacts with E2, like RBX1 (it plays a dual role) (Hua and Kao, 2006). It is believed they might have more general role as they are expressed in all the tissues (Hua and Kao, 2006).
Ubiquitination was also reported from the sporophytic SI (see Figure 1.13.a). In *Brassica napus* SI an U-box protein ARC1 (*Armadillo repeat containing protein 1*) was identified, which is a close homologue to the *A. thaliana* PUB17 and *Nicotiana* ACRE276, both playing an important role in hypersensitive response (Stone et al., 2003, Yang et al., 2006). The positive role of ARC1 was identified by the antisense suppression of *ARCI* in the stigma that lead to pollen tube growth of incompatible *Brassica* pollen (Stone et al., 1999). During the SI in Brassicaceae, ARC1 binds to the SRK (S-locus receptor kinase), a female determinant of *Brassica* SI, and functions downstream of activated SRK by interacting with Exo70A1 (see Figure 1.4.5.a) (Stone et al., 2003, Samuel et al., 2009). ARC1 was also demonstrated to colocalize with the subunits of the proteasome and the increase in ubiquitinated proteins was observed in the SI pistils (Stone et al., 2003). Exo70A1 is thus negatively regulated by SI through ARC1-mediated ubiquitination (Samuel et al., 2009).
1.7 SUMMARY AND OBJECTIVE

To summarize, in the introduction I have presented in details the SI system, which represents my main research. I also gave the background on plant reproduction, the role of pollen tube, PCD, HR, plant cell-cell signalling, plant receptors, use of *A. thaliana* as a model system and UbP degradation. The reason for such a broad range of topics is that this thesis presents research on three inter-related projects, studying the mechanism of self-incompatibility (SI) in *Papaver rhoeas* and *Arabidopsis thaliana*. My objectives were:

- Elucidating the possible involvement of ubiquitin proteasome (UbP) signalling pathway during SI in *Papaver* pollen

  The aim was to investigate the link between the two different mechanisms and to establish the role of UbP degradation during *Papaver* SI-induced PCD. Functional involvement of UbP pathway was investigated using inhibitors of proteasomal degradation, MG132, measuring levels of ubiquitination, inhibition of pollen tube growth, pollen viability and signalling to the caspase-like activity.

- Analysis of *Papaver* pollen S-determinant, PrpS and interaction between PrpS and PrsS

  The aim was to establish the nature of the *Papaver* pollen determinant and to characterize the PrpS binding to PrsS.

- Characterization and functional analysis of *Papaver* S-determinants in *A. thaliana*.

  This project was initiated in order to test whether the *Papaver* SI system might work when the S-determinants were moved to another species. Several lines of investigation were followed in order to achieve that. Initially, the segregation analysis and the expression analysis of PrpS and PrsS was carried out. Functional
analysis studies were then conducted \textit{in vitro} on transgenic \textit{A. thaliana} pollen expressing PrpS-GFP fusion proteins and \textit{in vivo}. \textit{In vitro} analysis investigated the major characteristics of the poppy SI response, such as \textit{S}-specific inhibition of pollen tube growth, formation of punctate actin foci and \textit{S}-specific loss of viability. Following successful \textit{in vitro} analysis, \textit{in vivo} analysis was attempted by performing crosses between \textit{A. thaliana} expressing PrsS protein and PrpS-GFP, respectively.
CHAPTER 2.

Materials and Methods
2.1. Plant material: *Papaver rhoeas*

2.1.1. *Papaver* plant cultivation

Seeds of *Papaver rhoeas* L. (Shirley) of both known and unknown pedigree were first allowed to germinate in humidified Petri dishes for ~2 days. Germinated seeds were potted in John Innes No. 1 potting compost and kept in a greenhouse at 15 °C. Resultant plants were thinned to one per pot after growing to a height of between 5 and 10 cm. Around 8 weeks later, plants were allowed to harden outside for around 4 weeks and were then transplanted to a field in rows with approximately 50 cm between plants and 90 cm between rows (Wheeler, 2001).

2.1.2. Determination of *Papaver S*-genotype

The *S*-haplotype was determined by pollinating each plant with pollen of known *S*-genotype. Once plants were in flower, two flowers, 1-2 days prior to anthesis, were emasculated using forceps and covered with a cellophane bag, to prevent entrance of insect pollinators. Emasculated flowers were pollinated the next day by application of pollen with a fine paintbrush directly onto the stigmatic rays. After pollination, cellophane bags were replaced over the flower and stigmas were harvested one day after pollination. The pollinated portion of the stigma was removed with a scalpel, placed in aniline blue, and left overnight to allow the stigmas to soften. A sample of the stigma was then placed on a microscope slide and observed under a microscope using UV illumination (see Figure 2.1.a,b). All families grown were two-class families and thus were pollinated with two classes of pollen, one to provide a fully incompatible pollination and one to provide a half-compatible pollination. Those
stigmas exhibiting fully incompatible pollination had easily identifiable callose in the pollen grains and few, if any, pollen tubes. Those exhibiting half-compatible pollinations had 50 % reacting as above whilst the remainder had no callose in the grain and long pollen tubes with callose plugs at intervals along the length of the tube.

Figure 2.1: Examples of aniline blue stained stigma squashes (a) compatible pollination and (b) incompatible pollination in poppy as visualised using UV-illumination.

2.1.3. Collection of Papaver pollen

Flowers with a long stem at 1 day prior to anthesis were harvested approximately 15 cm below the bud. Petals and sepals were removed while buds were placed in a cellophane bag and hung upside-down on the bench in the aerated glasshouse overnight, leaving anthers to shed pollen. Pollen was released by vigorous shaking, and collected by tipping into gelatin capsules. Capsules were then dried over silica gel at room temperature (RT) for about 1 h before freezing at -20 °C.
2.1.4. *Papaver* pollen tube growth in vitro

Pollen was hydrated for at least 30 min in a moist chamber at 25 °C. The shape of pollen grains was used to assess the state of hydration. When desiccated, pollen grains take an elliptical form, when hydrated pollen grains appear spherical. Following hydration, pollen was re-suspended in liquid germination medium (GM). Then pollen grains were plated on solid GM (GM plus 1.2 % agarose) in 9 cm Petri dishes. Normally, for 10 mg of pollen, 1.0 – 1.2 mL of GM was required. Pollen was grown for at least 45 min and assessed for percentage germination prior to experimentation. Following growth, pollen was treated as required for the experiment.

<table>
<thead>
<tr>
<th>Liquid Growth Medium (GM):</th>
<th>Growth Medium Salt Stock:</th>
</tr>
</thead>
<tbody>
<tr>
<td>13.5 % (w/v) sucrose</td>
<td>2 % (w/v) H$_3$BO$_3$</td>
</tr>
<tr>
<td>0.01 % H$_3$BO$_3$</td>
<td>2 % (w/v) KNO$_3$</td>
</tr>
<tr>
<td>0.01 % KNO$_3$</td>
<td>2 % (w/v) Mg(NO$<em>3$</em>)$_2$·6H$_2$O</td>
</tr>
<tr>
<td>0.01 % Mg(NO$<em>3$</em>)$_2$·6H$_2$O</td>
<td>7.2 % (w/v) CaCl$_2$·H$_2$O</td>
</tr>
<tr>
<td>0.036 % CaCl$_2$·H$_2$O</td>
<td></td>
</tr>
</tbody>
</table>

The solution was made as 13.5 % sucrose and then 0.5 % of GM salts stock was added to make up liquid GM.

**Solid GM:**
liquid GM with addition of 1.2 % agarose

2.1.5. Production of *Papaver* seeds

Emasculated *Papaver* flowers were pollinated with pollen of known S-genotype, covered with cellophane bag and left for approximately 6 weeks. After approximately 6 weeks seeds were collected from seed pods and stored in paper bags at 4 °C.
2.2. Plant material: *Arabidopsis thaliana*

2.2.1. *A. thaliana* cultivation

Equivalent of 50 μL of seeds of transgenic *A. thaliana* and untransformed Col-0 were sterilised with 500 μl of 20 % commercial bleach and continuously mixed with gentle shaking for 20 min. Seeds were then washed 4 times with 500 μL of sterile distilled water (SDW) and mixed by shaking for 15 min each time, then, centrifuged quickly to decant. Finally, seeds were resuspended in 200 μL of fresh SDW. Meanwhile, plates with Murashige and Skoog (MS) media were prepared by pouring autoclaved media with selection marker (50 μg μL⁻¹ Kanamycin, 25 μg μL⁻¹ Basta) for transgenic seeds or without for untransformed Col-0 seeds. Using a cut end sterile pipette tip seeds were plated in sterile conditions. The plated seeds were kept at 4 °C for two days in order to break dormancy and to synchronise seed germination before moving them to constant temperature room at 22 °C for two weeks. The amount of grown seedlings on the plates was determined for segregation analysis.

Transgenic *A. thaliana* lines with poppy S-determinants and untransformed Col-0 seeds were planted in pots labelled individually with yellow stick for genetically modified plants and white stick for untransformed plants. Immediately after transferring seedlings to pots, the tray was covered with cling-film for 2 days. *A. thaliana* plants were grown in pots, containing Levington M2 compost with Silvapearl in the greenhouse at 20 – 22 °C and under 16/8 h photoperiod conditions. Seeds were sown every two weeks. Plants were usually potted every two weeks and it normally took two to four weeks for the plants to flower. When plants flowered, each pot was placed in plant sleeve to contain the pollen spreading.
The plants were numbered from the first generation, with the number indicating original line, and the subsequent number of each generation after the dot. Generation numbers were separated by dot (.) or by the hyphen (-) for easier indication.

Murashige-Skoog (MS) media:
2.2 g L^{-1} of MS powder
pH adjusted to 5.6-5.8
1 % of agar
Media was autoclaved before use.

2.2.2. *A. thaliana* plant conditions

*A. thaliana* pollen is notorious for its difficulties in germination and only recently some detailed studies on *A. thaliana* pollen tube germination emerged (Fan et al., 2001, Boavida and McCormick, 2007, Bou Daher et al., 2009, Johnson-Brousseau and Mccormick, 2004). The difficulties regarding *A. thaliana* in vitro pollen germination were connected with its tricellularity (Boavida and McCormick, 2007, Bou Daher et al., 2009), and so far, majority of pollen being studied was from bicellular species (poppy, tobacco).

However, despite the studies performed on *A. thaliana* pollen germination, the work on *A. thaliana* was more demanding than *Papaver* pollen. There were numerous differences in the research procedures of pollen assays in *A. thaliana* and *Papaver*, the most obvious one was pollen collection. While poppy pollen collection occured once a year in a bulk batches with pollen being stored at -20 °C, fresh *A. thaliana* pollen was collected on a day of the experiment, as it was difficult to obtain a bulk batch. *A. thaliana* was producing new flowers every day and pollen was collected from day 0 flowers - stage 13 (Smyth et al., 1990). It was observed and reported that with the older flowers and older plants pollen germination rates
were decreasing (Boavida and McCormick, 2007), so in order to ensure reproducibility of results, only newly opened flowers were harvested for pollen collection.

2.2.3. Collection and germination of *A. thaliana* pollen

In the morning, 40 *A. thaliana* flowers (stage 13) were collected in a 500 μL microfuge tube containing liquid *A. thaliana* pollen GM (AtGM). Pollen was harvested by vigorous shaking and then the flowers were transferred into a second tube containing AtGM in order to obtain maximal amount of pollen. The process was repeated once more before the flowers were discarded. The AtGM with the pollen grains was centrifuged for 2 min at 3500 rpm and the excess liquid medium was removed. The pollen was pooled together, and resuspended in fresh AtGM and centrifuged again (2 min, 3500 rpm). Following centrifugation, the excess liquid was removed so that the final volume left in the tube for pollen germination was 100 μL. Pollen was pre-germinated for at least 45 min and assessed for percentage germination prior to experimentation. Following growth, pollen was treated as required for the experiment.

*Arabidopsis thaliana* liquid germination medium (AtGM):

- 18 % sucrose
- 0.01 % H$_3$BO$_3$
- 1.0 mM CaCl$_2$
- 1.0 mM Ca(NO$_3$)$_2$
- 1.0 mM MgSO$_4$

Sucrose was dissolved prior to the addition of the other salt components. pH adjusted to 7.0 and sterilised by filtration before use.
2.2.4. Collection of *A. thaliana* seeds

Trays with *A. thaliana* plants, which had fully formed siliques and no more flowering buds were lifted from the table so their water supply was disabled and plants were allowed to dry for about two weeks. Seeds were collected by gently shaking *A. thaliana* plants on the paper surface. Before placing them in a microfuge tube, seeds were sieved so as to remove remaining plant debris. Seeds from each plant were collected in separate clearly labelled tubes.

2.3 Experiments

2.3.1. Production of recombinant PrsS proteins

Transformant *E. coli* cells expressing PrsS₁, PrsS₃ and PrsS₈, separately, were plated from a glycerol stock, onto plates containing LB-agar medium with 50 μg mL⁻¹ ampicillin (LB-Amp) and incubated at 37 °C for about 16 h. The next day the colony was grown for 16 h in a 2 L flask containing 200 mL of LB-Amp at 37 °C on a shaker (200 rpm). Half of the culture (100 mL) was transferred into another 2 L flask with fresh 100 mL of LB-Amp. Protein over-expression was induced with 0.5 mM IPTG and incubated for 6 h at 37 °C with shaking at 200 rpm. A sample of the culture can be used for analysis of induction by SDS-PAGE. Culture was centrifuged at 5000 rpm at 4 °C for 10 min in Beckman centrifuge and pellets were stored at -20 °C until needed.

For the analysis, the pellet was resuspended in 200 mL of cold lysis buffer and centrifuged for 10 min at 5000 rpm at 4 °C. The pellet was resuspended in 20 mL of ice-cold lysis buffer, transferred in smaller tubes and then, lysozyme was added at a final concentration of
0.2 mg mL\(^{-1}\) and 0.25 mM PMSF. Reaction mix was incubated at 4 °C for 1.5 h before 3 mM sodium deoxycholate was added with 0.125 mM PMSF and incubated at 37 °C for 30 min, during which time the solution became very viscous. After that, samples were sonicated on ice for at least 5 times for 30 s pulses at 10 amplitude microns with at least 30 s break in between to allow cooling between each pulse. Following the sonication step, the solution was centrifuged at 5000 rpm for 20 min at 4 °C. The pellet was washed with 20 mL of cold lysozyme buffer. The whole cycle of pellet washing, sonications and centrifugation was repeated for 5-7 times, the pellet being frozen over night at -20 °C where necessary.

Proteins in inclusion bodies in the pellet were then solubilised in 6 M guanidine hydrochloride and 500 mM 2-mercaptoethylamine by shaking in orbital shaker for 4 h at 22 °C. In contrast with the previous protocol, which instructed to solubilize for 2 h, it was found that protein yield was increased by solubilising for 4 h. Following solubilisation, the suspension was centrifuged (15 min, 5000 rpm) at 20 °C and the supernatant was kept for protein refolding in a cold refolding buffer. Refolding took place at 4 °C by adding protein aliquots very slowly into the buffer while being stirred. The solution was incubated at 4 °C for 16 h with constant agitation. Refolded proteins were dialysed in three exchanges of at least 5 L of the cold dialysis buffer: in the morning, afternoon and overnight. The next day, the last set of the dialysis buffer was discarded (except for ~100 mL). Pressure was removed from the dialysis tubes by removing the clips. The tubes were then placed in the beaker with ~2 cm of dialysis buffer. Over that, the solid PEG6000 was generously sprinkled while the beaker was gently shaken. More PEG6000 was regularly added during the day until protein had concentrated to ~10 % of its original volume. When the desired volume was reached, PEG was washed from the tubes with distilled water, and proteins were aliquoted 1 mL to microtubes and snap-freezed in liquid nitrogen. Proteins were stored at –80 °C until required.
At every stage of the protocol, small samples were taken to check protein concentrations either on a protein gel (by SDS-PAGE) or with Bradford assay (Bradford, 1976).

**Lysis Buffer:**
- 50 mM Tris HCl, pH 8
- 100 mM NaCl
- 1 mM EDTA

**Refolding Buffer:**
- 100 mM Tris
- 2 mM EDTA
- 500 mM L-arginine hydrochloride
- 10 mM cystamine dihydrochloride
- 5 % glycerol
- pH 8

**Dialysis Buffer:**
- 50 mM Tris HCl, pH 8
- 100 mM NaCl
- 2 mM EDTA

### 2.3.2. Treatments of pollen tubes

Following successful pollen germination, pollen tubes were treated with PrsS proteins for SI response, incubated with various drugs or pretreated with drugs and then with PrsS proteins.

**PrsS proteins:**

Recombinant PrsS proteins (Kakeda et al., 1998) were dialysed over night at 4 °C from Tris buffer into GM (for experiments with poppy pollen) or AtGM (for experiments with Arabidopsis pollen) using dialysis tubing with 12000 – 14000 Mw cutoff. In 1 L of GM or AtGM, 1 mL of PrsS proteins could be dialysed. After dialysis, the protein concentration was determined by Bradford assay (Bradford, 1976) using bovine serum albumin as standard. For an induction of the SI response, PrsS proteins were added to pollen growing in *vitro* at a final concentration of 10 µg mL⁻¹ unless stated differently.
Treatment with N-(benzyloxycarbonyl)-leucinyl-leucinyl-leucinal (MG132):
Pollen tubes growing \textit{in vitro} were pretreated with 40 \(\mu\text{M}\) MG132 for 30 min prior to the addition of recombinant PrsS proteins and left for 5 h. Controls comprised of 40 \(\mu\text{M}\) MG132 treatment and were left for the same time. DMSO was used as a solvent control.

Treatment with Ac-DEVD-CHO:
Tubes were pretreated with 100 \(\mu\text{M}\) Ac-DEVD-CHO for 60 min prior to addition of S-proteins and left for 5 h. Controls comprised of 100 \(\mu\text{M}\) Ac-DEVD-CHO and were left for the same time.

2.3.3. Pollen protein extraction

Pollen tubes grown under specified conditions were collected from the Petri dish using a pipette with a cut tip, placed into a 1.5 mL microtubes (poppy pollen) and centrifuged (1 min, 13200 rpm) in a tabletop microcentrifuge. In the case of the Arabidopsis pollen experiments, centrifuging was for 2 min at 4000 rpm. Then the supernatant was discarded and an equal volume of 2X Tris extraction buffer with protease inhibitor or TM extraction buffer was added to the volume left in the microtube to make a final 1X concentration. Samples were roughly grounded using plastic grinders in the microtube before being snap-frozen in liquid N\(_2\) and stored at -20 °C until required.

Sample protein extracts were prepared by sonication 5 x 5 s at 10 amplitude microns (poppy pollen) or 5 x 2 s (Arabidopsis pollen) and kept on ice during sonication to prevent overheating. After sonication, samples were kept on ice for 15 min and then centrifuged (30 min at 13200 rpm at 4 °C). The supernatant was recovered into a microtube and then
centrifuged again (10 min, 13 200 rpm at 4 °C). The pellet was frozen at -20 °C. Clear supernatant was carefully recovered into a microtube. Samples were kept on ice during all the steps.

The protein concentration was determined by Bradford assay (Bradford, 1976) and then samples were used for SDS-PAGE or caspase-like activity assays.

<table>
<thead>
<tr>
<th>2X Tris extraction buffer:</th>
<th>Extraction buffer for caspase assay:</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 mM Tris-HCL pH 8</td>
<td>50 mM NaOAc</td>
</tr>
<tr>
<td>200 mM NaCl</td>
<td>10 mM L-cysteine</td>
</tr>
<tr>
<td>2.0 mM EDTA</td>
<td>10 % glycerol</td>
</tr>
<tr>
<td>1.0 M sucrose</td>
<td>0.1 % Chaps</td>
</tr>
<tr>
<td>2 X final protease inhibitor cocktail</td>
<td>pH 5 and pH 6</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>2x Tris Extraction buffer for TM proteins:</th>
<th>Extraction buffer for TM proteins:</th>
</tr>
</thead>
<tbody>
<tr>
<td>2x Tris extraction buffer</td>
<td>30 mM Hapes</td>
</tr>
<tr>
<td>0.2 % Triton X-100</td>
<td>2 % Triton X-100</td>
</tr>
<tr>
<td></td>
<td>300 mM NaCl</td>
</tr>
<tr>
<td></td>
<td>30 % Glycerol</td>
</tr>
<tr>
<td></td>
<td>5 mM MgCl₂</td>
</tr>
<tr>
<td></td>
<td>2 mM β-mercaptoethanol</td>
</tr>
<tr>
<td></td>
<td>pH 7.5</td>
</tr>
</tbody>
</table>

### 2.3.4. Bradford Assay

Protein concentration was estimated through the Bradford assay (Bradford, 1976) using BSA as standard. Protein assay reagent (BioRad, UK) (which contains Coomassie® Brilliant Blue G-250) was added to diluted protein samples, and its absorbance measured at 595 nm with a spectrophotometer.

For most of the experiments, protein concentration needed to be calculated as accurately as possible, so two replicates were measured for each sample and also compared with a BSA.
2.3.5. SDS-Polyacrylamide gel electrophoresis (SDS-PAGE)

2.3.5.1. Preparation of SDS-PAGE

Gels were cast using the BioRad self-assembly kits using standard procedures.

2.3.5.2. Protein staining on the poliacrylamide gel

After electrophoresis, gels were soaked for 1–2 h at room temperature in a Coomassie staining solution and then, destained by incubating the gel in a destaining solution for 2 h or until the bands were clearly apparent. Finally, gels were dried using vacuum drier for 1.5 h at 70 °C.

Coomassie staining solution: 0.1 % Coomassie Blue R-250 45 % (v/v) methanol 10 % glacial acetic acid

Destaining solution: 30 % (v/v) methanol 10 % (v/v) glacial acetic acid

2.3.6. Western blot

2.3.6.1. Protein transfer

After protein separation by SDS-PAGE the resolving gel was then transferred onto a sponge and two sheets of filter paper (Whatman) soaked in the protein transfer buffer. A nitrocellulose membrane (Hybond-C extra, Amersham) was cut to size and laid over the gel. A glass rod was used to remove any air bubbles. Two more sheets of filter paper (Whatman) were laid on top of the membrane and finally a sponge was laid on it. The layered arrangement was sandwiched between two electroblotting pads (BioRad). This was then inserted into the electroblotting tank (BioRad), which was filled with protein transfer buffer.
An ice block was also placed in the tank to prevent a overheating due to the high voltage used for blotting. BioRad power packs were used to blot the gel at 350 mA for 2 h. Following transfer, the nylon membrane was removed and placed in blocking solution at 4 °C for 16 h.

**Protein transfer buffer:**
- 20 % (v/v) methanol
- 0.2 M glycine
- 25 mM Tris

**Blocking solution:**
- 10 % (v/v) 10x TBS
- 0.01 % Tween
- 5 % (w/v) milk powder

**10 x Tris Buffered Saline (TBS):**
- 24.22 gL⁻¹ Tris
- 80 gL⁻¹ NaCl
- pH 7.6

**2.3.6.2. Protein immunodetection**

After blocking the membrane, it was incubated with primary antibodies (diluted at varying concentrations depending on the antibody being used), in a blocking solution for 2 h at room temperature. Then, three washes of 10 min each blocking solution were carried out and then, incubated with the secondary antibody. If the primary antibody was raised in mouse, an alkaline phosphatase-conjugated anti-mouse immunoglobulin was used as the secondary antibody (alkaline phosphatase detection), while if the primary one was raised in goat, a horse radish peroxidase-conjugated anti-goat immunoglobulin was used (enhanced chemiluminescence detection) diluted 1:5000 in blocking solution for 1 h. The blot was then washed 1x 10 min in blocking solution and 3x 10 min in 1X TBS-Tween.
Primary antibodies:
Monoclonal anti-ubiquitin (Sigma); titre 1:5000
Monoclonal anti-ubiquitin (Santa Cruz); titre 1:5000
Monoclonal anti-ubiquitin (donated by Dr Ari Sadanandom); titre 1:2500
Poly-clonal anti PrpS₁ C terminus (60C); titre 1:500 or 1 :1000

Secondary antibodies:
Anti-mouse HRP; titre 1:5000
Anti-mouse AP; titre 1:5000
Anti-rat AP; titre 1:5000
Anti-goat HRP; titre 1:5000

2.3.6.2.1. Enhanced chemiluminescence detection

ECL western blotting reagents were used to detect the HRP conjugated secondary antibody. Solutions A and B (Yakunin and Hallenbeck, 1998) were mixed 1:1 and poured over the blot for 1 min. Excess reagents were removed. Blots were analysed using FluorS Multi-imager and Quantity One software (Biorad).

Solution A
100 mM glycine, pH 10 (with NaOH)
0.4 mM luminol
8 mM 4-iodophenol

Solution B
0.12 % (w/w) H₂O₂ in SDW
2.3.6.2.2. Light intensity evaluation of western blots (ECL detection)

Light intensity was evaluated on images taken with the Fluor-S Multiimager system using Quantity One software. First, a rectangular volume was created around the area of interest and then a volume analysis report was requested for this area. Results came out as Volume in units counts per square millimeter (CNT*mm²) or Adjusted Volume (CNT*mm²). Adjusted volume refers to volume minus background volume.

2.3.6.2.3. Alkaline Phosphatase detection

66 µl NBT (nitro blue tetrazolium) and 33 µl BCIP (5-bromo-4-chloro-3-indolyl-phosphate) was added to 9.9 ml alkaline phosphatase detection buffer. The solution was poured over the blot and left in darkness until a colour change was observed where the Alkaline phosphatase-conjugated secondary antibody was present.

**Alkaline phosphatase detection buffer:**

- 100 mM NaCl
- 5 mM MgCl2
- 100 mM Tris
- pH 9.5

2.3.6.3. Modified western blot for detection of membrane proteins

Since the proteins are likely to aggregate during boiling, the protocol for protein loading on the SDS-PAGE gel was modified so the sample was not boiled prior to loading on the gel. Omitting the boiling step, SDS-PAGE was undertaken as described in section 2.3.5.1. Protein transfer in western blot was done as described in section 2.3.6.1. The modification
was at the end of the transfer when the blot was incubated in a stripping buffer for 15 min at 55 °C with mild agitation (Kaur and Bachhawat, 2009). The stripping buffer was washed from the blot with several washes with 1x TBST at RT until the β-mercaptoethanol smell was no longer detected. Then, the membrane was placed in a blocking solution at 4 °C overnight and immunodetection was undertaken as described in section 2.3.6.2.

**Stripping buffer:**
100 mM β-mercaptoethanol
2 % SDS
62.5 mM Tris-HCl pH 6.7

### 2.3.7. Slot-blot binding assay

Four overlapping 15-mer peptides, spanning the predicted 35 amino acid extracellular domain of PrpS₁ and a scrambled peptide were synthesized by Alta Bioscience (University of Birmingham, UK). These peptides were bound to a polyvinyl difluoride (PVDF) membrane using a slot-blotter. The membrane was properly labelled and pre-wetted with methanol for 10 s. Then, it was incubated with MiliQ water for 10 min on a shaker and finally incubated in PBS for 10 min with constant gentle shaking. The slot-blot filtration manifold was attached to a vacuum pump and after the insertion of the membrane, the slot-blot system was assembled and the vacuum pump was turned on. Different amounts of peptides (10 μg, 1.0 μg and 0.1 μg) in the final volume of 50 μL were bound to the membrane. The scrambled peptide control was used at the same concentrations, while the recombinant PrsS at concentration 0.1 μg. The membrane carrying the PrpS peptides was immediately rinsed with PBS and then incubated with 5 % skimmed milk in PBS for 1 h at room temperature. The membrane was then incubated with 20 μg mL⁻¹ recombinant PrsS₁
protein previously dialysed, as described above, against PBS at room temperature for 16 h to allow binding. Then, membrane was washed with PBS and blocked with 5 % milk in TBST for 1 h at room temperature. The membrane was then incubated for 2 h with the primary antibody raised against PrsS\textsubscript{1} to detect S protein binding to PrpS. Following incubation, the membrane was washed with 5 % skimmed milk in TBST and then incubated with anti-rabbit antibody conjugated with alkaline phosphatase. Detection was done with NBT/BCIP in alkaline phosphatase detection buffer as described above.

**Peptides:**

PrpS\textsubscript{1} A1 - VKLLGLVLHRLSFSE

PrpS\textsubscript{1} A2 - LHRLSFSEDQKWVA

PrpS\textsubscript{1} A3 - DQKWVAFGTAAICD

PrpS\textsubscript{1} A4 - TAAICDVLPLLVPKNML

PrpS\textsubscript{1} A1 scrambled 1 - ELGVKLHSLSLRFL

PrpS\textsubscript{1} A3 scrambled 1 - FTVDVKDCAAAAWGQI

PrpS\textsubscript{8} A1 - LKLLGWVLQHVTVE

PrpS\textsubscript{8} A1 scrambled 1 - GLTWLQLKEVHTVL

**Phosphate Buffer Saline (PBS):**

137 mM NaCl
2.7 mM KCl
10 mM Na\textsubscript{2}HPO\textsubscript{4}
2 mM KH\textsubscript{2}PO\textsubscript{4}
Primary antibodies:
Polyclonal anti-PrsS1; titre 1:6000
Polyclonal anti-PrsS3; titre 1:5000
Polyclonal anti-PrsS8; titre 1:4000

Secondary antibody:
Anti-rabbit AP; titre 1:5000

Prior to use, both antibodies were cleaned up using an Immobilised \textit{E. coli} lysate kit, Pierce, Rockford (see next chapter 2.3.8).

\textbf{2.3.8. Purification of antibodies using Immobilised \textit{E.coli} lysate kit}

Possible non-specific antibodies contained in the primary antibody preparations were eliminated by chromatography using a column with immobilised \textit{E. coli} total proteins. The antibody solution was passed through an \textit{E. coli} protein column, previously equilibrated with 10 mL TBS buffer (BupH\textsuperscript{TM} Tris buffered saline dissolved in 500 mL of SDW). After adding a 1 mL crude antibody solution to the column, an extra 100 µL TBS was added and fractions were collected in a 1.5 mL microtube. Protein concentration of all fractions was determined through Bradford assay (Bradford, 1976). Fractions with the highest protein concentration were pooled. Finally, the column was washed with at least 10 mL regeneration buffer, followed by 10 mL TBS plus 0.02 % sodium azide for storage.
2.3.9. Pollen tube length measurement

Pollen tubes were grown and given the appropriate treatment and mounted on glass slides. Pollen was observed using a Nikon Eclipse Ti:300 microscope attached to a cooled coupled device (CCD) camera supplied by Applied Imaging, UK. Capture and analysis of images was achieved with a Nikon NIS elements 3.2 image analysis system. Images were saved as jp2 files. Tube lengths were measured (50 tubes per treatment; \( N = 3 \)) with the NIS elements 3.2 software. Measurements were exported into a Microsoft Excel file where statistical analysis was performed.

2.3.10. Programmed cell death – caspase-like activity assay

To determine the presence of caspase-like activity, we used the fluorogenic caspase-3 substrate Ac-DEVD-AMC (Calbiochem). Pollen (Papaver or Arabidopsis) was hydrated and grown as described in sections 2.1.4 and 2.2.3, respectively. After ~1 h of growth, pollen tubes were treated as indicated in each experiment. Pollen was left in GM containing the indicated treatment for 5 h at room temperature as this period gives the highest caspase activity if PCD was occurring (Bosch and Franklin-Tong, 2007). The pollen was then collected into 1.5 ml microtubes and the protein extracted as described in section 2.3.3 and the protein concentration was determined as described in section 2.3.4. The protein concentration was adjusted to 1 \( \mu g \mu L^{-1} \) by diluting the extract in the sodium acetate extraction buffer (pH 6) and protein concentration was measured again. Samples with 10 \( \mu g \) were extracted with pH 6 caspase extraction buffer and were loaded into 96 well plate in duplicates, incubated with 50 \( \mu M \) Ac-DEVD-AMC (Calbiochem),
fluorogenic caspase-3 substrate in caspase buffer and adjusted to pH 5, since this is the optimum pH for caspase action in poppy pollen (Bosch and Franklin-Tong, 2007). If protein extracts were less than 1 µg µL⁻¹, the appropriate volume of protein extract was added into the well of the 96 well plate to make 10 µg of protein. The other samples in that experiment were made up to the greatest volume by adding the appropriate amount of pH 6.0 sodium acetate buffer so that all wells had the same volume of pH 6 buffer and protein. Fluorescence was monitored at 460 nm using a time-resolved fluorescence plate reader (FLUOstar OPTIMA; BMG LABTECH) every 15 min over a time period of 5 h. The caspase activity for each sample was calculated by subtracting the fluorescence reading of the first cycle from the final (21st) cycle reading. Results are presented as percentage caspase activity relative to the DMSO or GM control.

2.3.11 Cell Death
2.3.11.1. Cell Death – Viability test

Poppy pollen tubes were grown and treated with appropriate drugs, then incubated with 10 µg fluorescein diacetate (FDA) per mL of liquid GM for 5 min at room temperature and mounted on glass slides. Pollen was visualized under a Nikon Eclipse Ti:300 microscope attached to a cooled coupled device (CCD) camera supplied by Applied Imaging, UK. Capture and analysis of images was achieved with a Nikon NIS elements 3.2 image analysis system. Images were saved as jp2 and tif files. Pollen tubes were counted (30 tubes per treatment; N=3) in the following categories: (1) total, (2) green (alive) and (3) unstained (dead).
Arabidopsis pollen viability assay was based on using Evans blue dye. FDA was not an appropriate stain as the pollen was expressing GFP and the wavelengths overlapped. Pollen was stained with 0.05 % Evans blue for 15 minutes. After incubation, the sample was washed four times with SDW in order to remove excess dye. Aliquots of 20 µL were mounted on microscope slides and pollen was visualised using brightfield microscopy (Nikon Eclipse Tε300). Scoring categories were: (1) total, (2) unstained (alive) and (3) dark stained (dead). Each experiment was done in triplicate with 180 pollen counted each time. Statistical analysis by Chi square was performed in Microsoft Excel.

2.3.11.2. Pretreatment with tetrapeptide inhibitor Ac-DEVD-CHO

*Arabidopsis thaliana* pollen was pretreated with 100 µM Ac-DEVD-CHO for 1 h before SI was induced by adding stigmatic PrsS₁ or PrsS₃ recombinant proteins to PrpS₁-GFP and PrpS₃-GFP expressing pollen growing in vitro, respectively, at final concentration 20 µg/mL. Negative control was comprised of pollen, incubated in Ac-DEVD-CHO only, untreated pollen, grown in AtGM only or pollen with induced SI reaction without DEVDase inhibitor. Pollen was incubated for 8 h and then stained as described in previous section 2.3.11.1.

2.3.12. Actin labelling

Arabidopsis pollen was collected and germinated as described in section 2.2.3. Samples were fixed in a 1.5 microtube with 124 µg mL⁻¹ m-Maleimidobenzoyl-N-hydroxysuccinimide ester (MBS) for 6 min and then paraformaldehyde (PFA) was added to a final concentration of 2 % and the samples were incubated for 1.5 h at 4 °C. After fixation, PFA was eliminated.
with 3 washes of 1 mL TBS, removing the supernatant after each wash by centrifuging at 3600 rpm for 2 min. After the last wash, samples were incubated with TBS plus 0.1 % Triton for 40 min to permeabilise the pollen tubes. Tubes were stained with 66 nM Rhodamine-Phalloidin and samples were left for 16 h at 4 °C or 2 h at room temperature. Then, 10 μL of sample was pipetted onto a microscope slide and 5 μL of Vectashield with DAPI was added before slides were covered with a cover slip and sealed with transparent nail polish. Samples were visualised using epifluorescence or confocal microscopy. Epifluorescence was performed on a Nikon Tε300 fluorescence microscope. The slides were searched for intact pollen tubes. The nuclei were then visualised using the DAPI filter before switching to a TRITC filter to see if there was any Rhodamine-Phalloidin labelling of F-actin. Capture and analysis of images was achieved with a Nikon NIS elements 3.2 image analysis system. Images were saved as jp2 and TIF files and then manipulated using Image J and Adobe Photoshop.

Confocal laser scanning microscopy was performed by Katie Wilkins on Leica SP2 Inverted Confocal system. Full z-series sections of the pollen tubes were taken, 0.5 μm thick. GFP images were taken using Argon 488 nm laser, using 5 % of the power, emission bandwidth was 500 – 530 nm using 550.1 gain. Rh-Ph images were taken using 543 nm laser, using 100 % power, emission bandwith was 565 – 600 nm, using 550.1 gain.

Images from the confocal were saved as tiff files then viewed and manipulated using ImageJ software.

**Scanning settings:**
- 4x average scan
- 400 hz scan
- 512 x 512 pixel frame
- 63x oil immersion lens
2.3.13. Leaf mesophyll protoplast production

Leaves from 3-4 weeks old Arabidopsis plants were cut with a surgical knife into very thin strips on a clean paper surface and transferred to the Petri dishes containing 15-20 mL of enzyme solution. Leaves were incubated in a Petri dish sealed with Parafilm and wrapped in aluminium foil at 22 °C for 16 h. Care was taken to keep the Petri dish as still as possible not moving or shaking it at any point. Then, the enzyme solution was pipetted with a sterile Pasteur pipette very carefully to avoid disturbing the leaves, otherwise the protoplasts would be released too early. Protoplasts were released by adding 10 mL K3 media on the plate, which was then gently swirled. Protoplast containing medium was passed through a nylon filter in a sterile 20 mL plastic tube with a round bottom. The tube was left to stand still for 40 – 60 min and during this time the protoplasts floated to the top of the solution. The upper layer with protoplasts was transferred into a new round bottom tube using sterile Pasteur pipette. Then, protoplasts were washed with 5 mL K3 medium, which was carefully pipetted into the tube. The tube was left to stand again for ~1 h to allow protoplasts to float again. At this stage, small aliquot was taken to assess protoplasts viability using FDA, while the main part of protoplasts were transfected using 5 – 20 µg of plasmid pEarleyGate purified by Midiprep kit (Qiagen). 250 µL of suspended protoplast solution was added per tube with DNA and an equal volume of PEG solution was added to the mixture of DNA and protoplasts and incubated for 30 min at RT. The PEG solution was diluted by adding 2 mL K3, and tubes allowed to stand for 1 – 2 h to let the protoplasts float to the top. The solution underneath the protoplasts was carefully withdrawn and a new aliquot of 2 mL K3 medium was added to the protoplasts and left to incubate for 16 h in the dark at RT. The following day transient expression of the gene of interest was verified.
K3 medium: Prepared from following stock solutions
10x B5 medium with vitamins
200x MES (0.1 g/mL)
500 x myo-inositol (0.05 g/mL)
100x NH₄NO₃ (25 mg/mL)
100x CaCl₂*2H₂O (75 mg/mL)
100x D-xylose (25 mg/mL)
0.4 M sucrose
pH adjusted to 5.6 – 5.7 using 1.0 M KOH
Solution was filter sterilized and stored at 4 °C.

“Enzyme solution”:
0.5 % cellulase (R-10)
0.2 % macerozyme (R-10)
prepared in K3 medium, filter sterilized
and stored at -20 °C.

Protoplasts suspension solution medium
0.4 M Mannitol
20 mM CaCl₂*2H₂O
5 mM MES pH 5.7
filter sterilized and stored at 4 °C.

PEG solution
40 % PEG 4000 (Fluka)
0.4 M mannitol
100 mM Ca(NO₃)₂
warmed in the 60 °C water bath
pH 7.0 adjusted with 1.0 M KOH
Filter sterilized and aliquoted, stored at -20 °C

PEG solution
40 % PEG 4000 (Fluka)
0.4 M mannitol
100 mM Ca(NO₃)₂
warmed in the 60 °C water bath
pH 7.0 adjusted with 1.0 M KOH
Filter sterilized and aliquoted, stored at -20 °C

2.3.14. PCR screening for the presence of inserts in transgenic plants

Leaf disks were obtained from Arabidopsis transgenic plants and untransformed Col-0 as a
c control in small PCR microtubes. To extract DNA, 50 μL of an extraction buffer was added
and the leaf disk was grinded using a sterile filter tip, incubated at 95 °C for 10 min and
cooled on ice for 2 min. After that, 50 μL of a dilution buffer was added, and the mixture
was vortexed and centrifuged (1 min, 13200 rpm). For each PCR reaction 1 μL of DNA was
used. The presence of inserts was anlysed by electrophoresis on a 0.8 % agarose gel.

Oligonucleotide primers were supplied by MWG-Biotech.

Extraction Buffer
0.1M Tris-HCl, pH 9.5
0.25 M KCl
0.01 M EDTA
Solutions were prepared by aseptic work and filter sterilized before use.

Dilution Buffer
3 % BSA
PCR mix:
12.5 μL ReddyMix™ PCR Master Mix (Thermo Scientific)
1 μL forward primer
1 μL reverse primer
1 μL DNA
9.5 μL dH₂O

PCR conditions:
93 °C – 3 min  1x
93 °C – 1 min
57 °C – 1 min  35x
72 °C – 1 min
72 °C – 10 min  1x

Used primers:
Forward PrsS₁: 5’-GGAGCATTGGCATCCATTGCCG-3’ Tm = 64 °C
Reverse PrsS₁: 5’-CCATTATCTTCCAGAGGCACTGGG-3’ Tm = 64.4 °C
Forward PrsS₃: 5’-CGATCCACTGCCAATCAGAAGACG-3’ Tm = 64.4 °C
Reverse PrsS₃: 5’-GTGGAGCACCTTCCGCGTCG-3’ Tm = 67.6 °C
Forward PrsS₈: 5’-GGTAATGGCCATAGCATCGGG-3’ Tm = 61.8 °C
Reverse PrsS₈: 5’-CATCCGTCTGTCTTCCACAGGC-3’ Tm = 61.8 °C
Forward clon. PrpS₁: 5’-CCATGCCCCGAAGTGGAAGTGTTG-3’ Tm = 66.1 °C
Reverse clon. PrpS₁: 5’-CCTTAAGCTTGAGTTATAAGATGAGGGGAATCC-3’ Tm = 67.0 °C
Forward PrpS₃: 5’-CCATGCTTTACGTGGAAAGACC-3’ Tm = 62.4 °C
Reverse PrpS₃: 5’-GGCTGCAGAAGTGGCTTCATC-3’ Tm = 61.8 °C
Forward PrpS₈: 5’-CCCTATTTGGATCCGCACTTGCC-3’ Tm = 64.2 °C
Reverse PrpS₈: 5’-GAGGATTCAGAGGAGTTGCC-3’ Tm = 64.0 °C
2.3.15 DNA analysis by agarose gel electrophoresis

For the gel, agarose was fused in 0.5X TBE by heating the solution until it was completely clear. Then, 0.5 µg mL\(^{-1}\) ethidium bromide was added to the molten agarose. Gels were poured and electrophoresed using Hybaid or Biorad electrophoresis kits. Images of gels were captured using a FluorS Multi-imager and analysed using Quantity One software if necessary. An estimation of size was achieved by parallel electrophoresis of an aliquot of 1 kb ladder (Invitrogen).

**10 x TBE:**
- 0.9 M Tris
- 0.9 M Orthoboric acid
- 25 mM EDTA

**DNA loading buffer:**
- 40 % (v/v) glycerol
- 0.25 % (w/v) bromophenol blue

2.3.16 Semi-quantitative RT-PCR

2.3.16.1 RNA extraction

Total RNA was extracted using RNeasy Mini Kit (Qiagen). All materials and water was pretreated with 0.05 % or 0.1 % diethylpyrocarbonate (DEPC), respectively. DEPC reacts with histidine residues on the proteins and thus it inactivates RNases. However, it can react with RNA as well. Therefore, it was necessary to autoclave DEPC incubated solutions before the experiment. DEPC is heat-sensitive and is degraded by autoclaving.

For each extraction, 20 *A. thaliana* (transgenic or untransformed) flowers were collected. Flowers were frozen in liquid nitrogen and grinded thoroughly in microtubes using a plastic rod. β-mercaptoethanol was diluted 1:100 in RLT buffer and 450µl of the solution added to the grounded flowers and mixed by vortexing. The sample was then added to a lilac QIAshredder column spin column and centrifuged (2 min, 13200 rpm). The flow-through
was transferred to a new microtube, 0.5 V of 100 % ethanol was added, the solution mixed by pipetting, then immediately transferred to a pink RNeasy column and centrifuged (15 s, 13200 rpm). The flow-through was discarded and 700 µL RWI buffer added to the new column before centrifugation (15 s, 13,200 rpm). The flow-through was discarded and the column transferred to a new collection tube. 500 µL RPE buffer was added to the column, centrifuged (15 s, 13,200 rpm) and the flow-through discarded. Another 500 µL of RPE buffer was added to the column again and centrifuged (2 min, 10000 rpm) to ensure the membrane was dry. The column was transferred into a new microtube and centrifuged (1 min, 13200 rpm). Finally, the column was transferred to a final microtube, 30 µL of RNase free water added and centrifuged (1 min, 13200 rpm). Then, another 10 µL of RNase free water was added to completely elute everything and centrifuged (1 min, 13200 rpm). 5 µL of RNA was used to analyse by electrophoresis and the rest stored at -80 °C until required.

2.3.16.2. RNA analysis by agarose gel electrophoresis

Analysis of RNA samples was accomplished by electrophoresis on agarose gels using 5 µL of RNA loading dye. No ladder was used due to the possible contamination with RNases. 1 % agarose gel was made up in 0.5x TBE using RNase free SDW and poured in RNase free gel tank. Electrophoresis settings and gel visualisation are described in chapter 2.3.15.1. RNA was stained with ethidium bromide, which was previously added to the agarose solution at a final concentration of 0.5 µg mL⁻¹.
2.3.16.3. DNase treatment of RNA

In order to remove contaminating DNA, 10 μL of RNA was mixed with 10 μL of DNase solution and incubated at RT for 15 min. 2 mM EDTA was added and the mixture was incubated at 65 °C for 10 min. Then, 80 μL of RNase free water was added and phenol extraction proceeded by addition of equal volume of phenol. The solution was mixed and centrifuged (5 min, 13200 rpm). The upper aqueous layer was removed and transferred to a clean tube and an equal volume of chloroform was added. Again, the sample was vortexed and centrifuged (5 min, 13200 rpm). The top layer with aqueous phase was removed, transferred to the fresh tube and 1 μL of glycogen (Roche) was added, which serves as a carrier to promote nucleic acids precipitation. RNA was precipitated by adding 2.5 V of 70 % ethanol, vortexed well, incubated at 70 °C for 30 min and centrifuged (10 min, 13200 rpm). Supernatant was carefully removed. RNA pellet was washed with 70 % ethanol. The pellet was dried in a vacuum chamber for 15 min and resuspended in 20 μL of RNase free water.

**DNase solution:**
- 2 μL 10x DNase I buffer
- 1 μL recombinant RNasin (Promega)
- 1 μL DNase (Invitrogen)
- 6 μL RNase free SDW

2.3.16.4. cDNA synthesis by RT-PCR

Isolated total RNA was used in RT-PCR reaction One-step RT-PCR kit (Qiagen). cDNA was analysed by electrophoresis as described in section 2.3.15.1. Sample loading was
compared to glyceraldehyde-3-phosphate dehydrogenase C (GAPC), whose cDNA was also amplified and used as RT-PCR standard and loaded in parallel on the same gel.

**RT-PCR mix:**
2 μL RNA
10 μL 5x buffer
2 μL dNTP
1 μL forward primer
1 μL reverse primer
1 μL recombinant RNasein
2 μL enzyme mix
31 μL RNase free dH2O

**RT-PCR conditions:**
50 °C – 30 min   1x
95 °C – 15 min   1x
94 °C – 1 min
57 °C – 1 min
72 °C – 1.5 min   30x
72 °C – 10 min   1x

**Primers:**
GAPC forw: 5’-CACTGACAAAGACAAGGCTGCAGC-3’  Tm = 64.4 °C
GAPC rev: 5’-CCTGTGTGTGCTGCAAGTACAG-3’  Tm = 64.2 °C
CHAPTER 3

Investigation of the possible involvement of the ubiquitin-proteasomal pathway during SI in *Papaver* pollen
3.1. INTRODUCTION

As mentioned in the introduction (section 1.6), the Ubiquitin-mediated 26S proteasomal (UbP) degradation of proteins is required to maintain the cellular homeostasis and regulation of various cellular functions. Protein substrates are covalently attached to a small protein, ubiquitin, by the sequential action of three enzymes. Ubiquitin is first activated by an ubiquitin-activating enzyme E1 and transferred to the ubiquitin-conjugating enzyme E2. E2 binds to the ubiquitin-protein ligase E3 that catalyzes the formation of a covalent bond of ubiquitin molecules sequentially attached through Lys-48 of the previous ubiquitin (Hershko and Ciechanover, 1998). A chain of at least four ubiquitins is required for degradation by the 26S proteasome. Monoubiquitinated proteins or proteins tagged with the ubiquitin chain through the Lys-63 are not targeted for degradation but they are involved in various events such as subcellular localization or protein activation or protein-protein interaction (Dielen et al., 2010).

The specificity of the UbP pathway is mainly determined by the wide range of diverse E3 ligases that recognise substrates through the specific protein recognition domains (see introduction for details). In A. thaliana there are more than 1400 genes that encode different components of the UbP pathway and among them around 1300 encode different predicted E3 ligases (Smalle and Vierstra, 2004). There are four major groups of E3 ligases: monomeric HECT-domain E3 ligases and three groups that contain RING domain; monomeric RING/U-box E3 ligases and complex Cullin RING E3 ligases and APC E3 ligases. The Cullin RING ligases are further divided based on the different subunits of Cullin that they contain: the SCF complex contains SKP1, Cullin-1 protein F-box for protein specificity and RBX1; the complex with Cullin-3 subunit contains RBX1 and BTB1/POZ.
domain, and Cullin-4 based complex contains RBX1, DDB1 and WD (Vierstra, 2003, Vierstra, 2009).

The 26S proteasome is a large cylindrical multisubunit complex, consisting of the 20S central core domain, that is comprised of four stacked rings of $\alpha$ and $\beta$ subunits (each ring is composed of seven proteins), and the 19S regulatory “cap” structure, that can be divided into lid and base components. The 19S regulatory particle recognizes polyubiquitin tags attached to the target protein substrates, initiates the degradation process in the core and also removes the Ub chains (Voges et al., 1999, Kurepa and Smalle, 2008).

UbP degradation was demonstrated as playing a role in plant-pathogen interactions and some aspects of plant reproduction. The UbP pathway is involved in plant-pathogen interaction at several different levels. Plant-pathogen interactions start with the perception of the PAMPs, which are mediated by plasma membrane located pattern recognition receptors. This is the first step towards PAMP-triggered immunity, achieved through the cascade of downstream events (reviewed in Trujillo and Shirasu, 2010). The ubiquitination pathway, but not the 26S proteasome, is involved in the basal host resistance of barley epidermis that is attacked by powdery mildew (Dong et al., 2006). Plant pathogens developed several strategies by which they interfered with the plant UbP pathway. They can recruit E3 ligases and target crucial defense plant proteins for degradation by the 26S proteasome. For example $P$.syringae AvrPtoB acquired E3 ligase activity through which it conjugates the Ub chain on plant Fen or FLS2 proteins that stand guard against pathogen invasion, targets them for proteasomal degradation, and thus creates the way to infect the plant (Göhre et al., 2008, Rosebrock et al., 2007). Pathogens also developed a way to inhibit plant 26S proteasomal degradation, for example the $P$. syringae protein SylA inhibits the $\beta$ 1/2/5 subunits of the core particle of 26S proteasome (Groll et al., 2008).
Yang et al., (2006) reported that the E3 ligase, PUB17 from *A. thaliana* and its functional tobacco homologue ACRE276 act as positive regulators of cell death and defense responses in Solanaceae and Brassicaceae. In addition, (Sadanandom et al., 2008) reported the identification of the BTB/POZ domain transcriptional repressor, AtPOB1, that interacts with the ARM repeat of PUB17 to control disease resistance in plants. Plants that were deficient in POB1 produced spontaneous cell death, demonstrating its critical role in plant cell death.

Another important involvement of the 26S proteasome in hypersensitive response was demonstrated by (Hatsugai et al., 2009). They demonstrated a direct link between proteasome activity and plant PCD by identifying that the proteasomal subunit PBA1 acts as a long sought plant caspase-3-like enzyme. Proteasome and its caspase-3-like activity are required during bacterial infection to degrade a yet unknown negative regulator of membrane fusion, thus enabling the fusion of the plant plasma membrane and the vacuole membrane. In this way pores are created that enable the discharge of vacuole content and defence proteins outside the cell, inhibiting bacteria infection and causing HR cell death.

Ubiquitination and proteasomal degradation were also demonstrated as playing a role in some aspects of plant reproduction: free ubiquitin is required for pollen tube adhesion and guidance in lily (Kim et al., 2006) and proteasomal degradation is necessary for normal pollen tube development in *Picea* (Sheng et al., 2006) and kiwi (Speranza et al., 2001). Moreover, ubiquitination and proteasomal degradation are essential during S-RNase based gametophytic SI in Solanaceae, Rosaceae and Plantaginaceae, and in sporophytic SI in Brassicaceae (for recent reviews see Meng et al., (2011), Tantikanjana et al., (2010), Yee and Goring, (2009)). During an S-RNase-based incompatible response, self and non-self S-RNases enter pollen tube. The hypothesis whose *S*-specificity was not yet demonstrated in all GSI species is that pollen determinants recognize the non-self S-RNases and target them
for the proteasomal degradation while self S-RNases are not recognized and are allowed to degrade the RNA of an incompatible pollen, therefore preventing self-fertilization (Kao and Tsukamoto, 2004, Meng et al., 2011, Liu et al., 2009, Qiao et al., 2004b, Hua and Kao, 2006). In the S-RNase based SI system the pollen S-determinant is an S-locus F-box protein (SLF or SFB), a component of a SKP1-Cullin1-F-box (SCF) complex (Sijacic et al., 2004, Lai et al., 2002, Entani et al., 2003, Ushijima et al., 2003). In *Antirrhinum*, the SLF is a conventional SCF complex with a novel SKP1-like protein SSK1 (Huang et al., 2006), while in *Petunia* the SLF is part of a novel E3 ligase complex. The other components of the PiSLF complex in *Petunia* are PiCUL1-G and PiSBP1 (*P. inflata* S-RNase Binding Protein1), a RING-HC protein in addition to PiSLF (Hua and Kao, 2006). However, the novel E3 ligase identified does not contain SKP1 or RBX1.

The sporophytic SI mechanism found in Brassicaceae also involves components of UbP pathway. The ARC1 protein, an E3 ligase that is a member of PUB family of proteins, acts as a regulator of the *Brassica* SI response in the stigmatic papillae (Stone et al., 2003). ARC1 mediates the ubiquitination of EXO70A1, a putative component of an exocyst system, which regulates secretion (Samuel et al., 2009). In addition, an ubiquitin specific protease, a homologue of AtUBP22, was identified in rye pistils and its role was implicated in the SI in grasses (Hackauf and Wehling, 2005).

The self-incompatibility (SI) and hypersensitive response (HR) are important examples of signaling pathways triggered in response to incompatible interaction between two cognate proteins, resulting in PCD in plants. The mechanisms of HR and SI in *Papaver* share many similarities, such as: increased levels of [Ca\(^{2+}\)], Ca\(^{2+}\)-dependent phosphorylation of proteins, generation of reactive oxygen species and cessation of cytoplasmic streaming and caspase-3-like induced PCD (Geitmann et al., 2004). Ubiquitin-dependent proteasomal (UbP)
degradation is an important component of PCD during HR and has also been reported to play an important role in other SI systems, like GSI in Petunia and Pyrus, and SSI in Brassica, but has not yet been reported during SI in Papaver.

The majority of studies on the UbP pathway use the inhibitors of the proteasome activity due to their quick entering of cells and rapid reversibility of their action (Kisselev and Goldberg, 2001). The most desirable proteasome inhibitors are those specifically targeting the proteasome without affecting activities of other serine or cysteine proteases. Among the proteasome inhibitors, MG132 represents a potent cell-permeable proteasome inhibitor and is widely used. MG132 is a peptide aldehyde Z-Leu-Leu-Leu-CHO that inhibits the proteasome activity by covalently binding to the active site of the β-subunits in the core particle (Zhang et al., 2009). The specificity of MG132 was demonstrated in numerous studies on different systems, for example using a biochemical approach by tracking degradation of several proteins known to be substrates for proteasome degradation in yeast cells (Liu et al., 2007a). In plants, the use of the proteasome inhibitor MG132 was found to inhibit length of tracheary elements in Zinnia cell cultures and also in A. thaliana (Zhao et al., 2008). (Vacca et al., 2007) reported that MG132 prevented cells from PCD induced with HS in tobacco BY2 cells. In growing pollen tubes, MG132 affects pollen tube morphology (Sheng et al., 2006, Speranza et al., 2001), and mitochondrial dynamics and causes vacuolization in the pollen tube (Sheng et al., 2010).

AIMS

The aim of this study was to investigate whether there is a link between Papaver SI and UbP pathway and if so what role does the UbP degradation play during the pollen SI reaction in
Papaver. This work was carried out in collaboration with Dr. Ari Sadanandom from University of Durham, whose research is focused on the role of ubiquitin-mediated proteolysis during disease resistance in plants (Yang et al., 2006, Sadanandom et al., 2008). We aimed to establish a link between ubiquitination and other PCD events that occur in the pollen tube during SI reaction. This could provide an important information about cell death signalling and might also provide further insights into two mechanistically similar cell responses, SI and HR. In order to investigate the involvement of the UbP pathway during SI-induced PCD pathway in Papaver pollen, the SI response was characterized in the presence of the proteasomal degradation inhibitor MG132. Pollen viability, pollen tube growth inhibition, and the level of protein ubiquitination during the SI reaction were determined.

3.2. RESULTS

3.2.1. The proteasome inhibitor MG132 affects pollen germination and tube growth

In order to investigate whether the UbP pathway has any role in poppy pollen germination and growth, a pharmacological approach using the reversible proteasome inhibitor MG132 was used. Initially we wished to determine the concentration of the MG132 that would be effective without inhibiting pollen tube growth or causing the alterations to pollen tube morphology. The effect on normal pollen tube growth of different concentrations of MG132, in range of 10 to 100 µM (see Figure 3.1), was determined. Different concentrations of MG132 were applied to pre-germinated poppy pollen tubes in vitro. Pollen was left to incubate for 30 min or 3.5 h when the pollen tube lengths of 39 tubes were measured (See
Figure 3.1). Figure 3.1 presents the effect of different concentrations of MG132 on the length of pollen tubes.

The average pollen tube length of the untreated control after 3.5 h incubation was 461 μm and it decreased when pollen was pretreated with MG132 (see Figure 3.1). The difference in pollen tube length between untreated and 10 μM was not significant (P=0.085; nonsignificant), nor was the difference in length significant between pollen that was pretreated with 10 or 40 μM MG132 (P=0.122; nonsignificant). When comparing the pollen tube length of pollen that was pretreated with 40 or 60 μM MG132, the difference was highly significant (P=7.8*10^{-6}; **). When pollen was pretreated with 80 or 100 μM MG132 the pollen tube growth was inhibited significantly (P=1.4*10^{-14} and P=5.3*10^{-19}, ***, respectively), see Figure 3.1. Higher concentrations of MG132 (80 and 100 μM), caused severe pollen tube swelling and balloon-like pollen tube tips. So they were not considered further. Some signs of pollen tube tip swelling were also observed at 60 μM.
MG132 but not at 40 μM MG132. Therefore 40 μM MG132 was chosen as the most effective concentration to inhibit proteasomal degradation without causing major morphological changes in further experiments. This concentration is also consistent with the reports from the literature (Wang et al., 2009b, Sheng et al., 2006, Sheng et al., 2010, Speranza et al., 2001, Liu et al., 2007a). Reports from the other pollen tube systems like *Picea* (Sheng et al., 2006) and kiwi (Speranza et al., 2001) show that MG132 alters pollen tube morphology, causes tube tip swelling, tube branching and germination of more than one tube per grain but the removal of inhibitor restored growth. These effects were not so dramatic in case of poppy pollen germination, however tube swelling and branching was detectable when pollen was pretreated with 80 or 100 μM MG132 concentrations of MG132.

3.2.2. Effect of inhibition of proteasomal activity on pollen tube growth

After establishing the optimal concentration of MG132 for these studies and the effect it had on pollen tube growth, we examined whether the proteasomal activity was involved in poppy SI. Pollen was pretreated with 40 μM MG132 for 30 min and then SI was induced by incubating pollen with recombinant PrsS proteins for 3 h. The length of the pollen tubes was measured to examine whether the inhibition of proteasomal activity by MG132 would affect the inhibition of pollen tube growth caused by the SI response and rescue pollen from SI (see Figure 3.2).
Mean length of *Papaver* pollen tubes pretreated with 40 μM MG132 and then SI induced (MG-SI) was 186 ± 15 μm (n=50 pollen tubes, N=3 repeats), which was not significantly different compared to the *Papaver* pollen with induced SI (172 ± 15 μm; P=0.314; nonsignificant). The required additional control would be length of pollen incubated only with 40 μM MG132 but as it can be seen in Figure 3.1., that 40 μM does not significantly affect length when compared to non-treated. These results suggest that MG132 has no effect on the SI induced inhibition of pollen tube growth (see Figure 3.2). When compared to the mean length 389 ± 30 μm for the untreated (UT) control (n=50 pollen tubes, N=3 repeats), both values were highly significantly different (P=3.13*10^{-22} compared to MG-SI treated pollen tubes and P=8.67*10^{-25} compared to SI treated pollen tubes; ***)}, see Figure 3.2. Taken together, these data demonstrate that inhibition of proteasomal activity cannot rescue the pollen tube growth after the induction of SI.
3.2.3. Effect of inhibition of proteasomal activity on pollen tube viability

Since pollen tube length measurements did not give any indication of a link between SI and proteasome inhibition, we decided to test whether MG132 treatment can rescue the viability of pollen tubes after SI challenge. SI results in cell death, so if ubiquitination and proteasomal degradation were involved in SI, then the inhibition of proteasomal activity by MG132 might prevent pollen cell death and would therefore increase pollen viability compared to SI induced pollen.

Pollen tube viability was assessed using fluorescein diacetate (FDA). FDA penetrates through the cell membrane and inside the cell intracellular esterases cleave off the diacetate group producing the highly fluorescent product fluorescein (Breeuwer et al., 1995). The fluorescein will accumulate in cells which possess an intact membrane, so green fluorescence can be used as a marker of cell viability, whereas cells that do not possess an intact membrane or an active metabolism do not exhibit green fluorescence. Pollen tubes were grown and treated with MG132 and incompatible PrsS-proteins as described in the previous section. Pollen tubes were incubated with 10 µg/mL FDA in liquid GM for 5 min and examined, as described in the materials and methods section (see Figure 3.3).
Untreated pollen tubes had a mean viability of 62 %, which was highly significantly different to the 6.7 % viability of SI treated pollen (P=1.23*10^{-16}; ***). MG132 partly rescues pollen from SI induced cell death. When pollen tubes were pretreated with 40 µM MG132 prior to SI induction, we observed 325 % increase in pollen viability. The difference between SI and MG-SI treated pollen tubes was significantly different (P=5.53*10^{-05}; ***) and so was the difference between the UT control and the MG-SI treated pollen tubes (P=1.37*10^{-06}; ***). The data presented in this section demonstrate that MG132 can partly rescue the SI-induced cell death of Papaver pollen tubes. Because MG132 inhibits proteasomal activity, this suggests that ubiquitination-dependent proteasomal degradation is a component of Papaver SI.

3.2.4. Proteasomal degradation and PCD in incompatible poppy pollen

In order to investigate further whether proteasomal degradation might be functionally involved in SI induced PCD we measured caspase-like activity in incompatible pollen tubes
that were pretreated with the inhibitor of proteasomal activity MG132. SI triggers PCD involving DNA fragmentation and DEVDase activity in incompatible *Papaver* pollen (Thomas and Franklin-Tong, 2004). Caspase-like activities stimulated by SI have been characterised more directly, and in more detail, using AMC-based peptide caspase substrates that act as fluorogenic indicators for caspase activities, by exhibiting fluorescence upon specific protease cleavage (Bosch and Franklin-Tong, 2007). The prediction was that if ubiquitination and proteasomal degradation were involved in SI-induced PCD then the proteasome inhibitor would prevent PCD in incompatible pollen tubes and caspase-like activity would be lower compared to non-treated, SI-induced pollen tubes. We measured the caspase-3-like/DEVDase activity of protein extracts from SI challenged pollen tubes treated in the presence and absence of MG132 using Ac-DEVD-AMC tetrapeptide (see Figure 3.4).

![Figure 3.4: DEVDase activity of pollen tubes pretreated with 40 μM MG132 and induced SI (MG-SI). UT – activity of untreated control of pollen tubes grown in GM only, 40 μM MG – activity of pollen that was incubated with MG132, SI - DEVDase activity of pollen tubes incubated with recombinant stigmatic PrsS proteins. N=3 repeats. Error bars represent ± Standard Deviation of means.](image)

Pollen tubes grown in GM only (UT) or in GM with 40 μM MG132 exhibited very low DEVDase activity (see Figure 3.4). DEVDase activity of the pollen tubes incubated with 40 μM MG132 only was 14 % lower compared to UT control and was not significantly
different (P=0.143; nonsignificant). DEVDase activity increased by 52 % in pollen tubes where the SI response was induced by recombinant stigmatic PrsS proteins, and this increase was significant compared to the UT control (P=0.021, *). Pollen tubes that were pretreated with 40 µM MG132 before the SI was induced exhibited 26 % lower DEVDase activity compared to the SI samples. The inhibition of proteasomal activity affected DEVDase activity (P=0.046; *). Pollen tubes that were pretreated with 40 µM MG132 before the incompatible response was triggered were also significantly different compared to pollen tubes that were incubated with 40 µM MG132 only or with UT control (P=0.034 and P=0.014, respectively; *). These data, presented in Figure 3.4., demonstrate that the inhibition of proteasomal activity by MG132 had a small but significant effect on pollen tubes as it partly inhibited the DEVDase activity. It suggests a potential link between DEVDase activity and proteasomal activity in incompatible *Papaver* pollen tubes, even though the DEVDase activity stimulated by SI was not very high and MG132 did not reduce activity very much.

### 3.2.5. Effect of proteasomal inhibition by MG132 on the ubiquitination levels in incompatible pollen tubes

In the previous sections we provided preliminary evidence that proteasomal degradation might play a role during the SI reaction in poppy pollen tubes. As ubiquitination via Lys48 is tightly connected with the proteasomal degradation we wished to determine whether the levels of ubiquitination might change upon SI in *Papaver* pollen. Proteins were extracted from poppy pollen tubes pretreated with 40 µM MG132 before the induction of SI by
recombinant stigmatic PrsS and controls, and run on a SDS-PAGE, blotted and then probed with α-ubiquitin antibody. Figure 3.5.a shows the pattern of ubiquitination of total protein extracts. Figure 3.5.b shows corresponding relative light intensity evaluation of the blot.

![Figure 3.5.](image)

**Figure 3.5.:** Ubiquitination levels of poppy pollen pretreated with 40 μM MG132 and SI induced (a) Representative image of western blot of MG-SI protein extracts probed with ubiquitin antibody; (b) percentage of average relative light intensity of ubiquitinated protein extracts; N=4, Error bars represent ± St. Error. GM – extract from untreated pollen; 40MG GM – extract from pollen incubated in 40 μM MG132; SI – extract from pollen that had induced SI reaction; 40 MG SI – extract from pollen that was pretreated with 40 μM MG13 before the induction of SI by PrsS proteins.

The ubiquitination signal was measured on the whole band of the western blot. The untreated pollen sample was given value 100 %. When tubes were grown in GM with the proteasome inhibitor MG132 the level of ubiquitination increased by 133 % due to the probable inhibition of proteasomal degradation, so more ubiquitinated proteins appeared to be accumulated in the pollen tubes. When SI was induced, the level of ubiquitination was more intense by 39 % compared to the control UT sample, suggesting that ubiquitination was increased by the SI response. The strongest signal was obtained when samples were pretreated with proteasome inhibitor before inducing SI and despite the increase in ubiquitination by MG132 on its own, this could implicate a possible role for ubiquitin-independent proteasomal degradation during SI responses.
3.2.6. Proteasomal degradation acts upstream or in parallel with caspase-3-like activity

In order to determine whether proteasomal degradation of proteins acts upstream, downstream or together with caspase-like activity, MG132 was added to the protein extracts and DEVDase activity was measured. If proteasomal degradation acts downstream of DEVDases, then it would be expected that there would be no changes in the level of DEVDase activity, but if it acts upstream or in cooperation with caspase-3-like enzymes, then we would expect to see an increase or decrease in DEVDase activity.

Proteins of pollen tubes treated with incompatible PrsS-proteins and control pollen tubes were extracted as described in the materials and methods section. In these extracts, DEVDase activity was already stimulated by SI before the extracts were mixed with 40 µM MG132 and the control sample was mixed with DMSO, which was the solvent for MG132 (see Figure 3.6).

Figure 3.6.: Caspase activity of pollen tubes with induced SI and added 40 µM MG132 in the pollen tube extract (MG-SI). SI presents the DEVDase activity of pollen incubated with the recombinant stigmatic PrsS proteins; SI-DMSO presents activity of pollen that had added DMSO in the SI pollen tube extract; N=3 repeats. Error bars represent ± Standard Deviation of means.
As can be seen from Figure 3.6, the DEVDase activity of SI extracts with added inhibitor of proteasomal activity MG132 was significantly decreased by 15 % compared to SI induced DEVDase activity (P=0.026; *). The DMSO solvent had no effect on the DEVDase activity when added to the SI challenged pollen protein extract (P=0.963, NS), indicating that the proteasome inhibitor MG132 reduces the DEVDase activity.

These data suggest that inhibition of proteasomal activity by MG132 in incompatible Papaver pollen tubes has a small but significant effect on activated caspases and that it acts upstream or alongside of caspase-3 like activation.

3.2.7. Caspase-3 inhibitor decreases ubiquitination

In order to make a firmer connection between PCD in poppy SI and ubiquitination, an experiment using the caspase-3 inhibitor, Ac-DEVD-CHO, was performed. Pollen tubes were pretreated for 1 h with Ac-DEVD-CHO and then challenged with incompatible stigmatic recombinant PrsS-proteins for 3 different time points: 1 h, 2 h and 4 h. Total protein extracts were separated by SDS-PAGE gel, and blotted and probed with an anti-ubiquitin antibody (see Figure 3.7). Figure 3.7.a shows the ubiquitinated protein pattern in pollen tube samples with induced SI at three different time-points. The ubiquitination levels were higher with SI samples compared to samples that were pretreated with the inhibitor of caspase-3-like activity, Ac-DEVD-CHO and then SI challenged.

The relative signal intensity of the ubiquitination pattern was measured using Quantity One software (see Figure 3.7.b). Ubiquitination in the Papaver pollen that was incubated with the incompatible stigmatic PrsS recombinant proteins increases with the time of SI challenge.
When caspase-3-like activity was inhibited, there was ~20 % less ubiquitination of proteins, however, with the increasing time of SI challenge, the ubiquitination levels increased as well.

![Western blot](image1)

**Figure 3.7.:** Protein ubiquitination levels in poppy pollen tubes pretreated with an inhibitor of caspase activity (Ac-DEVD-CHO) and SI challenged at three different time points: 1 h, 2 h and 4 h. Untreated pollen tubes germinated in GM are a negative control at t0 and 4 h (blue outline) (a) western blot probed with ubiquitin antibody, tubulin presents a control for equal loading; (b) relative light intensity of ubiquitination levels, blue outline presents untreated control, red outline presents samples with induced SI and yellow outline presents samples pretreated with Ac-DEVD-CHO and then SI induced. (N=1)

The experiment was carried out only once, so a firmer conclusion would require more repeats. However, taken together these data indicate that during SI in incompatible *Papaver* pollen tube increased ubiquitination of yet unknown protein targets occur, which are probably targeted for degradation by the 26S proteasome. This SI-specific ubiquitination of
proteins is connected with the caspase-3-like activity, which, based on previous results, acts upstream or in parallel with the UbP pathway in the incompatible *Papaver* pollen tube.

### 3.2.8. Involvement of an E3 ligases in *Papaver* SI

The E3 ligase AtPUB17 and its tobacco homologue NtACRE276 were demonstrated to play a functional role in the HR cell death in tobacco and *A. thaliana*, mediated by the resistance proteins (Yang et al., 2006). In a screen for interactors of the ARM domain of PUB17, (Sadanandom et al., 2008) isolated the POB1 protein. AtPOB1 belongs to the family of BTB/POZ transcriptional repressors and is localised in the nucleus during HR. Dr Ari Sadanandom (University of Durham) provided antibodies against PUB17 and POB1 proteins. We wished to investigate further the involvement of UbP pathway in the SI in *Papaver*. If this was the case, there might be some functional homologues of PUB17 and POB1 activated in the *Papaver* pollen tube system, recognized by the antibodies. If the proteins could be identified, we could investigate further their possible involvement in the SI in *Papaver*.

While anti-PUB17 antibody (PUB17 with Mw 83 kDa) did not recognise any specific protein (Figure 3.8.a), anti-POB1 antibody (POB1 with Mw 65 kDa) recognised distinct proteins in poppy pollen tube extracts (Figure 3.8.b).
Figure 3.8: Protein expression of putative homologues of AtPUB17 and AtPOB1 E3 ligases during SI in *Papaver* (a) AtPUB17 antibody did not recognise any specific protein homologue at 83 kDa during SI in *Papaver*. (b) AtPOB1 (Mw 65 kDa) recognised a putative homologue at 63 kDa. Another two protein bands at 37 and 35 kDa were recognised by AtPOB1. Decrease of signal with time of SI suggests that SI triggered their degradation. Tubulin was used for equal loading.

PUB17 antibody resulted in a smear, with some weaker bands visible at low molecular weights. Signal intensity of the poppy pollen extract detected with AtPUB17 decreased with the time of SI incubation, but this was also the case with untreated pollen. However, when testing whether the AtPOB1 antibody could recognise an equivalent protein during SI in *Papaver* pollen, we observed a distinct band at 63 kDa and two lower ones at 37 and 35 kDa. Whereas the higher molecular weight band probably corresponds to the POB1 protein (Mw 65 kDa), the lower bands might be degradation products (Figure 3.8.b), suggesting that SI triggers its degradation.

*A. thaliana* AtPOB1 plants were resistant to virulent bacteria, they accumulated ROS and exhibited HR cell death (*A. Sadanandom, personal communication*). AtPOB1 represses the expression of genes that regulate defence against pathogen *Pseudomonas* (Sadanandom et al., 2008). If drawing a parallel between HR and SI systems, AtPOB1 homologues recognised by the POB1 antibody could function by repressing genes required for the SI
response. Upon SI, putative poppy POB1 homologues may be targeted for degradation, gene expression is activated and functional SI can occur. Interestingly, AtPOB1 proteins are localised in the nucleus (Sadanandom et al., 2008). Their localisation has not yet been examined in the Papaver pollen, however if it is nuclear as well, this could explain their slow degradation, as it is known that DEVDase activity in Papaver becomes localised in the vegetative cell and generative nucleus at 5 h SI (Bosch and Franklin-Tong, 2007), although it was rarely observed in the pollen at 1 or 2 h post SI. This could imply that POB1 homologues in Papaver might be responsible for repressing caspase-3-like activity in compatible pollen and are being subsequently degraded in incompatible pollen tubes.

3.3. DISCUSSION

Ubiquitin-dependent proteasomal degradation pathway plays a role during SI in Papaver pollen and is most likely an important component of Papaver SI. In the present study, a pharmacological approach with the proteasome inhibitor MG132 was used to demonstrate the role for UbP pathway. A link between PCD and the UbP pathway in incompatible Papaver pollen was also demonstrated. MG132 is a membrane-permeable peptide aldehyde with chymotrypsin-like inhibitory activity. The treatment of poppy pollen with the selective proteasome inhibitor MG132 slightly altered tube morphology and decreased the length of pollen tubes in a concentration-dependent manner.

UbP pathway is involved in kiwifruit pollen tube organization, germination and maintenance of polarized pollen tube growth as ubiquitin and its conjugates were localized
mainly in the apex of pollen tubes (Scoccianti et al., 2003, Speranza et al., 2001). When proteasomal degradation was inhibited by MG132 dramatic changes to the morphology of pollen tube were observed, such as inhibition of pollen tube growth and pollen tube tip swelling and branching, and germination of more than one tube per grain, which was also observed in *Picea* pollen (Scoccianti et al., 2003, Sheng et al., 2006, Speranza et al., 2001). However the effects of the inhibitor were reversible since the removal of inhibitor restored growth. Those effects were not so dramatic in case of *Antirrhinum* pollen, although some pollen tube tips appeared to be swollen (Qiao et al., 2004b). Similar response was observed in *Papaver* pollen germination with severe pollen tube alterations at concentrations of MG132 exceeding 40 μM. The alterations in pollen tube morphology suggest that UbP activity is required for normal pollen tube growth and development in *Papaver*. As pollen tube growth was affected by MG132 at higher concentrations, 40 μM concentration was chosen for the experiments on *Papaver* pollen.

It is well established that the UbP pathway, and more precisely E3 ubiquitin ligases play a role in the sporophytic SI (Yee and Goring, 2009, Tantikanjana et al., 2010) and in the S-RNase based gametophytic SI (Meng et al., 2011). MG132 inhibited compatible pollination in *Antirrhinum*, but had little effect on incompatible pollination *in vitro* or *in vivo* (Qiao et al., 2004b). During compatible pollination the level of ubiquitinated S-RNases was significantly higher than during incompatible pollination in *Antirrhinum* and *Petunia* (Hua and Kao, 2006, Qiao et al., 2004b). The S-RNase degradation model was based on these findings. It was postulated that pollen determinant of S-RNAse based SI, SLF, specifically detoxifies non-self S-RNases via the UbP pathway allowing compatible pollinations and thus indicating that non-self pollen tubes might have been protected by specific degradation

In \textit{Papaver}, the ubiquitination was implicated to play a role during the SI response, rather than during the SC response. In incompatible pollen tube extract, an increased ubiquitination of yet unknown proteins was detected that is connected with the caspase-3-like activity, which acts downstream or in parallel with the UbP pathway.

The 26S proteasomal degradation was also demonstrated to play a role during SI in \textit{Brassica}, where activated SRK targeted E3 U-box ligase ARC1 to the proteasome, thus implicated to mediate the UbP pathway (Stone et al., 2003). Suppression of ARC1 in \textit{Brassica} or incubation of \textit{Brassica} stigmas with the proteasome inhibitors can lead to the breakdown of SI response (Stone et al., 1999, Stone et al., 2003). Following incompatible pollination, ARC1 targets substrate proteins, presumably compatibility factors, and their degradation by the 26S proteasome results in pollen rejection (Samuel et al., 2011).

Ubiquitin-dependent protein degradation is also important during plant-pathogen interactions and plant responses to the environmental factors (Dielen et al., 2010). Vacca et al.,(2007) demonstrated that decreased cell viability due to heat shock (HS) in tobacco BY2 cells was rescued by pretreatment with MG132, showing that the impairment of the proteasome function results in the prevention of cell death. Inhibition of the proteasomal degradation by MG132 also rescued the loss of viability in yeast cells undergoing acetic acid induced PCD (Valenti et al., 2008), and tungsten induced loss of viability in pea root cells (Adamakis et
The inhibition of proteasomal degradation inhibited length of tracheary elements in Zinnia cell cultures and also in A. thaliana (Zhao et al., 2008). Thus, the examples stated above indicate that proteasomal activity is involved in and is required for cell death in plant cells.

A similar picture was observed in Papaver SI, when the viability of incompatible pollen tubes was significantly restored when proteasomal degradation was inhibited by MG132 confirming the involvement of proteasomal degradation in the SI response in Papaver pollen. The use of proteasome inhibitors also allowed for the demonstration of the role of proteasomal degradation in PCD.

A direct link between proteasome-dependent degradation and plant PCD was established by Hatsugai et al., (2009). By using caspase-3 inhibitors, proteasome inhibitors and RNAi lines of the proteasome β-subunits, which possess proteolytic activity β1 (PBA), β2 (PBB) and β5 (PBE), they demonstrated that the proteasome was directly involved in bacteria-induced vacuole membrane and plasma membrane fusion and consequently hypersensitive cell death. Moreover, they identified that the proteasomal subunit PBA1 acts as a caspase-3-like enzyme.

The data, presented above confirm that caspase-3-like activity of SI samples was significantly higher compared to that in the untreated control pollen, while caspase-3-like activity of pollen tubes in which proteasomal degradation had been inhibited before induction of SI was significantly decreased although not to same level as control. This demonstrates that caspase-3-like activated PCD in poppy SI requires proteasome activation. Taking into account that the proteasomal subunit PBA1 acts as a caspase-3-like enzyme in A.
thaiana, it is tempting to suggest that the same happens in Papaver (Hatsugai et al., 2009). This was not tested as this work was conducted before the report by Hatsugai et al., (2009) was published; nevertheless, some parallels can be drawn.

Molecular mechanisms of SI share similarities with the histocompatibility in animals as well as with the hypersensitive response in plant-pathogen incompatible interaction as all these mechanism require high degree of recognition specificity to recognise between self and non-self. Recognition specificity during plant-pathogen response requires highly polymorphic resistance genes, and it involves a specific interaction between receptor at the epidermal cell surface and its cognate peptide ligands from pathogen (Coll et al., 2011). When a microbial pathogen invades the plant tissue, a defence mechanism is initiated in the plant that triggers signalling cascade of downstream events leading to inhibition of growth of plant cells in order to prevent spread of the pathogens. Downstream signalling events are strikingly similar to downstream events during Papaver SI: influx in cytosolic calcium, alterations of the actin cytoskeleton, activation of ROS, activation of caspase-like proteins and cell death. Some molecules were also implicated to play a role during SI and pathogen response, such as tobacco S-like RNase NE, that share similarities with RNases determining the specificity of the S-RNase based SI, inhibited the hyphal elongation of plant pathogens (Hugot et al., 2002, Dodds et al., 1996). Additionally, thioredoxin, that is implicated to negatively regulate Brassica SRK in the absence of SCR, was demonstrated to also negatively regulate Cladosporum fulvum-9 (Cf-9) (Cabrillac et al., 2001, Rivas et al., 2004). The incompatible interaction between Cf9, a membrane anchored glycoprotein and Avr9, a secreted protein, elicit a hypersensitive response and cell death. The interaction between Avr9 and Cf9 was also used when AtPUB17 protein was identified (Yang et al., 2006). PUB17 is an E3 ligase
that acts as positive regulator of cell death and defense responses in Solanaceae and Brassicaceae (Yang et al., 2006). PUB17 restored functional HR in plants that had HR impaired due to silencing of Cf9 (Yang et al., 2006). The incompatible interaction between Cf9, a membrane anchored glycoprotein and Avr9, a secreted protein, resulted in cell death. This is similar to the incompatible interaction between the pollen transmembrane protein, PrpS, and secreted stigmatic protein, PrsS, which also results in cell death in *Papaver* pollen. A collaboration with Dr. Ari Sadanandom, resulted in testing anti-PUB17 antibody during SI response in *Papaver*, as well as another E3 ligase, named POB1, that was identified as PUB17 interacting protein (A. Sadanandom, *personal communication*). Using antibodies against AtPOB1 the potential poppy homologues of POB1 were detected and their abundance changed over time, implicating a role for POB1 homologues during *Papaver* SI. Nuclei localised AtPOB1 act as a negative regulator of PCD, as plants that were deficient in POB1 produced spontaneous cell death (Sadanandom et al., 2008). *A. thaliana* AtPOB1 plants were resistant to virulent bacteria, they accumulated ROS and exhibited HR cell death (A. Sadanandom, *personal communication*). AtPOB1 represses the expression of genes that regulate defence against pathogen *Pseudomonas* (Sadanandom et al., 2008). If drawing a parallel between HR and SI systems, AtPOB1 homologues in *Papaver*, recognised by POB1 antibody, could function by repressing genes required for SI response. Interestingly, AtPOB1 proteins are localised in the nucleus (Sadanandom et al., 2008). Their localization was not yet examined in the *Papaver* pollen, however, if it is nuclear as well, this could explain their slow degradation, as it is known that DEVDase activity in *Papaver* peaks at 5 h of SI and becomes localised in the vegetative cell and generative nucleus (Bosch and Franklin-Tong, 2007), while it was rarely observed in the pollen at 1 or 2 h post SI. If POB1-like proteins act as negative regulators of
PCD by inhibiting caspase-3-like enzymes in normal pollen, then their degradation could trigger caspase-3-like activity.

This could implicate that POB1 homologues in *Papaver* might be responsible for repressing caspase-3-like activity in compatible pollen and are being subsequently degraded in incompatible pollen tubes. The identification of POB1 homologues during *Papaver* SI and their further analysis will establish a further evidence for comparison between two mechanisms, SI and HR.

Based on the results obtained, several models were created that could help to explain a link between UbP degradation and caspase-3-like activity in SI-induced PCD in *Papaver* pollen (Figure 3.9-3.10). These models assume that proteasome activation acts in parallel with DEVDase activity, because a decrease in DEVDase activity was observed where the proteasome inhibitor MG132 was used with the SI induced pollen tube protein extracts.

A model presented in Figure 3.9 is based on the discovery by Hatsugai et al., (2009), and it assumes that proteasome contains caspase-3-like activity, as they demonstrated in *A. thaliana* where proteasomal subunit PBA1 is a caspase-3-like enzyme. Although the experiments for this project were conducted before the publication by Hatsugai et al., (2009), some of the results regarding the involvement of the UbP pathway in *Papaver* SI can be explained by this model.
Figure 3.9.: Model of proteasomal degradation and caspase-3-like activation assumes that proteasome contains caspase-3-like activity. During the SI (yellow star) the proteins targeted for degradation are tagged with Ub via Lys48 residue. When the chain is long enough, proteins are degraded by the 26S proteasome and the model assumes that among them are also proteins with DEVD motif, recognised by DEVDases within the proteasome (yellow circles). During an SI we observed an increase in ubiquitinated proteins as well as increase in DEVDase activity. Substrate for caspase-3-like enzymes is represented as a fluorescent probe attached to DEVD tetrapeptide used in caspase-like assays, Ac-DEVD-AMC.

The Ac-DEVD-AMC substrate (used for illustration), substrates for caspase-3-like enzymes, and additional unknown proteins, are first tagged by ubiquitin molecules and targeted for the proteasomal degradation. Thus we can explain the increase in ubiquitination levels observed by western blot in SI induced pollen extracts. Once in the proteasome, the catalytic subunit (i.e. caspase-3-like enzyme) cleaves the substrate. When the proteasomal degradation is inhibited by MG132, the decrease in DEVDase activity is observed, which is explained by the model (Figure 3.6. and 3.9). MG132 inhibits proteasomal degradation of short-lived proteins, so it is assumed that proteasomal degradation is not completely inhibited, which would explain why there is not a complete decrease in DEVDase activity (Figure 3.6). In order to test this model we would need to use more specific inhibitors of proteasomal activity, blocking the catalytic subunit, such as Ac-APnLD-CHO (Hatsugai et al., 2009). However, this model cannot explain why there is less ubiquitination in pollen if we inhibit DEVDase activity by Ac-DEVD-CHO before the SI induction (Figure 3.7). If the model is correct, we should observe more ubiquitinated proteins as the proteasomal degradation and
caspase-3-like activation are inhibited, unless the inhibitor Ac-DEVD-CHO only inhibits DEVDase activity and does not affect proteasomal degradation by other catalytic subunits. Hatsugai et al., (2009) demonstrated that during the bacterial infection β1 and β5 subunits of proteasome with trypsin and chymotrypsin activity also play a role, and that their activity is reduced upon the inhibition of caspase-3-like activity. However, this might not be the case in our model, and therefore, the proteasomal degradation of other proteins that are not substrates for caspase-3 like enzymes might still continue. However, further experiments would be required to confirm any of these predictions.

Another model that could be proposed is based on the discovery by Hatsugai et al., (2009), who reported membrane fusion between the vacuolar membrane and the plasma membrane leading to leakage of vacuolar contents into the apoplast, thereby containing the bacterial infection. The membrane fusion is presumably suppressed by a negative regulator, which is degraded by the 26S proteasome upon bacterial infection. A similar mechanism could function during *Papaver* SI (Figure 3.10). Results obtained with the antibodies against AtPOB1 E3 ligases indicate the existence of a *Papaver* POB1 homologue that is affected by SI (Figure 3.8). AtPOB1 is a negative regulator of PAMP triggered immunity and effector triggered immunity and plays an important role in regulating HR cell death in *A. thaliana* and *N. benthamiana* plants upon fungal infection with *C. fulvum* (A. Sadanandom, personal communication). Taken together, based on the results obtained with the POB1 antibody during SI in *Papaver* pollen and the report by Hatsugai et al., (2009), an additional model was created. This model is set out in Figure 3.10. It presumes the existence of a negative regulator of DEVDase activity, which could be a putative POB homologue.
Normal pollen tube growth

POB1 homologue; negative regulator of DEVDase activity

POB1 homologue

Ub

SI

Ub

Ub

Ub

Ub

Ub

Ub

MG132

(continues on the next page)
**Figure 3.10.** Model that assumes the existence of negative regulator of DEVDase activity, possibly POB1 homologue (a) model assumes that during *Papaver* pollen tube growth, the hypothetical negative regulator of DEVDase activity, possibly POB1 homologue, is regulating the inactivity of DEVDases (b) During SI, negative regulator is tagged by UB through Lys48 residue and targeted for 26S proteasomal degradation and DEVDase activity is increased. Presumably there are also feedback loops between DEVDase activity and proteasomal degradation and DEVDase activity and negative regulator, (c) upon pretreating pollen with proteasomal inhibitor MG132 before SI, the accumulation of Ub is observed as well as decrease in DEVDase activity. Negative regulator is still ubiquitinated but not degraded as proteasomal activity is inhibited (d) when pollen in pretreated with DEVDase inhibitor Ac-DEVD-CHO prior to SI, DEVDase activity is inhibited and this inhibition could feedback to negative regulator, so it is not ubiquitinated anymore. Therefore negative regulator resumes its role and act in concert with DEVDase inhibitor to downregulate DEVDases. Assuming that the proteasomal activity acts upstream of DEVDase activity it is not affected, the degradation of other ubiquitinated proteins targeted during SI is undisturbed.

The model presented in Figure 3.10.a assumes that during normal undisturbed pollen tube growth, DEVDase activity is inhibited by a negative regulator, that could be a *P. rhoeas* POB1 homologue. Upon SI (Figure 3.10.b) the negative regulator is tagged by a chain of ubiquitins, connected by Lys48 and degraded by the 26S proteasome, thus releasing DEVDase activity. The model also assumes a feedback loop between the negative regulator, the 26S proteasome and DEVDase activity. If the proteasomal degradation is inhibited by MG132, then, as known from the results, DEVDase activity is reduced and due to the feedback loop, the ubiquitination of the negative regulator might be reduced, thus re-activating the inhibitor of DEVDase activity (Figure 3.10.c). However, ubiquitination of
other unknown proteins in the pollen tube still occurs but, due to the inhibition of 26S proteasome, their degradation is disabled, therefore the accumulation of ubiquitinated proteins can be observed on a western blot as an increase in ubiquitination signal (Figure 3.5). Upon the addition of the tetrapeptide inhibitor Ac-DEVD-CHO (Figure 3.10.d), the DEVDase activity is decreased, influencing the decrease in ubiquitination of the negative regulator (Figure 3.10.d). However, the proteasomal degradation of ubiquitinated proteins might not be affected, which could explain why the ubiquitination signal of pollen upon SI appeared stronger compared to pollen that was pretreated with Ac-DEVD-CHO before SI induction (Figure 3.7). In order to confirm or disprove this model additional experiments are required which would examine more closely the 26S proteasome activity as well as the POB1 substrate upon inhibition of the DEVDase activity.

Ubiquitination and proteasomal degradation are necessary for normal development of the pollen tube, and for degradation of damaged, outnumbered and unwanted proteins in the cell (Sheng et al., 2010, Sheng et al., 2006, Speranza et al., 2001). They are also strongly involved in the HR of plants to pathogens as well as in other types of SI responseses (Dielen et al., 2010, Trujillo and Shirasu, 2010, Chen et al., 2010, Yee and Goring, 2009, Vierstra, 2009). Taken together our data indicate that the proteasomal degradation acts together with, or upstream of, caspase activation and that there is a potential link between caspase activity and proteasomal degradation (see Figure 3.11).
Figure 3.11.: Schematic model of the SI response in *Papaver*. Upon the interaction of secreted stigmatic PrsS proteins and pollen transmembrane protein PrpS, a downstream signalling network is triggered, starting with an increase in K\(^+\) and [Ca\(^{2+}\)], who is signalling to the inhibition of pyrophosphatase activity and pollen tube tip growth, activation of MAPK, depolymerization of actin and microtubule cytoskeleton and appearance of punctate actin foci, activation of ROS and NO and activation of caspase-like activities, leading to the PCD of the pollen tube. The model presents DEVDase activity that most likely acts in parallel with proteasomal degradation. DEVDases are presented as yellow circles within the proteasome, as identified by Hatsugai et al., (2009).

However, in order to establish a clearer position of the UbP pathway on the SI timeline in *Papaver* pollen, and to draw further parallels between SI and HR, additional experiments would be required investigating proteasome inhibition, finding targets of ubiquitination during SI and specifying the role of the *P. rhoeas* POB1 homologue. In order to specifically elucidate the relationship between proteasome and DEVDases, and to determine whether DEVDase activity is part of the proteasome or acts separately, biochemical and imaging assays should be performed.

Gu et al., (2010) developed a membrane permeable fluorescent probe, MV151, containing a vinyl sulphone reactive group and Bodipy fluorescent group, that bind to the plant proteasome in an activity-dependent manner and could be used for live cell imaging of proteasome activities as well as quantification by protein gels during SI in *Papaver* pollen in...
the presence of the proteasome and the DEVDase inhibitor. In this way the proteasome could be also tested for other caspase-like activities. Antibodies against different subunits of the proteasome could also be obtained and tested. (Kisselev et al., 2003) measured different substrate specific sites of the proteasome by using isolated proteasomes from rabbit muscle or yeast. We could adopt this method and analyse a proteasome isolated from *Papaver* pollen for caspase-like activities using inhibitors and fluorescent tagged tetrapeptides using the plate reader during SI. Live cell imaging could be performed using CR(DEVD)$_2$ probe for DEVDase activity and MV151 probe for proteasome activity (Gu et al., 2010, Bosch and Franklin-Tong, 2007).

By performing biochemical assay of binding DEVDases on the biotinylated column and detecting the protein on a blot with antibody against PBA1, we could initially test whether the observation that the PBA1 subunit of proteasome exhibit DEVDase activity (Hatsugai et al., 2009) is specific for *A. thaliana* leaves or is more general within different plant species. Additionally, the studies of involvement of the UbP pathway during SI in *Papaver* pollen could be tested during SI in *A. thaliana* pollen expressing the *Papaver* pollen determinant PrpS (for details see next chapters). If the transgenic *A. thaliana* model were to produce similar results, then the genetic manipulations and mutations of different components could be performed and tested for response.

In order to identify targets of the proteasomal degradation a pull down assays could also be performed in a similar manner to that described above using biotinylated ubiquitin. The protein band could be excised and subjected to tandem mass spectroscopy (MS) which could reveal the nature of proteins specifically ubiquitinated during SI which are not ubiquitinated in normal untreated pollen.
The role of POB1 should also be analysed more closely. Western analysis using POB1 antibody should be repeated during time progression of the SI. In order to identify a link between putative PrPOB1 and DEVDase activity, POB1 should be also tested using inhibitors of proteasome activity and DEVDase activity. To test the model in Figure 3.10, where POB1 is presented as a negative regulator of SI, it would be interesting to see if it would be identified on MS. The POB1 sequence should be obtained and the protein could be inhibited by using designed POB1 antisense oligonucleotides. If POB1 does act as a negative regulator of SI then an increase in DEVDase activity should be observed even in untreated *Papaver* pollen. It would also be interesting to establish a link between POB1, whose *A. thaliana* and *N. benthamiana* homologues are implicated in plant pathogen responses, and ROS during SI in *Papaver* pollen.

By performing some of the additional experiments, the possible role of ubiquitination and proteasomal degradation should be more firmly linked to the SI in *Papaver* pollen and further parallels should be established in molecular mechanisms of SI and HR.
CHAPTER 4

Investigation of the interaction between

PrpS and PrsS
4.1. INTRODUCTION

As mentioned in introduction (section 1.4.3), the recent discovery of PrpS, the *Papaver* pollen S-determinant (Wheeler et al., 2009) has identified PrpS as a small, 20 kDa transmembrane protein, associated with the plasma membrane. It represents a novel class of receptor as it has no homology to any known protein in the database (Wheeler et al., 2009). In *Papaver*, SI is comprised of interaction between the secreted stigmatic PrsS proteins that act as a ligand to the receptor on the pollen tube. Identification of PrpS therefore strongly supports the long-standing hypothesis of receptor-ligand – like interaction (Wheeler et al., 2009).

To establish whether PrpS is functionally involved in the SI response, peptides were first tested with *in vitro* SI bioassays. Pollen with the haplotype PrpS$_1$/PrpS$_3$ was challenged with the incompatible recombinant stigmatic PrsS$_1$ and PrsS$_3$ proteins, which had been incubated with the PrpS$_1$ peptides. The pollen was ‘rescued’ when peptides were able to block the receptor-ligand type interaction and so far this was the first indication that the predicted extracellular region is involved in recognition and that PrpS might mediate pollen inhibition (Wheeler et al., 2009).

The main aim of this part of the project was to further characterize the binding between PrpS and PrsS in order to demonstrate that PrpS is the pollen S-determinant. There are several approaches to characterize and identify regions and residues of protein-protein binding interaction. Some of them, such as mutagenesis, were not possible due to difficulties to transform *Papaver* plants. Other approach that is widely used for protein-protein interaction is yeast-two-hybrid assay. Some preliminary investigations of interaction between PrpS and
PrsS were conducted by Natalie Hadjiosif, that identified a large number of false positive interactions as the PrsS proteins seem to auto-activate the system, while PrpS appeared toxic for the cells, presumably due to hydrophobicity of the protein. Considering that PrpS is a transmembrane protein, most likely adopted yeast-two-hybrid screens would be required, such as split-ubiquitin (Iyer et al., 2005). However, due to the time constraints we decided to test the interaction by the slot-blot binding assay, also known as far western assay, by which the protein of interest is bound on the membrane (on the blot) and not separated by electrophoresis as with western blot. The membrane is then incubated with binding partner and the potential interactions detected by specific antibody against binding partner. The far-western technique using peptide arrays is widely used for the mapping of epitopes and for the receptor-ligand interactions (Volkmer et al., 2011). Using peptide array and dot blot assay, a subdomain on Brassica stigmatic SRK was identified, that has an important role in binding of the pollen SCR (Kemp and Doughty, 2007). Furthermore, using dot blot assay, they demonstrated an affinity for binding between Brassica SRK microsomal membrane proteins and recombinant pollen SCR protein in an S-specific manner (Kemp and Doughty, 2007).

We wished to confirm whether one key criteria for the pollen S-determinant, which was binding to the pistil S-determinant, and the S-specific interaction between PrpS and PrsS was investigated. Initially, the secondary structure predictions of PrpS structure were analysed using various transmembrane protein prediction programmes and the extracellular domain was identified as a result of that search. By designing and synthesizing peptides that represent parts of the potential ligand binding domains of the extracellular region of PrpS, we wished to confirm the S-specific interaction between PrpS and PrsS in vitro.
This study formed a part of the publication (Wheeler et al., 2009) and the review (Wheeler et al., 2010). My contribution was the analysis of the *Papaver* pollen *S*-determinant, PrpS. I analysed the predicted transmembrane structure of the PrpS protein, and demonstrated using far-western ligand-blotting that a predicted extracellular domain interacts in an *S*-specific manner with the stigmatic PrsS proteins. The papers are inserted in the Appendix III.

4.2. RESULTS: The *S*-specific interaction of PrpS and PrsS

4.2.1. Analysis of predicted structure of PrpS

In order to test if the PrsS and PrpS interact, it was first necessary to obtain the predicted structure of the transmembrane protein PrpS.

The majority of membrane protein prediction programmes base their searches on the fact that transmembrane (TM) segments in proteins can be distinguished by continuous stretches of hydrophobic amino acid residues (Krogh et al., 2001, Hirokawa et al., 1998, Tusnády and Simon, 2001). Figure 4.1 shows the structure of the pollen PrpS₁ protein as predicted by the prediction programme TMHMM (http://www.cbs.dtu.dk/services/TMHMM/).
As PrsS proteins are secreted by the stigma it was expected that the interaction would occur on the extracellular part of the PrpS protein, as the extracellular part of the PrpS is exposed to the secreted stigmatic PrsS.

According to the first TMHMM prediction, PrpS₁ has 3 transmembrane domains and an extracellular loop segment, comprised of 35 amino acids, from position 63-97, which seems the most promising site for the interaction with the PrsS proteins (Figure 4.1.b).

The N-terminus of PrpS is possibly a signal peptide and therefore cleaved off (Petersen et al., 2011). The significance of this part of the protein being extracellular is not discussed further, but the work presented here discusses on the central extracellular part of the PrpS protein (35 aa), where the interaction most likely occurs. According to the TMHMM
prediction, 4 overlapping peptides, covering the extracellular domain sequence of extracellular domain, were synthesized by AltaBiosciences (Figure 4.2).

Peptide A1: VKLLGLVLHRLSFSE
Peptide A2: LHRLSFSEDQKWVA
Peptide A3: DQKVVAFGTAAICD
Peptide A4: TAAICDVLVPKNML
Scrambled A3 peptide: FTVDVKDCAAAGQI

Figure 4.2: Amino acid sequence of four overlapping peptides. Peptides represent PrpS1 extracellular 35aa domain predicted by TMHMM prediction programme and at the bottom scrambled A3 peptide.

Peptide A3 and its randomized version were already used in order to establish the functionality of the PrpS in the SI response. They were first tested for the effect on the SI phenotype (Figure 4.3) (Wheeler et al., 2009).

Figure 4.3: In vitro SI bioassay. (a) PrsS proteins were added to pollen and SI reaction occurred – pollen was inhibited; (b) PrsS proteins were added to pollen together with A3 peptide and pollen was “rescued”; (c) when scrambled A3 peptide was added to pollen and SI reaction occurred it indicates that scrambled peptide had no effect and pollen was inhibited (Wheeler et al., 2009).

Figure 4.3.b (work conducted by Kim Osman) illustrates that the addition of the 15-mer peptide A3, corresponding to parts of the predicted extracellular domain of the PrpS1, rescues pollen tube growth in the competition bioassay with PrsS1, whereas randomized version had no effect in the bioassay (Figure 4.3.c) (Wheeler et al., 2009). Since the in vitro bioassay experiment was carried out (Figure 4.3), two more PrpS alleles had been identified:
PrpS3 and PrpS8. Alignment and prediction of their structures, as predicted by TMHMM is presented in Figure 4.4.

![Figure 4.4](http://www.ebi.ac.uk/Tools/clustalw2/index.html)

As prediction programmes give just predictions, we studied the sequence analysis of all three PrpS alleles using various prediction programmes in order to obtain a more “realistic” picture of the predicted structures.

PrpS sequences were analysed using following transmembrane protein prediction programmes: TMHMM2.0, PredictProtein, SOSUI, HMMTOP, TMpred, TM-Finder, SPLIT4, ConPred II, Phobius (Hirokawa et al., 1998, Tusnády and Simon, 2001, Krogh et al., 2001, Rost and Liu, 2003, Deber et al., 2001, Juretić et al., 2002, Arai et al., 2004, Käll et al., 2004, Hofmann and Stoffel, 1993). The TMHMM prediction programme, whose prediction was first used to determine structure, is based on the hidden Markov model (Krogh et al., 2001). It incorporates hydrophobicity, charge bias, helix lengths, and grammatical constraints into one model (Krogh et al., 2001). TMHMM success rate in
discriminating soluble from membrane proteins is claimed to be higher than 99% in proteins without a signal peptide, but it can yield false positives and false negatives (Krogh et al., 2001). Figure 4.5 shows an alignment of PrpS1 using different prediction programmes. Synthesized peptides that correspond to the extracellular 35 aa segment were designed according to the TMHMM prediction. Two other programmes, SOSUI and PredictProtein predict that part of the “35 aa region” is extracellular and part transmembrane, while the TMPredict and HMMTOP programmes predict this part to be intracellular. All the prediction programmes give very similar topology for PrpS8 but, surprisingly, they predict an inverted structure with 4 TM segments for PrpS3.

Figure 4.5: Structural prediction of PrpS1 using different prediction programmes: TMHMM, SOSUI, PredictProtein, HMMTOP, TMPred, TMFinder and SPLIT4. Amino acids are colour-coded: pink – extracellular, red – transmembrane, blue – intracellular, black – no orientation given.
Membrane-protein topology-prediction methods are typically based on sequence statistics and contain hundreds of parameters that are optimized using known topologies of membrane proteins (Bernsel et al., 2008). Although the predictions differ, they all predict that PrpS has transmembrane helices and the alignment of the PrpS sequences suggests that all three proteins share a similar topology. Predictions indicated 3, 4, 5 or 6 TM domains for PrpS₁ but PrpS most likely has four transmembrane segments, as it is a good number to make a four helix bundle in the membrane. 3 and 5 TM proteins do exist but are rarer (Dr. A. Lovering, personal communication). Additional argument for four TM domains is the GxxxG motif that is involved in number of helix oligomers, that is also present in the PrpS₁, PrpS₃ and PrpS₈ sequence toward the C-terminus, and is also aligned with Drosophilla protein Flower, for which they demonstrated the formation of a channel (see Appendix I) (Dawson et al., 2002, Senes et al., 2004, Yao et al., 2009). So, if this motif is to potentially play a role in oligomerization in PrpS, then it should be within the fourth TM domain. Also the TMHMM diagram for PrpS₁ in Figure 4.1.a shows a fourth highly hydrophobic region at the C-terminus between aa 125 and 170. At this position there is a predicted TM domain in PrpS₃ and it is highly possible that it is applicable to the other alleles as there are stretches of hydrophobic acids in the TMHMM predicted “intracellular” domain in PrpS₁ and PrpS₈. This area is 39 % identical at the aa level between the three alleles and 74 % similar for PrpS₁ and 78 % similar for PrpS₈. The alignment of these 4 TM domains is such that a loop is extracellular as we have some evidence that it is involved in binding with PrsS (Wheeler et al., 2009), which has extracellular access to PrpS. Based on the predictions of the 4 prediction programmes, cartoon on Figure 4.6 shows 4 different topologies. Interestingly, the PredictProtein prediction shows very similar topology of PrpS₁ to what we think it might be. Peptide A₃ is located between aa 77-91. This peptide gave the strongest interaction using
far-western assay as well as competition bioassay. Nevertheless, despite all four programmes presented in Figure 4.6 identifying a major extracellular domain apart of TMHMM on whose prediction the synthesis and design was based, only PredictProtein predicted region of peptide A3 to be extracellular, SOSUI predicts only part of this peptide extracellular and part transmembrane, while HMMTOP predict it intracellular (Figure 4.6). Therefore, we must keep in mind that predictions are just that: predictions and not experimentally defined structures.

**Figure 4.6.** Cartoon of possible structural topologies of PrpS1 as revealed by different prediction programmes (a) TMHMM, (b) SOSUI, (c) PredictProtein, (d) HMMTOP. Cartoon Colour-coding: pink – extracellular, red – transmembrane, blue – intracellular. Numbers indicate the amino acid residue for PrpS1.

### 4.2.2. Binding assays

The aim of this experimental study relating to PrpS was to establish whether the PrpS and PrsS bind each other and whether the binding was S-specific. As all the above predictions indicate an extracellular loop segment, we carried out further experiments using the
synthesized 15-mer peptides, spanning over the extracellular domain of PrpS1. We used the ‘slot-blot’ method, a technique used in molecular biology to detect biomolecules also known as “far western”.

The peptides representing the extracellular domain of PrpS1 were applied directly to a PVDF membrane and the potential binding between PrpS1 peptides and recombinant PrsS1 protein was detected with antibody raised against PrsS1.

Figure 4.7. shows a representative slot-blot with all the peptides (see also Figure 4.2). From the figure we can see differential binding with all the peptides, except with the peptide A2, which gave no binding. No binding was observed with the scrambled peptide.

![Figure 4.7.](image)

Figure 4.7.: Binding assay with differential loading 10 μg, 1 μg and 0.1 μg of peptides A1, A2, A3, A4 and scrambled A3 peptide as control. Binding was detected with anti-PrsS1 at a concentration of 1:2000.

A problem that was initially observed with the slot-blot assay was relatively high background on the PVDF membrane. In order to reduce the background optimization was required, such as: adding extra washes to the membrane, cleaning up the antibodies and using higher titre of the primary antibody.
The peptide A3 gave positive results with *in vitro* bioassays so it was important to attempt to demonstrate the differential specific binding and allelic specificity. Despite the results in figure 4.7, the work was continued mostly with peptide A3. As the peptides synthesized corresponded to PrpS₁, we attempted to demonstrate the S-specificity of peptide A3 through binding with PrS₃ and detection with PrS₃ antibody (see Figure 4.8).

![Figure 4.8](image.png)

**Figure 4.8.** Binding assay with differential loading 10 μg, 1μg and 0.1 μg of A3 peptide and scrambled A3 peptide as a control. Peptide A3 designed over PrpS₁ was bound on the membrane, which was then incubated with PrS proteins. (a) membrane was incubated with PrS₁ proteins and binding was detected with anti-PrS₁, (b) membrane was incubated with PrS₃ proteins and binding detected with anti-PrS₃ (both at a concentration of 1:2000).

Figure 4.8 shows that the membrane incubated with anti-PrS₁ gave higher background than the membrane incubated with anti-PrS₃. The PrS₁-treated membrane had a clearer signal and differential loading was observed with peptide A3. Although there is a weak signal with 10 μg of scrambled peptide, this disappears when the background is subtracted. On the other hand, the PrS₃ blot showed an unusual pattern. At the first glance it seems that we have differential loading not only with peptide A3, but also with the scrambled peptide. Both blots were processed at the same time with the same reagents and solutions. The only difference
was in the primary antibodies. Closer inspection showed that there was also some signal in the position of the slots that were empty (i.e. which did not have any peptide loaded). Furthermore, the signal does not seem to be throughout the position of the slot but is confined to the edge. These indentations could also be observed in later blots, but only when detected with anti-PrsS₃ antibody (see representative Figure 4.9).

**Figure 4.9:** One of many repeated binding assays with differential loading 10 µg, 1 µg and 0.1 µg of A3 peptide and scrambled A3 peptide as a control. Peptide was bound on the membrane, which was then incubated with PrsS proteins. Recombinant PrsS₁ and PrsS₃ were loaded as the control at a concentration of 0.1 µg. (a) following binding, membrane was incubated with PrsS₁ proteins and binding was detected with anti-PrsS₁ (α-PrsS₁; 1:6000), (b) membrane was incubated with PrsS₃ proteins and binding detected with anti-PrsS₃ (α-PrsS₃; 1:500) antibody. E – empty slots.

The blot on Figure 4.9 shows the S-specific binding of A3 peptide with PrsS₁. Differential loading of peptide A3 is demonstrated and no signal with the scrambled peptide. The PrsS₁ antibody specifically recognizes the recombinant PrsS₁ protein and not the recombinant PrsS₃ protein. Several different titres were tested by western blot in order to try to get specific signals. Antibodies were used at concentrations that give an equivalent intensity of signal for 0.1 µg of peptides. Antibodies were also column purified to remove unspecific interactions. The blot probed with the anti-PrsS₃ antibody does not give a straightforward
result. If we focus only on the signals for peptide A3, then we might conclude that we have
differential loading. But we can also observe differential binding with the scrambled peptide.
Also, the recombinant PrsS3 protein is recognised specifically by the PrsS3 antibody,
whereas the recombinant PrsS1 protein gives a very weak signal that can be attributed to the
background. If we look at the blot as a whole, then we see quite strong signals in the
positions of the empty slots.

All the data so far shows that the anti-PrsS3 antibody does not bind specifically enough, as it
binds to the edges of empty wells and often gave a very high background signal.
Furthermore, alleles 1, 3 and 8 have 73 % aa identity in the area of peptide A3 (see Figure
4.10). For this reason we moved the focus of our attention to peptide A1 (Figure 4.2.) where
the aa identity between alleles is 27 %, so the region is more variable.

![Clustal W alignment of amino acid sequence of predicted extracellular loop of 3 different alleles of PrpS protein. Identical amino acids are marked with "*", conserved substitutions with "::" and semi-conserved substitutions with ".:."
](image)

In order to test for reciprocal S-specific interaction, we chose the area of peptide A1 in PrpS1
and PrpS8. Peptide A1 and its corresponding scrambled control from this less similar region
of extracellular domain were designed and synthesized by AltaBiosciences (Figure 4.11.).

![Peptides A1 for PrpS1 and PrpS8 and their scrambled versions, synthesized by AltaBiosciences
](image)
Peptides were tested using the binding assay for the S-specific interaction. The PrpS1 and PrpS8 peptides were bound to the membrane and incubated with the corresponding pistil PrsS1 and PrsS8 proteins, respectively. Binding was detected with anti-PrsS1 and anti-PrsS8 antibodies and can be seen in Figure 4.12.

![Figure 4.12](image)

**Figure 4.12.:** Binding assay between pollen PrpS1 and PrpS8 A1 peptides and recombinant stigmatic PrsS1 and PrsS8 proteins. Peptides were bound on the membrane at three different concentrations: 10 μg, 1.0 μg and 0.1 μg. The membrane was then incubated with PrsS proteins. Recombinant PrsS1 and PrsS8 were loaded as the controls at a concentration of 0.1 μg. (a) following binding, membrane was incubated with PrsS1 proteins and binding was detected with anti-PrsS1 (α-PrsS1; 1:6000), (b) membrane was incubated with PrsS8 proteins and binding detected with anti-PrsS8 (α-PrsS8; 1:4000) antibody.

We can see from Figure 4.12 the S-specific binding between PrpS1 and PrsS1 but not between PrpS8 and PrsS1. There was no binding detected between the scrambled control peptides and PrsS1 and very weak binding between 10 μg of the PrpS1 A1 scrambled control peptide and PrsS8. This provides the first formal evidence that PrpS and PrsS interact in an S-specific manner.
4.3 DISCUSSION

In this chapter we demonstrated with the binding assay the $S$-specificity for the interaction between PrpS and its ligand PrsS using peptides overlapping the extracellular domain of PrpS. Slot-blot binding assay is one of the commonly used techniques to show for protein-protein interactions (Southwick and Long, 2002). A modified version of slot-blot assay, a dot blot was also used by Kemp and Doughty, (2007), with which they documented binding in a haplotype-specific manner between stigmatic membrane proteins and recombinant SCR protein. Pollen recombinant SCR$_{29}$ was iodinated before incubation with a membrane with bound stigmatic microsomal membrane proteins from stigmas S$_{29}$S$_{29}$ and S$_{63}$S$_{63}$ of Brassica. They confirmed the $S$-specific binding of SCR$_{29}$ to S$_{29}$S$_{29}$ stigmas with fivefold higher affinity than to stigmatic membrane of different allele (Kemp and Doughty, 2007). Our assay demonstrated in a similar manner the $S$-specific binding between PrsS and PrpS, however, we demonstrated more specifically the binding between the extracellular domain of the transmembrane protein PrpS$_{1}$ and the recombinant ligand PrsS$_{1}$ and not just binding of a membrane extract with the recombinant protein (Kemp and Doughty, 2007, Wheeler et al., 2009).

Brassica-like SI is triggered upon the interaction between pollen coat protein $S$-locus Cysteine Rich (SCR), that binds to and activates $S$-locus Receptor Kinase (SRK), transmembrane protein encoded in stigmatic papillae (Kachroo et al., 2001, Takayama et al., 2001). A. thaliana is already used as a model for Brassica-like SI, expressing SRK/SCR gene pairs from A. lyrata and C. grandiflora (Nasrallah et al., 2002, Nasrallah et al., 2004, Boggs et al., 2009b). Use of this model system and mutagenesis enabled the identification of the 6-7 important amino acid residues in the extracellular domain of the SRK, determining
the specificity of the receptor-ligand interaction (Boggs et al., 2009a). These residues occur in two separate clusters located in the highly polymorphic sites of eSRK and were found on the same position in two variants tested.

The PrpS-PrsS interaction could be also demonstrated using different techniques, such as modified yeast-two-hybrid assays: split-ubiquitin or reverse Ras recruitment system (Stagljar and Fields, 2002). The reverse Ras recruitment system, for example, is based on Ras pathway in yeast. Membrane protein of interest (PrpS) is expressed in the membrane, while the interaction protein (PrsS) is fused to the mRas, that is cytoplasmic. The successful interaction between PrpS-PrsS should then result in the growth of yeast at 36 °C, which is otherwise too high (Stagljar and Fields, 2002). Alternative way to demonstrate PrpS-PrsS protein interaction would be pulldown assays, where PrsS could be immobilized as a bait on a His-column, and used to pull out the PrpS protein in the pollen extract. However, next steps are also the analysis of the PrpS residues that are important for the interaction with PrsS in the extracellular domain, and that are important for the intracellular interaction with yet unknown proteins. Additionally, the detailed analysis of TM domains could provide information’s regarding the oligomerization of the protein, or its potential to form a pore or a channel.
CHAPTER 5

Functional analysis *in vitro* of the *Arabidopsis thaliana* expressing *Papaver rhoeas* PrpS₁ and PrpS₃
5.1 INTRODUCTION

As described in the introduction, section 1.4.3., *Papaver rhoes* avoids self-fertilization on a molecular level by cell-cell specific self-recognition and rejection of self-pollen with transmembrane receptor PrpS, localised in the pollen plasma membrane, that lands on the stigma, which secretes small cysteine-rich proteins PrsS (Wheeler et al., 2009). Such SI interaction in the incompatible *Papaver* pollen triggers the cascade of downstream molecular events in pollen that ultimately lead to the inhibition of pollen tube growth and PCD of pollen tube (Thomas and Franklin-Tong, 2004).

We wished to see if the *Papaver* S-determinants PrpS and PrsS could be functionally transferred to other species, initially to *Arabidopsis thaliana*, triggering *Papaver*-like SI response. This would provide an important model for studying *Papaver* SI, as genetic manipulations are very difficult in *Papaver*, while *A. thaliana* is well amenable to the transformation by *Agrobacterium*, and extensive mutant collections already exist and are commercially available. The functionality of *Papaver* SI in other species could also have a long term potential for the F₁ hybrid breeding in crops. Many agriculturally important species, such as wheat, rye, barley or rice are self-compatible (SC) and for plant breeding and production of hybrids other ways of elimination of self-fertilization are essential, such as manual emasculation of anthers or use of cytoplasmic male sterility (for example very important for rice hybrid breeding) (Dwivedi et al., 2008, Cheng et al., 2007, Pelletier and Budar, 2007). These breeding programmes can be very expensive and lengthy to reach the end result so the stable introduction of SI that would interrupt SC might be positive and useful for future production of F₁ hybrids.
As mentioned above, the first step towards in the research of functionality of PrpS and PrsS in another angiosperm was the transformation into *A. thaliana*, a plant model organism, whose sequenced genome was sequenced (Initiative, 2000). *A. thaliana* is a SC species from the Brassicaceae family, which lost SI through mutation in the male specificity gene *SCR*, while female specificity gene *SRK* remained intact (Tsuchimatsu et al., 2010, Tang et al., 2007).

*A. thaliana* became the transgenic model of the Brassica SI system by transformation with SRK-SCR receptor ligand gene pair isolated from self-incompatible crucifers *Arabidopsis lyrata* or *Capsella grandiflora* (Nasrallah et al., 2002, Boggs et al., 2009b, Rea et al., 2010, Nasrallah et al., 2004). Their studies on transgenic *A. thaliana* focused on the analysis of the natural variation for the expression of SI, the evolutionary switch between outbreeding and inbreeding in *A. thaliana*, analysis of residues that determine SI specificity and identification of genes required for SI signalling and potential overlaps with other plant signalling pathways (reviewed in (Rea et al., 2010). Several different accessions of *A. thaliana* were transformed with SRK-SCR gene pair with SI response tested and they identified that not all accessions exhibit same degree of SI and they exhibit different levels of polymorphism at the S-locus and the SI modifier loci, leading to conclusion that inbreeding status of *A. thaliana* was probably developed multiple times by independent mutations in different accessions (Sherman-Broyles et al., 2007, Boggs et al., 2009c). *A. thaliana* presents an excellent system to study molecular mechanisms due to easy transformation using *Agrobacterium*, which can be difficult or impossible in some species, for example *Brassica* or in our case, *Papaver*. Transgenic *A. thaliana* expressing SRK-SCR gene pair was used for analysis of residues that are important for SRK specificity and therefore SI response. Site-directed mutagenesis approach was used on polymorphic amino acid residues in the extracellular domain of SRK
and six were identified to mediate the ligand-specific SI activation (Boggs et al., 2009a). Mutational analysis in transgenic *A. thaliana* expressing SRK-SCR also enabled the identification of mutation in RNA-dependent RNA polymerase RDR6, that acts as a negative regulator of SI and causes stigma exertion and simultaneous enhancement of SI, which implicated a broader role for SRK in pistil development (Tantikanjana et al., 2009). Further unexpected results were obtained when crossing *A. thaliana* plants expressing *SRKb-SCRb* genes that exhibit stable SI response with *A. thaliana* strains that contained inactive *PUB17* or *APK1b* genes, which were most similar to *Brassica ARCl* and *MLPK*. Those crosses resulted in unabashed SI response, which means that the role of ARCl and MLPK might not be as crucial for SI as previously reported (Rea et al., 2010, Kitashiba et al., 2011).

*Papaver*-based SI *A. thaliana* model was generated by *Agrobacterium*-mediated transformation of the *Papaver* *S*-determinants into *A. thaliana*. The transformations of *Papaver* *S*-determinants into *A. thaliana* were carried out by Barend de Graaf (BG plant lines) and Huawen Zou (HZ plant lines). This chapter describes in detail work with transgenic *A. thaliana* lines expressing PrpS1-GFP (BG16) and PrpS3-GFP (HZ3) and to lesser extent also 35S:PrpS1 (BG3). These were the lines that were available and chosen to work on at the start of this project. The constructs are presented in the Table 5.1.

**Table 5.1:** The cartoon illustrations of the constructs of the poppy SI determinants that were used for *Agrobacterium* mediated transformation into *A. thaliana*.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Resistance Marker</th>
<th>Name</th>
<th>Construct</th>
</tr>
</thead>
<tbody>
<tr>
<td>PrpS1</td>
<td>Kan</td>
<td>BG16</td>
<td><img src="image" alt="Construct BG16" /></td>
</tr>
<tr>
<td>PrpS3</td>
<td>Kan</td>
<td>HZ3</td>
<td><img src="image" alt="Construct HZ3" /></td>
</tr>
<tr>
<td>PrpS1</td>
<td>Kan</td>
<td>BG3</td>
<td><img src="image" alt="Construct BG3" /></td>
</tr>
</tbody>
</table>
To achieve tissue specific expression of the male \(S\)-determinants \(\text{PrpS}_1\)-GFP and \(\text{PrpS}_3\)-GFP in the pollen of \textit{A. thaliana}, a strong, pollen specific promoter from tobacco, \textit{ntp303}, was used (Weterings et al., 1995). For BG3 line, \(\text{PrpS}_1\) expression was cloned under the control of the cauliflower mosaic virus 35S promoter (Hull, 1983) that has a constitutive expression throughout the plant. In order to detect the transformed plant cells, the kanamycin (Kan) resistance cassette was on the T-region of the construct acting as a selection marker. In all lines except BG3, \(\text{PrpS}\) was expressed as a fusion protein with green fluorescence protein (\textit{GFP}) towards its C-termini.

The results of the study, presented in this chapter, resulted in a publication where I am joint first author. The paper is presented in the Appendix III.

\textbf{AIMS}

Part of this PhD study was the functional analysis of \textit{Papaver} SI in \textit{A. thaliana}. Several approaches were taken: \textit{in vitro} and semi-\textit{in vitro} described here, and \textit{in vivo} (described later in Chapter 6). The initial studies were the characterization and segregation analysis based on the inheritance of the resistance markers Kan and reporter marker GFP of \(\text{PrpS}_1\)-GFP and \(\text{PrpS}_3\)-GFP expressing \textit{A. thaliana} lines, and identification of homozygous progenies of transgenic lines with the highest expression of \textit{Papaver} \(S\)-determinants. The use of the homozygous lines would optimise the reproducibility of the study. Once the highest expressing homozygous lines were identified, the functional analysis studies were conducted. The aim was to conduct \textit{in vitro} studies on transgenic \textit{A. thaliana} pollen expressing \(\text{PrpS}_1\)-GFP and \(\text{PrpS}_3\)-GFP fusion protein. We wished to establish whether \(\text{PrpS-}\)
GFP in *A. thaliana* pollen responded to its cognate stigmatic recombinant proteins in the same manner as *Papaver* pollen. The SI response was induced in transgenic *A. thaliana* pollen expressing PrpS$_1$-GFP or PrpS$_3$-GFP and germinating in Petri dishes, with an addition of the cognate recombinant stigmatic PrsS$_1$ or PrsS$_3$ proteins. Various responses were analysed, such as: inhibition of pollen tube growth, decrease of pollen viability and the appearance of puncate actin foci. The outline of these experiments is presented on Figure 5.1. The S-specific response was ensured by analysing PrpS-GFP expressing *A. thaliana* pollen upon incubation with biologically inactive incompatible PrsS proteins, treatment with compatible proteins and incubation of wild type Col-0 *A. thaliana* pollen using *Papaver* stigmatic PrsS recombinant proteins.

![Figure 5.1](image-url)

**Figure 5.1:** Schematic diagram presenting the outline of the *in vitro* functional analysis experiments. *In vitro* SI reaction was induced in transgenic *A. thaliana* pollen expressing PrpS1-GFP or PrpS3-GFP and the several responses were tested, such as inhibition of pollen tube growth, decrease of viability and the appearance of punctate actin foci.

The inhibition of pollen tubes was analysed by measuring pollen tube length, analysed the organisation of the actin cytoskeleton with the use of rhodamine phalloidin and the viability of pollen tubes at different time points using Evans Blue dye as well as investigated pollen cell death and DEVDase activity of transgenic *A. thaliana* pollen.
5.2 RESULTS

5.2.1. Characterization of *A. thaliana* expressing *Papaver* SI system

At the start of the project, the seeds from up to 30 independent primary transformants of different lines were available: BG16 expressing PrpS$_1$-GFP, HZ3 expressing PrpS$_3$-GFP and BG3 expressing PrpS$_1$ using constitutive 35S promoter, whose characterisation is presented in this chapter and lines HZ1 expressing PrsS$_1$ and HZ2 expressing PrsS$_3$ and BG6 expressing whole *S*-locus whose characterisation is presented in the next Chapter 6.

Before carrying out the functional characterisation it was necessary to identify the highest expressing individual lines. Although lines were pre-screened on the selection medium, there was the possibility that they did not contain the insert corresponding to the *S*-genes within their genome. Therefore we screened for the plants with inserts by PCR on genomic DNA purified from leaf disks. The T$_2$ generation of the transgenic *A. thaliana* lines were tested for the presence of the inserts by PCR using genomic DNA as a template. Lines BG16 and HZ3 were tested for the presence of PrpS$_1$ and PrpS$_3$ respectively. Once having confirmed if the insert was present, only the plants that showed the insert by PCR were tested if they were expressing the PrpS by reverse transcription-PCR (RT-PCR).

We wished to identify at least two individuals with the highest expression of *Papaver* *S*-determinants within each transgenic *A. thaliana* line, so the further functional analysis could be accomplished.
5.2.1.1. PrpS expressing A. thaliana lines

Seeds of Arabidopsis::PrpS-GFP were initially screened on the germination medium MS containing kanamycin. Only the plants containing the PrpS-GFP transgene were expected to survive, so resistant and non-resistant seedlings were counted for segregation analysis. Heterozygous plants were expected to segregate in Mendelian fashion with 3:1 ratio between resistant and non-resistant seedlings, while homozygous plants were expected to be 100% resistant on the kanamycin containing media. Another marker utilised in the analysis of homozygous lines was the reporter gene GFP, expressed in transgenic A. thaliana pollen. It was expected that in homozygous lines GFP would be expressing in every pollen grain (100% expression). The results of the segregation analysis are presented in Tables 5.2 and 5.3.

### Table 5.2: Segregation of Kan resistance and GFP expression in progenies of transgenic A. thaliana lines expressing Papaver pollen determinant PrpS1

<table>
<thead>
<tr>
<th>Line (PrpS1-GFP)</th>
<th>Generation</th>
<th>KanR</th>
<th>KanS</th>
<th>N</th>
<th>Ratio</th>
<th>$\chi^2$ (3:1)</th>
<th>GFP (%) ± SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>BG16B# 25</td>
<td>T2</td>
<td>202</td>
<td>84</td>
<td>3</td>
<td>2.4:1</td>
<td>2.91</td>
<td>67 ± 6</td>
</tr>
<tr>
<td>BG16B# 25.1</td>
<td>T3</td>
<td>248</td>
<td>15</td>
<td>3</td>
<td>16.5:1</td>
<td>52.23</td>
<td>91 ± 1</td>
</tr>
<tr>
<td>BG16B# 25.1-1</td>
<td>T4</td>
<td>354</td>
<td>1</td>
<td>8</td>
<td>HO</td>
<td>95.35</td>
<td>97 ± 1</td>
</tr>
<tr>
<td>BG16B# 25.1-2</td>
<td>T4</td>
<td>134</td>
<td>0</td>
<td>2</td>
<td>HO</td>
<td>44.67</td>
<td>96 ± 2</td>
</tr>
<tr>
<td>BG16B# 25.1-1.x</td>
<td>T5</td>
<td>351</td>
<td>1</td>
<td>10</td>
<td>HO</td>
<td>114.7</td>
<td>100</td>
</tr>
<tr>
<td>BG16A# 19</td>
<td>T2</td>
<td>291</td>
<td>126</td>
<td>4</td>
<td>2.3:1</td>
<td>6.05</td>
<td>67 ± 5</td>
</tr>
<tr>
<td>BG16A# 19.1</td>
<td>T3</td>
<td>353</td>
<td>125</td>
<td>5</td>
<td>2.8:1</td>
<td>0.34</td>
<td>79 ± 2</td>
</tr>
<tr>
<td>BG16A# 19.8</td>
<td>T3</td>
<td>152</td>
<td>79</td>
<td>3</td>
<td>1.9:1</td>
<td>10.4</td>
<td>80 ± 2</td>
</tr>
<tr>
<td>BG16A# 19.8-2</td>
<td>T4</td>
<td>64</td>
<td>19</td>
<td>2</td>
<td>3.3:1</td>
<td>0.20</td>
<td>66 ± 1</td>
</tr>
<tr>
<td>BG16A# 19.8-3</td>
<td>T4</td>
<td>115</td>
<td>10</td>
<td>3</td>
<td>11.5:1</td>
<td>19.27</td>
<td>95 ± 1</td>
</tr>
<tr>
<td>BG16A# 8.3.1</td>
<td>T4</td>
<td>117</td>
<td>85</td>
<td>2</td>
<td>1.4:1</td>
<td>31.4</td>
<td>82 ± 2</td>
</tr>
<tr>
<td>BG16A#13</td>
<td>T2</td>
<td>53</td>
<td>7</td>
<td>3</td>
<td>7.6:1</td>
<td>5.69</td>
<td>87 ± 1</td>
</tr>
<tr>
<td>BG16A#13.1</td>
<td>T3</td>
<td>84</td>
<td>11</td>
<td>3</td>
<td>7.6:1</td>
<td>9.12</td>
<td>97 ± 2</td>
</tr>
<tr>
<td>BG16A#13.2</td>
<td>T3</td>
<td>118</td>
<td>12</td>
<td>3</td>
<td>10:1</td>
<td>17.24</td>
<td>98 ± 1</td>
</tr>
</tbody>
</table>
With the segregation analysis, the expected Mendelian segregation for the single insert was 3:1 (Kan<sup>R</sup>:Kan<sup>S</sup>) and the proportion of the GFP expressing pollen in heterozygous line would be 50 % (1:1 Mendelian segregation ratio). Such segregation was observed in many of the analysed PrpS-GFP expressing A. thaliana lines but not in all of them. After selfing, the T<sub>3</sub> generations sometimes exhibited ratio ~15:1 (BG16B#25.1, HZ3.1-15) which probably corresponds to the transgene insertion at two independent loci. But with the increased segregation, we also observed an increase in GFP expression in pollen. The GFP expression in pollen of PrpS<sub>1</sub>-GFP expressing A. thaliana was around 67 % in T<sub>2</sub> BG16B#25 and 91 % in the BG16B#25.1 line with the higher segregation ratio. After another round of selfing the line was homozygous in generation T<sub>4</sub> and also GFP expression in pollen was 97 % at that generation. As the total pollen was counted for the GFP expression, the pollen that was not expressing PrpS<sub>1</sub>-GFP exhibited distorted shape under microscope, pollen grains were smaller and it was not viable. Therefore we concluded that line, homozygous for the Kan resistance feature, was also homozygous for the GFP expression, as expected.

However, line BG16A#19 was not yet homozygous for the insert at generation T<sub>4</sub>. It shows 3:1 segregation ratio in generation T<sub>2</sub> and T<sub>3</sub> of BG16A#19.1, with 79 % of pollen expressing GFP. However, the T<sub>3</sub> line BG16A#19.8 that was also identified as the highest expressing line (described later in this Chapter), shows segregation ratio of 2:1 with 80 % of pollen expressing PrpS<sub>1</sub>-GFP. Although segregation ratio 2:1 seems to be associated with the homozygous lethality (Limanton-Grevet and Jullien, 2001), the T<sub>3</sub> generation of BG16A#19.8 exhibited 3:1 segregation (BG16A#19.8-2) with 66 % GFP expression, and 11.5:1 (BG16A#19.8-3) with 95 % GFP expression in pollen. The higher segregation ratio could be associated with transgene insertion at two loci, possibly linked (Feldmann et al., 1997). Line BG16A#19 was not yet homozygous for the insert at generation T<sub>3</sub>, however the
T₃ line BG16A#19.8 was also identified as a highest expressing line (described later in this Chapter). Additionally, the unusual segregation ratio of 1.4:1 was also observed for the T₄ generation of the line BG16.8#3.1, which was also identified to express PrpS₁-GFP in high concentrations, while the 82 % of the pollen was expressing PrpS₁-GFP.

Distorted segregation ratios were observed also in the line BG16A#13, which was added to the experiments later on, due to its very high GFP expression of more than 90 % in generation T₃. Its segregation shows 8:1 in T₁ with very high PrpS₁-GFP pollen expression of 87 % and 8:1 and 10:1 in T₃ with homozygous PrpS₁-GFP pollen expression of 97 % and 98 %, respectively. Such distorted segregation ratios were previously associated with the segregation at the S-locus in Petunia (Harbord et al., 2000) where distorted segregation ratios of 8:1 or higher were anticipated due to gamethophytic selection of S-alleles.

However, in the case of Papaver S-transgenes in A. thaliana, the location of the insert was not identified. Line BG16A#13 was having homozygous GFP expression in pollen in generation T₃, which did not show any abnormalities, but was still heterozygous for the Kan resistance marker. The explanation for this discrepancy could be that the two markers in line BG16A#13 follow the Mendelian segregation of independent segregation in T₃ generation because the markers are located on the opposite sides of the T-DNA construct so it could be possible for one inserted construct to lose the Kan marker. However, pollen from BG16A#13 exhibited normal germination rates compared to Col-0 and for that reason it was used in some preliminary experiments, such as caspase assays.

For functional analysis the progeny of lines BG16B#25 and BG16A#19 were selected (see also RT-PCR analysis later in this section).
The unusual high segregation ratios most often indicate the segregation of inserts at two linked loci. Whereas the segregation of inserts at two unlinked loci gives 15:1 ratio, the segregation at two linked loci could give rise to segregation ratios $3:1 < x < 15:1$ (Feldmann et al., 1997). If we assume multiple insertion and independent segregation of the separated fusion construct then we would be expecting greater GFP expression than single insert, which is what we observed. The variable transgene expression could also be due to the position effects, meaning that the actively transcribed regions of the genome or a transgene insertion near enhancer elements are more likely to be expressed. Therefore, a few seedlings with the insert near the highly expressed region could give rise to the observed intense expression of GFP (for example line BG16A#13 which exhibited very high GFP expression, while the Kan segregation was heterozygous 8:1 and 10:1 ratios). Another possible reason for distorted segregations of double T-DNA copy could results from epigenetic transgene silencing, which is common for repetitive sequences, potentially as a defence response against foreign DNA (Jorgensen et al., 1996, Muskens et al., 2000, De Buck et al., 2007, Matzke and Matzke, 1998). Double insertion of the gene often arrange as inverted repeats of which many are dominant silencing loci and can repress the expression of homologous genes, associated with increase in DNA methylation (Muskens et al., 2000). The post-transcriptional silencing of genes results from degradation of double stranded RNA (dsRNA) directed DNA methylation and dsRNA induced degradation of homologous RNAs (Muskens et al., 2000).

Table 5.3 presents the results of segregation analysis of PrpS$_3$-GFP expressing A. thaliana lines.
Table 5.3: Segregation of kanamycin resistance and GFP expression in progenies of transgenic A. thaliana lines expressing *Papaver* pollen determinant PrpS3. Statistical analysis was performed using Chi square analysis ($\chi^2$ value presented). Percentage of GFP expressing pollen is presented as mean value ± SEM. HO–homozygous. N.A.–not analysed.

<table>
<thead>
<tr>
<th>Line (PrpS3-GFP)</th>
<th>Generation</th>
<th>Kan$^R$</th>
<th>Kan$^S$</th>
<th>N</th>
<th>Ratio</th>
<th>$\chi^2$ (3:1)</th>
<th>GFP (%) ± SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>HZ3.1</td>
<td>T1</td>
<td>141</td>
<td>49</td>
<td>2</td>
<td>3:1</td>
<td>0.016</td>
<td>/</td>
</tr>
<tr>
<td>HZ3.1-3</td>
<td>T2</td>
<td>116</td>
<td>39</td>
<td>2</td>
<td>3:1</td>
<td>0.0022</td>
<td>71 ± 3</td>
</tr>
<tr>
<td>HZ3.1-15</td>
<td>T2</td>
<td>169</td>
<td>9</td>
<td>2</td>
<td>19:1</td>
<td>37.8</td>
<td>88 ± 3</td>
</tr>
<tr>
<td>HZ3.1-3.1</td>
<td>T3</td>
<td>98</td>
<td>39</td>
<td>2</td>
<td>2.5:1</td>
<td>0.87</td>
<td>83 ± 3</td>
</tr>
<tr>
<td>HZ3.1-15.1</td>
<td>T3</td>
<td>272</td>
<td>21</td>
<td>4</td>
<td>13:1</td>
<td>49.7</td>
<td>96 ± 1</td>
</tr>
<tr>
<td>HZ3.1-3.1-x</td>
<td>T4</td>
<td>269</td>
<td>28</td>
<td>7</td>
<td>7:1</td>
<td>21.7</td>
<td>96 ± 1</td>
</tr>
<tr>
<td>HZ3.1-15.1-x</td>
<td>T4</td>
<td>200</td>
<td>0</td>
<td>8</td>
<td>HO</td>
<td>66.7</td>
<td>98 ± 1</td>
</tr>
<tr>
<td>HZ3.2</td>
<td>T1</td>
<td>0</td>
<td>200</td>
<td>2</td>
<td>0:1</td>
<td>600</td>
<td>N.A.</td>
</tr>
<tr>
<td>HZ3.3</td>
<td>T1</td>
<td>132</td>
<td>54</td>
<td>2</td>
<td>2.4:1</td>
<td>1.61</td>
<td>N.A.</td>
</tr>
<tr>
<td>HZ3.4</td>
<td>T1</td>
<td>0</td>
<td>100</td>
<td>1</td>
<td>0:1</td>
<td>300</td>
<td>N.A.</td>
</tr>
<tr>
<td>HZ3.5</td>
<td>T1</td>
<td>96</td>
<td>56</td>
<td>2</td>
<td>1.7:1</td>
<td>11.37</td>
<td>N.A.</td>
</tr>
<tr>
<td>HZ3.7</td>
<td>T1</td>
<td>46</td>
<td>26</td>
<td>1</td>
<td>1.8:1</td>
<td>4.74</td>
<td>N.A.</td>
</tr>
<tr>
<td>HZ3.8</td>
<td>T1</td>
<td>53</td>
<td>32</td>
<td>1</td>
<td>1.6:1</td>
<td>7.25</td>
<td>N.A.</td>
</tr>
</tbody>
</table>

In the A. thaliana HZ3 line expressing PrpS3-GFP, the seeds of transformed plants were pooled together so the T1 do not represent independent primary transformants, but rather plants that were selected on the Kan containing MS medium from the pool of T0 seeds. Eight of T1 lines presented in Table 5.3 were grown on the MS-Kan medium, two out of those were Kan sensitive (HZ3.2 and HZ3.4) and only two lines, namely HZ3.1 and HZ3.3 showed the expected 3:1 segregation ratio based on the inheritance of Kan resistance. Lines HZ3.5, HZ3.6 and HZ3.7 show non-Mendelian segregation ratio of 2:1 and were not included in subsequent experiments.

The line HZ3.1 was analysed as well as its subsequent generations (presented on Table 5.3). One progeny of the line, HZ3.1-3 exhibits 3:1 ratio in generation T2 and T3, indicating it is still segregating following Mendelian inheritance, while in T4 it exhibits an abnormal 7:1 segregation ratio. However, despite following the Mendelian rules for Kan resistance gene, the PrpS3-GFP protein expression is 71 % in T2 and 83 % in T3, indicating the 3:1
segregation of PrpS3-GFP pollen expression. The other progeny of the line 3.1, HZ3.1-15 shows a 19:1 segregation in T2 with 88 % pollen GFP expression. 13:1 ratio in T3, which suggests that the PrpS3-GFP insert might be present at two loci (normally 15:1 Mendelian segregation) and it reached homozygous state by pollen PrpS3-GFP expression of 96 %. The HZ3.1-15 line reached homozygous state for the both markers (Kan and GFP) in generation T4. This line was also used in the experiments testing for the functional analysis, while line HZ3.1-3.1 was used only in preliminary tests due to its unusual segregation pattern (3:1 in T2, 3:1 in T3, 7:1 in T4), although the GFP expression in T4 was complete.

In order to identify lines with the highest expression of the transgene the RT-PCR was used. Initial PCR screening for the presence of PrpS1 and PrpS3 inserts was performed on the leaves of young plants, using gene specific primers and flowers of the plants that were positive for the insert were analysed by RT-PCR. Figure 5.2.a present the screening results on BG16 line expressing PrpS1-GFP, while Figure 5.2.b present HZ3 line expressing PrpS3-GFP.
Figure 5.2. (a) PrpS$_1$ expressing *A. thaliana* plants that were positive for inserts were analysed by the RT-PCR in order to identify the highest expressors (lines BG16.8#3, BG16A#19 and BG16B#25). (b) PrpS$_3$ expressing *A. thaliana* plants that were positive for inserts were analysed by RT-PCR in order to identify the highest expressors (line HZ3.1). GAPC was used as a standard and is presented at the lower part of each image. Reactions without DNA and with Col-0 DNA were used as a control.

The expression levels were analysed using RT-PCR, which was conducted on the flowers from the same plants used for genotyping and housekeeping gene GAPC was used as a control. Figure 5.2.a shows the results of the RT-PCR analysis on BG16 flowers. PrpS$_1$ primers were designed to align on 5’ and 3’ ends of the insert and a size of the obtained product was expected to be 583 bp. BG16B#25 exhibited uniform expression with all the samples analysed, while the expression of the BG16A#19 and BG16.8#3 was less uniform. The RT-PCR analysis was done on PrpS$_3$ expressing flowers using HZ3.1 line only, with the exception of the samples 7 and 13 (Figure 5.2.b). The reason for HZ3.3 not being analysed by RT-PCR was with time and material constraints at the time. PrpS$_3$ primers were designed to yield a product of 436 bp. Under the same conditions, wt Col-0 was used as a control but not yielding any product.
It can be observed in Figure 5.2.b that PCR product of PrpS3 gene resulted in double band. There could be a series of technical reasons for doublets in the gel, from the different concentrations between the forward and reverse primers, different annealing temperatures, to non-homogeneous distribution of the dye between molecules (Carlsson et al., 1995). However, technical issues were taken into the account and no problems were found with primers or melting temperatures. Primers were also used in genotyping PCR reaction where they produce normal single band. The DNA electrophoresis was repeated using 1 % agarose gel and running under lower currents for longer time but with the same result. In the process of transformation cDNA of PrpS3 was used; however the PCR bands presented in the Figure 5.2.b were obtained from RNA so this could likely be a splicing variants. PrpS1 and PrpS8 were already demonstrated to produce splice variants but not PrpS3 (Hadjiosif, 2008).

The expression of PrpS3 was not uniform with all the flowers, six flowers exhibited higher expression than the rest. Further selection of 5 samples was made initially, by quantifying the intensity of the bands using ImageJ software (see Figure 5.3) as the highest expressing lines were desired for the functional analysis.

**Figure 5.3.** (a) Top 5 highest expressing PrpS1 and PrpS3, and the corresponding GAPC control. Legend: 1-BG16.8#3.5, 2-BG16A#19.5, 3-BG16A#19.8, 4-BG16B#25.1, 5-BG16B#25.9, 6-HZ3.1-2, 7-HZ3.1-3, 8-HZ3.1-13, 9-HZ3.1-14, 10-HZ3.1-15. (b) Column chart presenting ratio between transgene signal and GAPC signal. Quantification was performed using ImageJ software. Highest expressors outlined in red.
cDNA samples of plants BG16.8#3.5, BG16A#19.5, BG16A#19.8, BG16B#25.1 and BG16B#25.9 from PrpS1 expressing flowers, and HZ3.1.2, HZ3.1.3, HZ3.1.12, HZ3.1.14 and HZ3.1.15 from PrpS3 expressing flowers were further re-analysed on the DNA gel and compared to GAPC control (Figure 5.3.a).

In order to get some idea of the quantification of expression of the transgene, the peak band intensity was measured using ImageJ software for each sample and its GAPC control (Figure 5.3.b). From each transgenic line, two plants with the highest expression identified by intensity ratio were used in subsequent functional experiments. The apparent lines with the highest expression of the transgene were: BG16A#19.8 and BG16B#25.1 expressing PrpS1-GFP, HZ3.1-3 and HZ3.1-15 expressing PrpS3-GFP, however lines BG16B#25.1 and HZ3.1-15 exhibited higher and homozygous expression of GFP (see Tables 5.2 and 5.3).

The progeny of these plants were used in further experiments to determine whether the PrpS-GFP was functional when expressed in *A. thaliana* pollen. Family tree of the PrpS1 and PrpS3 expressing lines BG16 and HZ3 are presented in the Appendix II.

Since the RT-PCR results showed that mRNA was present in the pollen of transgenic *A. thaliana* expressing PrpS1-GFP, we tried to confirm the PrpS1 expression at the protein level in BG16 pollen. For this aim we extracted proteins in two different buffers suitable for the extraction of transmembrane (TM) proteins: first buffer was 2x extraction buffer for TM proteins and second buffer was additionally optimised for extraction of TM proteins (*personal communication with Andy Lovering*). Proteins were extracted with a buffer with high Glycerol and Triton X-100 concentrations, and the sample was not boiled prior to loading on SDS-PAGE, as proteins are likely to aggregate during boiling. After the transfer of the gel onto the membrane, the blot was incubated in stripping buffer for short time before
detection with antibodies (see Materials and Methods). Proteins were separated by western blotting and detected using alkaline phosphatase (Figure 5.4).

**Figure 5.4:** Western blot analysis of pollen extract from BG16 pollen expressing PrpS₁ and controls, using two different extraction buffers. Buffer 1 (left side) was 2x extraction buffer and buffer 2 (right side) was optimised for extraction of TM proteins (with increased Glycerol and Triton X-100 concentrations; samples were not boiled prior to loading on a membrane; after the transfer, the blot was incubated in stripping buffer before detection with antibodies; see Materials and Methods). As controls were used *A. thaliana* non-transgenic Col-0 pollen extract, as positive control was used *Papaver* S₁S₃ pollen extract and as negative control *Papaver* S₃S₈ pollen extract. Samples were loaded on the membrane in duplicates, and after blotting the membrane was incubated in stripping buffer at 55 °C for 15 min before the incubation with antibodies. Primary antibody used was rat polyclonal anti-PrpS₁ (60C) in 1:1000 concentration and secondary was anti-rat-AP in 1:5000 concentration. PrpS₁ in *Papaver* extract is indicated by red star.

In *Papaver* S₁ and S₃ (S₁S₃) pollen extract several bands could be observed with both extraction buffers, but extraction buffer 2 gave clearer result. PrpS₁ size should be ~20 kDa so the lower strong band correspond to it. However, bands in *Papaver* S₁S₃ pollen extract were also detected at ~28 kDa mark, very strong at 30 kD and weaker at 36 kDa, The upper band could correspond to PrpS dimer. In BG16 pollen extract, we could observe bands at higher molecular weight ~28 and 30 kDa with the 28 kDa band being more intense than 30 kDa. There were also some weaker bands at ~10 kDa.

If PrpS forms multimers, this could indicate toward its potential function as a channel as there are some reports of the small receptors that multimerize in order to form a channel, for example small and novel transmembrane protein named Flower identified in *Drosophila* nervous system (Yao et al., 2009). Flower so far offers most analogies to PrpS proteins.
although they do not share much sequence homology but rather a topological similarity. And since the Flower function has been established and demonstrated to multimerize and form a Ca\textsuperscript{2+} channel, this might be a possibility for PrpS as well. However, in order to confirm the dimer or multimer using western blot further optimization might be required.

5.2.2. \textit{In vitro} functional analysis of \textit{Papaver} SI determinants in transgenic \textit{Arabidopsis thaliana} \textit{in vitro}

As described earlier several PrpS\textsubscript{1}-GFP and PrpS\textsubscript{3}-GFP expressing transgenic \textit{A. thaliana} lines were selected for further functional analysis: BG16B\#25.1-1 and BG16A\#25.1-1.x expressing PrpS\textsubscript{1}-GFP, and HZ3.1-3.1, HZ3.1-15.1 and HZ3.1-15.1-x expressing PrpS\textsubscript{3}-GFP. Lines segregated 3:1 with respect for the Kan resistance marker in heterozygous state and were expected to segregate 50 \% for GFP expression. When they reached homozygous state they were completely resistant to Kan and exhibited 100 \% GFP expression. Transgene mRNA expression was analysed using RT-PCR and the above selected lines were selected for their high expression.

Pollen from transgenic \textit{A. thaliana} expressing \textit{Papaver} SI pollen \textit{S}-determinant, PrpS, was used in functional analysis. The semi-\textit{in vitro} SI approach was adopted from the poppy SI system. \textit{A. thaliana} pollen expressing PrpS\textsubscript{1}-GFP or PrpS\textsubscript{3}-GFP germinated in the petri dish to which control or SI reactions were induced by the addition of recombinant stigmatic PrsS proteins (see Figure 5.5). Control reactions were comprised of untreated pollen, pollen that was treated with heat denatured (i.e. biologically inactive) incompatible stigmatic proteins and compatible stigmatic proteins, while incompatible reaction was comprised of incubating
PrpS₁-GFP and PrpS₃-GFP expressing *A. thaliana* pollen with PrsS₁ and PrsS₃ stigmatic recombinant proteins, respectively (see Figure 5.5).

**Figure 5.5.** Schematic diagram of the SI experiment *in vitro*. If PrsS₁ or PrsS₃ recombinant proteins are applied to Arabidopsis pollen expressing PrpS₁-GFP or PrsS₃-GFP (right side, red outline) there is the genetic match between pistil and pollen determinant and the result is incompatible interaction with inhibited pollen tube growth. Control reactions (left side, outlined with green) were comprised of untreated pollen, grown in GM only, of pollen incubated with inactivated heat denatured incompatible PrsS proteins or with compatible PrsS proteins. Compatible control reactions resulted in growth of pollen tubes.

The aim was to investigate whether characteristic features from *Papaver* SI could be observed in *A. thaliana* pollen expressing PrpS-GFP. Therefore, following SI induction *in vitro* the *S*-specific inhibition of pollen tube growth, the alterations of actin cytoskeleton, pollen viability and PCD were examined in PrpS-GFP expressing *A. thaliana* pollen (Geitmann et al., 2000, Thomas and Franklin-Tong, 2004, Jordan et al., 2000). A preliminary investigation was conducted on the response of *A. thaliana* mesophyl protoplasts using line BG3 expressing 35S::PrpS₁ and Col-0 transformed with 35S:: PrpS-GFP upon the incubation with recombinant stigmatic PrsS proteins.

### 5.2.2.1. *S*-specific inhibition of transgenic *A. thaliana* pollen tube growth

The inhibition of transgenic pollen tubes was analysed upon the induced SI response in pollen of transgenic *A. thaliana* lines BG16B#25.1-1 expressing PrpS₁-GFP, and HZ3.1-15.1 expressing PrpS₃-GFP. *A. thaliana* BG16B#25.1-1 and BG16B#25.1-2, and HZ3.1-15.1 and
HZ3.1-3.1 pollen along with untransformed control and *Papaver* S₁ and S₃ control pollen were pre-germinated for 1 h prior to the induction of SI and controls. Analysis was conducted by measuring the length of pollen tubes after overnight incubation (details in Materials and Methods section). *Papaver* S₁S₃ pollen was used as a control and was incubated in GM only (untreated – UT) or had induced SI reaction. *A. thaliana* untransformed pollen, also used as a control, was incubated in GM (UT) or with stigmatic recombinant PrsS₁ or PrsS₃ proteins. *A. thaliana* transgenic pollen expressing PrpS₁-GFP or PrpS₃-GFP was incubated in GM only (UT), had induced SI reaction by applying active PrsS₁ recombinant stigmatic proteins to PrpS₁-GFP pollen or PrsS₃ proteins to PrpS₃-GFP expressing pollen, while the control reactions comprised incubation with heat denatured (hd) PrsS proteins and compatible PrsS proteins. Pollen was pre-germinated for 1 h before the SI or controls were induced, as described above, and was then left to incubate for 20 h (overnight). Results of pollen tube length measurements are presented in Figure 5.6.
Untreated and SI *Papaver* pollen were used as controls. The average length of the *Papaver* pollen tubes grown in GM was 883±168 μm while the length of pollen tubes with induced SI was 40±13 μm, which was about 96 % shorter (N=3, n=20 pollen tubes, P = 1.2x10^{-43}, ***).

The untreated *A. thaliana* (UT) pollen tubes reached average length of 753±263 μm and were 15 % shorter compared to the poppy control pollen tube length (N=6; n=20 tubes; P=5.2x10^{-3}; **). However, when Col-0 pollen was treated with recombinant stigmatic PrsS₁ (758±307 μm; P=0.87; N.S.) or PrsS₃ (759±272 μm; P=0.89; N.S.), they had no effect compared to the untreated pollen grown in GM only.

BG16B#25.1-1 pollen, which expresses PrpS₁-GFP fusion protein, grown in GM only was not significantly different compared to the untreated Col-0 control (737±245 μm; P=0.73; N.S.) but when it was challenged with incompatible PrsS₁ proteins (see Figure 5.6, yellow bars outlined with red line), the pollen tube length exhibited 96 % decrease compared to the
BG16 UT (30±18 μm; P=5.2x10^{-66}; ***). To determine whether the PrsS₁ protein without its biological activity could induce this decrease, we inactivated PrsS₁ by heat denaturing for 5 min. The pollen tube length of BG16B#25.1-1 treated with biologically inactivated PrsS₁ was not significantly different from the UT BG16 pollen and was 96 % longer than the pollen treated with biologically active PrsS₁ (721±291 μm; P=0.75, N.S. compared to BG16-UT and P=7.3x10^{-58}, *** compared to BG16-PrsS₁). This indicates that biologically active PrsS₁ was required for the inhibition of pollen tube growth. To assess whether any active PrsS protein was able to inhibit the pollen tube of transgenic A. thaliana pollen expressing PrpS₁-GFP or if the inhibition was S-specific, the BG16B#25.1-1 pollen was incubated with PrsS₃ proteins. The resulting length of pollen tubes indicate that the PrsS₃ proteins had no effect on the BG16 pollen tubes since they were not significantly different to the BG16-UT and highly significantly longer than the BG16 pollen incubated with incompatible PrsS₁ (740±301 μm; P=0.95, N.S. compared to BG16-UT and P=4.1x10^{-58}, *** compared to BG16-PrsS₁). This demonstrates that PrpS₁-GFP expressing A. thaliana pollen was inhibited in an S-specific manner.

The experiment was repeated with HZ3.1-15.1 line, whose pollen expresses PrpS₃-GFP fusion protein. When HZ3.1-15.1 pollen was grown in GM, it was not significantly different compared to the Col-0 UT (723±318 μm; P=0.58; N.S.). To examine the SI inhibition of pollen tube growth, the PrpS₃-GFP pollen was challenged with PrsS₃ stigmatic proteins (Figure 5.6 green bar outlined with red line) and the pollen was 85 % shorter compared to the HZ3 UT (108±55 μm; P=5.5x10^{-47}; ***). HZ3.1-15.1 pollen, treated with heat-denatured (biologically inactive) PrsS₃ protein was not significantly different from UT pollen (736±299 μm; P=0.81, N.S. compared to HZ3-UT), but was significantly longer than the SI challenged
pollen, treated with biologically active PrsS₃ (P=7.6x10⁻⁵⁴, *** compared to HZ3+PrsS₃). The HZ3 pollen was also incubated with PrsS₁ proteins and the resulting length of pollen tubes indicates that the PrsS₁ proteins had no effect on the HZ3 pollen tubes. Pollen tubes incubated with PrsS₁ were not significantly different to the HZ3-UT and highly significantly longer to the HZ3 pollen incubated with incompatible PrsS₃ (746±314 μm; P=0.66, N.S. compared to HZ3-UT and P=4.7x10⁻⁵³, *** compared to HZ3-PrsS₃). This demonstrates that PrpS₃-GFP expressing A. thaliana pollen was inhibited in an S-specific manner.

Together, these data demonstrate that the inhibition of the PrpS₁-GFP expressing pollen tube growth in selected BG16 lines by recombinant PrsS₁ proteins or PrpS₃-GFP expressing pollen in selected HZ3 lines by recombinant PrsS₃ proteins acts in an S-specific manner and requires biologically active PrsS₁ or PrsS₃ proteins, respectively. Pollen tube inhibition assays that were carried on transgenic A. thaliana pollen expressing PrpS₁-GFP or PrpS₃-GFP proteins challenged with PrsS₁ or PrsS₃ recombinant proteins show that inhibition occurred very rapidly. Inhibited incompatible A. thaliana pollen tubes were very short which was expected and pollen tube inhibition was complete. With Papaver SI the arrest of the “incompatible” pollen tube growth is one of the first events upon the interaction of PrpS and incompatible PrsS protein, which is crucial for successful prevention of fertilization.

This S-specific inhibition response of transgenic A. thaliana pollen expressing Papaver PrpS proteins by PrsS proteins suggests that poppy male S-determinant, PrpS, is functional in A. thaliana.
5.2.2.2. PrpS-PrsS interaction stimulates formation of punctate actin foci in transgenic *A. thaliana* pollen

The results in previous section 5.4.2 confirmed that the Arabidopsis transgenic pollen expressing *Papaver* PrpS-GFP was inhibited in an *S*-specific manner by the addition of the recombinant stigmatic PrsS proteins. The formation of the punctate actin foci is one of the hallmark features of SI response in *Papaver* as actin cytoskeleton is a very early target for the SI signals (Snowman et al., 2002, Geitmann et al., 2000, Snowman et al., 2000, Poulter et al., 2010a). Therefore, in order to provide further evidence that the observed inhibition of *A. thaliana* pollen tube length was due to an SI response, we explored other key poppy SI-like events in *A. thaliana* pollen. We investigated whether actin filament modification was also observed in *A. thaliana* PrpS-GFP expressing pollen when treated with PrsS recombinant proteins.

We initiated the analysis of actin cytoskeleton in pollen of the transgenic *A. thaliana* lines BG16B#25.1-1 expressing PrpS1-GFP, and HZ3.1-15.1 expressing PrpS3-GFP, after 3 h SI challenge. BG16B#25.1-1 was incubated with recombinant stigmatic PrsS1 proteins and HZ3.1-15.1 with PrsS3 proteins. Controls consisted of untreated (UT) Col-0 and transgenic pollen; transgenic *A. thaliana* pollen treated with heat denatured (hd) recombinant proteins or compatible recombinant proteins, as with previous assay.

Untransformed *A. thaliana* Col-0 pollen, grown in GM only, had a visible array of F-actin filaments in pollen grains and pollen tubes (Figure 5.7.a) and the results also demonstrate that actin filaments in nontransgenic Col-0 pollen were not disturbed upon the addition of recombinant PrsS1 (Figure 5.7.b).
Filamentous actin arrays were also observed in the unchallenged pollen of transgenic line BG16 expressing PrpS1-GFP (Figure 5.7.c) or HZ3 expressing PrpS3-GFP (Figure 5.7.h). Characteristic punctate actin foci were observed in BG16 pollen that was incubated with PrsS1 proteins (Figures 5.7.d&e) and HZ3 pollen challenged with PrsS3 proteins (Figure 5.7.i). Foci were not present in any of the controls where incompatible and heat inactivated PrsS proteins were added (Figures 5.7.f &j) or when compatible PrsS control was applied (Figures 5.7.g & k). Taken together these data demonstrate that *A. thaliana* PrpS1-GFP and PrpS3-GFP expressing pollen exhibit the S-specific appearance of punctate actin foci, a hallmark feature of poppy SI.

*A. thaliana* BG16B#25.1-1 pollen expressing PrpS1-GFP was examined for the presence of actin filaments or punctate actin foci (see Figure 5.8) in order to quantify the S-specific appearance of foci observed under the microscope. Figure 5.8.a clearly shows that actin
filaments (blue bars) are predominant in all control treatments, while SI challenge on PrpS1-GFP expressing pollen exhibit mostly punctate actin foci (red bars). Taking into account only scored pollen with clearly visible actin structures, 88 % of the untreated non transgenic Col-0 pollen samples exhibit filamentous actin, which was not affected if Col-0 pollen was challenged with PrsS1 proteins (85 % filaments; P=0.46). Scoring filaments, BG16B#25.1-1 untreated pollen exhibited 88 % actin filaments (P=0.97 compared to Col-0 untreated; N=250), while BG16B#25.1-1 that was incubated with biologically inactivated PrsS1 or compatible PrsS8 exhibited 89 % actin filaments (P=0.78 and P=0.85, respectively, compared to BG16 untreated, N=250), indicating that biologically active PrsS1 only are required for the alterations in actin cytoskeleton. Quantification also revealed that SI challenge of BG16 with PrsS1 resulted in 81 % of pollen exhibiting punctate actin foci (Figure 5.8.c), which represents a highly significant 558 % increase compared to untreated BG16 pollen (P=8.9x10^{-45}; N=250).
Figure 5.8: Quantification of actin foci and filaments in PrpS\textsubscript{1}-GFP expressing pollen. (a) actin foci (red bars) and filaments (blue bars) presented on the same graph; (b) quantification of actin filaments; (c) quantification of actin foci. N=5, n=50; Error bars represent ± St. Error of means.

None of the non-transgenic controls or BG16 controls with inactivated incompatible proteins or compatible proteins showed actin foci pattern and were not significantly different from the untreated pollen in which foci are mostly observed. Together these data demonstrate the highly significant presence of punctate actin foci, specifically in SI challenged transgenic \textit{A. thaliana} BG16B\#25.1-1 pollen, expressing PrpS\textsubscript{1}-GFP. These data demonstrate that there is an \textit{S}-specific interaction between biologically active PrsS\textsubscript{1} and PrpS\textsubscript{1}-GFP, leading to the alterations of actin cytoskeleton in PrpS\textsubscript{1}-GFP expressing \textit{A. thaliana} pollen, that result in the appearance of punctate actin foci. Therefore a key characteristic of poppy SI was observed in transgenic \textit{A. thaliana} pollen, indicating that PrpS\textsubscript{1}-GFP transgene is functional in \textit{A. thaliana}.  

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5.2.2.3. Evidence for PCD in the *A. thaliana* transgenic pollen upon SI challenge

5.2.2.3.1. Pollen viability is decreased in an S-specific manner upon interaction between PrpS-GFP expressing *A. thaliana* pollen and recombinant stigmatic PrsS

We were interested to examine whether the interaction between *Papaver* pollen and pistil determinants affected Arabidopsis pollen viability. It has been shown in *Papaver* that the S-specific interaction between PrpS and PrsS decreases pollen viability and triggers PCD (Thomas and Franklin-Tong, 2004). The pollen viability in *Papaver* was analysed using FDA, which becomes fluorescent when taken up by metabolically active–viable–pollen (Breeuwer et al., 1995). However, we were unable to use the same drug in *A. thaliana* as it was not possible to distinguish between FDA signal of viable pollen and GFP signal of pollen expressing PrpS-GFP fusion protein. Therefore, we used the low toxicity non-permeating azo-compound dye Evans Blue (Shigaki and Bhattacharyya, 1999). Dye penetrated in non-viable cells, so they appear dark blue when inspected by the microscope, while it was excluded from viable cells with intact membrane so they appeared colourless.

Lines of PrpS1-GFP and PrpS3-GFP expressing *A. thaliana* pollen (BG16B#25.1-1 and BG16B#25.1-2, and HZ3.1-15.1 and HZ3.1-3.1) were used as for measuring pollen tube length and pollen was collected and treated as with previous experiments. Viability of pollen was inspected at three time points: 0 h, 8 h and 2 4h by adding 0.05 % Evans Blue to the sample. Dye was left to incubate for 10 min and washed with GM before pollen viability was assessed (see Figure 5.9). In poppy SI DNA fragmentation indicating PCD was measured after 8 h (Thomas and Franklin-Tong, 2004) although the maximum DNA fragmentation was observed at 14 h post SI (Jordan et al., 2000). *A. thaliana* pollen tubes
grew slower than *Papaver* with the optimized germination rates reported after 16 h, so we chose to investigate viability at 24 h post-SI.

![Graph showing viability (%) of A. thaliana transgenic pollen BG16 (yellow bars) and HZ3 (green bars) along with Col-0 (dark gray bars) and *Papaver* pollen control (light gray bars) at 0 h, 8 h and 24 h. Pollen was incubated with GM only (UT), with incompatible PrsS1 proteins for BG16 pollen or PrsS3 proteins for HZ3 pollen (SI, red outline), with heat-denatured PrsS proteins and with compatible PrsS1 for BG16 pollen and PrsS3 for HZ3 pollen. Error bars represent standard deviation of mean. N=5 repeats (N=3 for poppy), n=117±21 pollen counted.]

*Figure 5.9*: Viability (%) of *A. thaliana* transgenic pollen BG16 (yellow bars) and HZ3 (green bars) along with Col-0 (dark gray bars) and *Papaver* pollen control (light gray bars) at 0 h, 8 h and 24 h. Pollen was incubated with GM only (UT), with incompatible PrsS1 proteins for BG16 pollen or PrsS3 proteins for HZ3 pollen (SI, red outline), with heat-denatured PrsS proteins and with compatible PrsS1 for BG16 pollen and PrsS3 for HZ3 pollen. Error bars represent standard deviation of mean. N=5 repeats (N=3 for poppy), n=117±21 pollen counted.

The initial viability of the untreated *A. thaliana* non-transgenic Col-0 pollen at t₀ was 89±4 % (P=3.6x10⁻² compared to poppy UT). The viability of Col-0 pollen decreased by 17 % after 8 h. The PrsS proteins did not have any effect on the viability of untransformed pollen, viability of pollen treated with PrsS₁ for 8 h was 76±11 % (P=0.71 compared to Col-0 t=8 h) and viability of Col-0 treated with PrsS₃ for 8 h was 77±9 % (P=0.83 compared to Col-0 t=8 h).
In control *Papaver* pollen from $S_1S_3$ plants, the incubation with incompatible PrsS$_1$PrsS$_3$ proteins for 8 h resulted in highly significant 90 % decreased viability compared to the *Papaver* pollen that was incubated in GM for 8 h ($P=4.3\times10^{-19}$ compared to *Papaver* $S_1S_3$ UT $t=8$ h). With longer incubation time (24 h total), the viability of the control pollen decreased. Untreated $S_1S_3$ *Papaver* pollen exhibited 59±5 % viability, which is 39 % less than at 8 h but viability of $S_1S_3$ *Papaver* pollen incubated with PrsS$_1$ and PrsS$_3$ proteins did not significantly change compared to the 8h incubation 6±1 % ($P=2.3\times10^{-51}$ compared to *Papaver* $S_1S_3$ UT; ***). *A. thaliana* untreated non-transgenic control Col-0 pollen exhibited 50±11 % viability after 24 h incubation. The viability of Col-0 incubated with PrsS$_1$ or PrsS$_3$ for 24 h was not significantly different from the untreated Col-0 ($P=0.72$ and $P=0.78$, respectively).

Viability of transgenic BG16 pollen (expressing PrpS$_1$-GFP) grown in GM only was initially around 88±2 % ($P=0.98$ compared to Col-0 UT $t_0$; N.S.). After 8 h, the viability decreased for 16 % and reached 75±9 % ($P=0.83$ compared to Col-0 UT $t=8$h). When BG16 pollen was challenged with PrsS$_1$ proteins for 8 h, a significant decrease in viability to 30±9 % ($P=2.9\times10^{-10}$ compared to BG16 UT $t=8$ h) was observed. The heat-denatured PrsS$_1$ protein had no effect on BG16 pollen after 8 h incubation (viability 70±11 %; $P=0.50$ compared to BG16 UT $t=8$ h) nor did active PrsS$_3$ protein (viability 70±11 %; $P=0.48$ compared to BG16 UT $t=8$ h). The viability of BG16 pollen that was incubated overnight in the GM decreased by 29 % and reached 53±6 % of viability (total decrease in 24 h was 40 %) but was not significantly different from the Col-0 UT control at the same incubation time ($P=0.64$). Significant decrease in viability was observed when BG16 pollen was incubated with PrsS$_1$ proteins for 24 h with 14±2 % of viability ($P=6.9\times10^{-9}$ compared to the BG16 UT $t=24$ h). The decrease in viability compared to the untreated BG16 control was 73 % and 53 %
compared to the BG16 pollen incubated with PrsS$_1$ proteins for 8 h only. This could suggest that the cell death response in *A. thaliana* pollen is somewhat slower compared to *Papaver* pollen. The response was *S*-specific and observed only with active PrsS$_1$ proteins as incubation of BG16 with heat-denatured PrsS$_1$ or PrsS$_3$ for 24 h had no significant effect on pollen compared to the untreated BG16 control (P=0.75 and P=0.69, respectively).

Viability of PrpS$_3$-GFP expressing transgenic pollen HZ3 exhibited similar results to the observed from BG16. HZ3 grown in GM exhibited an initial viability of 89±2 % (P=0.92 compared to Col-0 UT t=0; N.S.). After 8 h, the viability decreased by 24 % and reached 69±7 % (P=0.40 compared to Col-0 UT t=8 h). The incubation of HZ3 pollen with recombinant PrsS$_3$ proteins for 8 h caused a significant decrease of viability to 36±14 % (P=6.5x10$^{-6}$ compared to HZ3 UT t=8 h). Only active PrsS$_3$ protein had an effect on the viability of HZ3 pollen. After 8 h incubation with heat-denatured PrsS$_3$ protein, the viability was 72±9 % (P=0.53 compared to HZ3 UT t=8 h). No effect on viability was observed when the HZ3 pollen was incubated in the presence of active PrsS$_1$ proteins that decreased viability of BG16 pollen (viability 69±10 %; P=0.83 compared to HZ3 UT t=8 h). After overnight incubation in GM the viability of HZ3 pollen decreased for further 26 % and reached 50±6 % (total decrease in 24 h was 44 %) but was not significantly different from the Col-0 UT control at t=24 h (P=0.93). With incompatible reaction between HZ3 pollen and recombinant PrsS$_3$ proteins, the significant decrease in viability was observed after 24 h incubation as it reached 21±6 % (P=2.2x10$^{-5}$ compared to the HZ3 UT t=24 h). The decrease in viability compared to the untreated HZ3 pollen incubated in GM for 24 h was 57 %, while further decrease compared to incompatible incubation of HZ3 pollen with PrsS$_3$ for 8 h was 40 %.

As with BG16 pollen, the response was *S*-specific and observed only with active PrsS$_3$
proteins. The incubation of HZ3 with heat inactivated PrsS3 or active PrsS1 for 24 h had no significant effect on pollen compared to the HZ3 control grown in GM only (P=0.75 and P=0.69, respectively).

The SI-induced decrease of viability in pollen from BG16B#25.1-1 line expressing PrpS1-GFP and HZ3.1-15.1 line expressing PrpS3-GFP, by the addition of the recombinant stigmatic PrsS1 or PrsS3, respectively, indicates that the S-specific interaction between PrsS and PrpS is functional and results in cell death of transgenic pollen.

5.2.2.3.2. Decrease of pollen viability involving DEVDase activity

A key downstream event after the S-specific interaction between the stigmatic PrsS proteins and pollen receptor PrpS in *Papaver*, is the activation of the caspase-like proteins ending up with the PCD of pollen (Thomas and Franklin-Tong, 2004). The PCD in *Papaver* was demonstrated with assays measuring DNA fragmentation, and with the use of caspase-3 specific tetrapeptide inhibitor Ac-DEVD-CHO, which abolished PCD (Thomas and Franklin-Tong, 2004, Jordan et al., 2000). Hence, we investigated whether DEVD-ase activity plays any role in the S-specific decrease of viability observed in transgenic *A. thaliana* lines BG16B#25.1-1.1, BG16B#25.1-1.5 and BG16B#25.1-1.7, expressing PrpS1-GFP and HZ3.1-15.1-1, HZ3.1-15.1-6 and HZ3.1-15.1-12, expressing PrpS3-GFP. Transgenic *A. thaliana* homozygous pollen from T4 generation plant lines was used for the experiment. *A. thaliana* plants from lines BG16B#25.1-1.1 and BG16B#25.1-1.7 were used as a source for PrpS1-GFP pollen, and lines HZ3.1-15.1-1 and HZ3.1-15.1-5 for PrpS3-GFP pollen. PrpS1-GFP and PrpS3-GFP expressing *A. thaliana* pollen, and controls *Papaver* S1S3
pollen and *A. thaliana* untransformed Col-0 pollen were pretreated with the inhibitor of DEVDase activity, tetrapeptide Ac-DEVD-CHO, before the induction of SI by the addition of recombinant stigmatic PrsS proteins. Additional control was comprised of pollen incubated in GM only (untreated – UT) or pollen that had induced SI reaction. Viability of pollen was investigated at two time points: 0h and 8h by adding 0.05 % Evans Blue solution to the sample. The results of these assays are presented in the Figure 5.10. When comparing percentages of viability between treatments, the control was assumed a 100 % activity.

*A. thaliana* non-transgenic pollen germinated in GM (UT) exhibited 90±4 % viability at t₀ and was only 1 % higher than viability of untreated *Papaver* pollen at t₀ (P=0.79 compared to poppy UT). After 8 h incubation the viability of Col-0 pollen decreased by 28 % having 65±4 % of viable pollen. The pretreatment with tetrapeptide Ac-DEVD-CHO did not have any effect on the non-transgenic pollen; viability was 62±6 % (P=0.66 compared to Col-0 UT pollen t=8 h). PrsS proteins did not have any effect on the viability of the Col-0 control pollen, viability of pollen treated with PrsS₁ for 8 h was 65±4 % (P=0.95 compared to Col-0 UT t=8 h) and viability of Col-0 treated with PrsS₃ for 8 h was 66±6 % (P=0.98 compared to Col-0 UT t=8 h).
The viability of untreated control S1S3 *Papaver* pollen decreased after 8 h incubation to 76±2 % while pollen that was pretreated with the caspase inhibitor Ac-DEVD-CHO for an hour and then incubated for 8 h exhibited 69±4 % viability (P=0.29 compared to S1S3 *Papaver* UT control t=8 h). The incompatible challenge on *Papaver* pollen decreased the viability to 6±2 % (P=1.3×10^{-23} compared to *Papaver* S1S3 UT t=8 h). However, viability was not lost when pollen was pretreated with caspase-3 inhibitor Ac-DEVD-CHO before induction of SI but significantly increased viability that was 43±18 % (P=1.6×10^{-9} compared to S1S3 *Papaver* SI t=8 h).

Initial viability of transgenic pollen expressing PrpS1-GFP grown in GM only was around 90±4 % (P=0.98 compared to Col-0 UT t0). After 8h, the viability decreased by 20 % and
was 72±8 % (P=0.28 compared to Col-0 UT t=8 h). The pretreatment of BG16 pollen with tetrapeptide Ac-DEVD-CHO did not significantly affect the viability (69±4 %, P=0.60 compared to BG16 UT t=8 h). SI challenge of BG16B#25.1-1.1 and BG16B#25.1-1.7 pollen with PrsS1 proteins for 8 h caused a significant decrease of viability to 22±11 % (P=5.3x10^{-13} compared to BG16 UT t=8 h), but viability increased significantly by 162 % when PrsS1-challenged BG16 pollen was pretreated with inhibitor Ac-DEVD-CHO (57±4 %; P=4x10^{-7} compared to BG16 SI t=8 h).

Similar viabilities were observed in HZ3.1-15.1-1 and HZ3.1-15.1-5 pollen expressing PrpS3-GFP. Initial viability of HZ3 pollen was 91±4 % (P=0.88 compared to Col-0 UT t0). After 8 h, the viability decreased for 22 % and reached 71±6 % (P=0.39 compared to Col-0 UT t=8 h). When HZ3 pollen was pretreated with tetrapeptide Ac-DEVD-CHO, the viability was not significantly affected (63±10 %, P=0.23 compared to HZ3 UT t=8 h). Challenge of PrpS3-GFP expressing pollen with stigmatic recombinant PrsS1 proteins for 8 h caused a significant decrease of viability to 21±9 % (P=7.8x10^{-13} compared to HZ3 UT t=8 h); however, viability increased significantly by 155 % if SI challenged HZ3 pollen was pretreated with inhibitor Ac-DEVD-CHO (52±2 %; P=3x10^{-6} compared to HZ3 SI t=8 h).

Taken together, these data demonstrate that PrsS triggers cell death in PrpS-GFP expressing pollen in an S-specific manner. Moreover, the cell death involves DEVDase activity and indicates that PCD might be triggered in incompatible A. thaliana pollen expressing PrpS1-GFP or PrpS3-GFP fusion protein when incubated with the cognate recombinant stigmatic PrsS proteins.
5.2.2.4. PCD in *A. thaliana* transgenic pollen expressing PrpS-GFP

As described in previous section, the viability of transgenic *A. thaliana* pollen expressing male PrpS$_1$-GFP or PrsS$_3$-GFP was reduced in an $S$-specific manner upon treatment with the recombinant stigmatic PrsS proteins. In order to ascertain whether the poppy SI response is fully functional in *A. thaliana*, analysis of PCD was addressed. Therefore, we initiated studies to analyse SI-induced PCD in transgenic *A. thaliana* pollen expressing PrpS$_1$-GFP and PrpS$_3$-GFP by attempting to measure caspase-3-like DEVDase activity more directly using the caspase-3 specific fluorogenic tetrapeptide Ac-DEVD-AMC. The stimulation of several specific caspase-like activities was demonstrated using caspase activity assays (Bosch and Franklin-Tong, 2007). They were based on measuring the release of the fluorophore from the specific peptide substrate tetrapeptide, e.g. caspase-3 optimal tetrapeptide recognition motif is DEVD. Substrate used in the fluorometric assay has DEVD conjugated to the fluorescent probe AMC (Ac-DEVD-AMC). The fluorescence can be detected upon cleavage of the AMC from the DEVD peptide by caspase-3-like proteins. Assays measuring caspase-like activity in *Papaver* pollen were optimised to use 10 $\mu$g of the total pollen protein extract in the assay. Due to the miniature size of *A. thaliana* flowers which produce less pollen compared to *Papaver* flowers and the fact that pollen required fresh collection every day, preliminary experiments were carried out on *A. thaliana* pollen, in order to investigate whether the caspase activity could be reliably measured with less than 10 $\mu$g of total *A. thaliana* pollen protein extract. In order to use the same concentration of pollen protein extract as it was used with *Papaver* pollen, the method was not suitable for *A. thaliana* pollen, as the pollen collection from *A. thaliana* flowers was very time consuming.
Figure 5.11 shows initial preliminary data using 3, 4 and 5 μg protein extract that encouraged us to pursue this assay further.

Figure 5.11: DEVDase activity of protein extract from BG16B#25.1-1 and HZ3.1-15.1 pollen that was incubated with GM only (UT), or had induced SI by addition of recombinant stigmatic PrsS₁ (BG16 S1) and PrsS₃ (HZ3 S3) proteins, respectively. Non-transgenic Col-0 incubated in GM only (UT) was used as a control (far left bar). (a) 5 μg of pollen extract; (b) 4 μg of pollen extract and (c) 3 μg of pollen extract.

Preliminary results in Figure 5.11 suggested that DEVDase activity in SI-induced samples of transgenic A. thaliana pollen expressing PrpS₁-GFP and PrpS₃-GFP was detected and
increased compared to the control sample, even when using the substrate in concentration as low as 3 μg. Since the protein concentration of our loaded pollen extract in the assay was on the lower limit of detection, this means that any changes in the very sensitive *A. thaliana* pollen balance appeared enhanced in the caspase assay.

In order to examine whether SI interaction in *A. thaliana* transgenic pollen triggered PCD, we used another approach and pretreated pollen with the tetrapeptide inhibitor Ac-DEVD-CHO before the induction of SI and measurement of caspase-like activity. *A. thaliana* transgenic homozygous T₄ generation plant lines BG16B#25.1-1.1 and BG16B#25.1-1.7 were used as a source for PrpS₁-GFP pollen, and lines HZ3.1-15.1-1 and HZ3.1-15.1-5 for PrpS₃-GFP pollen; pollen from untransformed Col-0 was used as control. Pollen was pretreated with 100 μM tetrapeptide inhibitor, Ac-DEVD-CHO and SI was induced by the addition of incompatible recombinant stigmatic PrsS proteins: PrsS₁ proteins added to PrpS₁-GFP expressing pollen and PrsS₃ proteins to PrpS₃-GFP expressing pollen. Controls comprised of transgenic and untransformed pollen that was untreated, incubated with the inhibitor Ac-DEVD-CHO only or incubated with PrsS₁ or PrsS₃ recombinant stigmatic proteins.

The caspase-3-like activity measurements are presented in Figure 5.12.
Figure 5.12: DEVDase activity in *A. thaliana* pollen with induced SI or pretreated with inhibitor Ac-DEVD-CHO prior to induction of SI. Grey bars correspond to untransformed Col-0 control that was untreated or incubated with Ac-DEVD-CHO or stigmatic PrsS₁ and PrsS₃ proteins. BG16B#25.1-1.1 and BG16B#25.1-1.7 pollen (represented by yellow bars) was challenged with incompatible PrsS₁ (BG16 S1) and was also pretreated with inhibitor Ac-DEVD-CHO prior to induction of SI (BG16 DEVD-CHO S1). Control was comprised of untreated BG16 pollen (BG16 UT) and BG16 pollen incubated with inhibitor only (BG16 DEVD-CHO). HZ3.1-15.1-1 and HZ3.1-15.1-5 pollen (represented by green bars) was challenged with incompatible PrsS₁ (HZ3 S3) and was also pretreated with inhibitor Ac-DEVD-CHO prior to induction of SI (HZ3 DEVD-CHO S3). Control was comprised of untreated HZ3 pollen (HZ3 UT) and HZ3 pollen incubated with inhibitor only (HZ3 DEVD-CHO). Error bars are ± SEM.

The cleavage of Ac-DEVD-AMC by SI-activated caspase-3-like proteins in *A. thaliana* transgenic pollen was significantly increased by 150±52 % in BG16 PrsS₁ treated pollen (yellow bar, P=0.04, *) and by 106±23 % in HZ3 PrsS₃ treated pollen (green bar, P=0.003, **). Pretreatment of pollen tubes with the tetrapeptide inhibitor significantly reduced the SI-induced DEVDase activity in BG16 pollen by 87±12 % (P=0.009, **) and by 62±23 % in HZ3 pollen (P=0.003, **). As shown in Figure 5.12, control treatment of *A. thaliana* pollen with tetrapeptide inhibitor Ac-DEVD-CHO had no significant effect on the DEVDase activity of pollen (P=0.90 for Col-0 pollen, grey bars; P=0.67 for BG16 pollen, yellow bars and P=0.63 for HZ3 pollen, green bars).
These data together presents the first evidence for activation of a caspase-3-like activity in PrpS₁-GFP and PrpS₃-GFP expressing *A. thaliana* pollen upon induction of SI. The PrsS induced PCD demonstrates that PrpS₁-GFP and PrpS₃-GFP are functional in *A. thaliana*.

We also wished to characterize the DEVDase activity by adding the tetrapeptide inhibitor Ac-DEVD-CHO to the pollen extracts in the assay to see if we could bring down the levels of DEVDase activity. SI was stimulated in *A. thaliana* transgenic pollen PrpS₁-GFP and PrpS₃-GFP and untransformed Col-0 control by the addition of stigmatic recombinant PrsS proteins. Following 5 h incubation, when maximal DEVDase activity was exhibited in *Papaver* pollen (Bosch and Franklin-Tong, 2007), the *A. thaliana* pollen protein extracts were made. To test the effect of caspase inhibitor on the substrate cleavage, 50 μM peptide inhibitor Ac-DEVD-CHO inhibitor was added to the assay and DEVDase activity was measured (Figure 5.13).

**Figure 5.13.** Effect of DEVDase inhibitor on the SI induced caspase-like activity. Light grey bars present DEVDase activity in *A. thaliana* BG16 #25.1-1 and HZ3.1-15.1 pollen challenged with PrsS₁ and PrsS₃ proteins, respectively (SI challenge outlined by red line). Non-transgenic Col-0 pollen with the same challenge was used as a control. Dark grey bars show DEVDase activity of pollen extract with the additions of Ac-DEVD-CHO inhibitor in the assay. N=3 repeats. Error bars present ± St. Error of means.
Results in Figure 5.13 show the increase of DEVDase activity by 66% in PrpS1-GFP expressing pollen challenged with PrsS1 and increase by 175% in PrpS3-GFP expressing pollen challenged with PrsS3. Despite the increase, the results of this preliminary experiment were not significantly different from the untreated controls (P = 0.47 for PrpS1 and 0.21 for PrpS3). However, when the inhibitor Ac-DEVD-CHO was used in the assay, the activity was reduced, indicating the activated caspase-3-like DEVDase activity. DEVDase activity was decreased significantly with the addition of the inhibitor Ac-DEVD-CHO in PrpS1-GFP and PrpS3-GFP expressing pollen that had SI induced, indicating that the PrsS-PrpS interaction stimulates high DEVDase activity in transgenic A. thaliana pollen.

As our data indicate, DEVDase activity could be decreased, however, the results were not significant. Experiment was repeated only three times so it might be possible that with more repeats and higher protein concentration we could observe significant response.

### 5.2.3. *A. thaliana* mesophyll protoplasts expressing PrpS1

The previous sections demonstrated that PrpS1-GFP and PrpS3-GFP were functional in pollen of transgenic *A. thaliana*. Preliminary experiments were conducted to explore whether *Papaver* PrpS1 receptor could be functional in other tissues as well. This would demonstrate that for a functional SI response, just PrpS and PrsS were required, while the downstream components required for the characteristic SI response were already present in the host cell.

These initial preliminary studies were done using *A. thaliana* mesophyll protoplasts as they offer versatile cell-based system. Initially, *A. thaliana* BG3, stable transformant line was used, expressing PrpS1 under the control of 35S promoter. Using RT-PCR the presence of PrpS1 mRNA was confirmed in three weeks old BG3 seedlings, however, the signal for
expression of the functional protein using western blot was not possible to detect. However, if the protein was expressed than it should be somehow responding to the recombinant PrsS1. We attempted a series of experiments trying to capture the response between two determinants: immunolocalisation, viability and caspase activity assay, but no response was detected. There could be various reasons for this:

1. PrpS1 protein might not be expressed in high enough levels in BG3 protoplasts and young seedlings.
2. If it was expressed it might not be properly folded and therefore not functional.
3. It might be expressed in a very low concentration and it might not be expressed in all the cells of seedlings.

Therefore we decided to use a system of transient DNA expression in A. thaliana non-transgenic Col-0 mesophyll protoplasts. We transfected Col-0 protoplasts with 35S:PrpS1-GFP, so the transfected protoplasts could be visualised by detecting GFP reporter gene and the response to PrsS1 would be anticipated. Following transfection protocol (see materials and methods) we isolated Col-0 protoplasts and transfected them with 35S:PrpS1-GFP. The transfection rates were very low, between 10-19 %, however the GFP signal was observed and detected (see Figure 5.14).

![Figure 5.14: GFP expression in Col-0 protoplasts transformed with 35S:PrpS1-GFP. (a) protoplasts expressing GFP on the right, left is brightfield image; (b) protoplasts with no GFP expression on the image on the right, left represents brightfield image of the same protoplast. The experiment was done together with Dr. Javier-Andres Juarez-Diaz.](image)
Using transformed Col-0 protoplasts we measured the viability of Col-0 35S:PrpS₁-GFP protoplasts that were incubated with recombinant PrsS₁ proteins at three time points: t₀, 24h and 48h. As a control we used Col-0 protoplasts transfected with H₂O. The preliminary experiment was repeated twice and viability was measured using 0.05 % Evans Blue. Results are presented on Figure 5.15.

Figure 5.15: Viability of *A. thaliana* Col-0 protoplasts transfected with 35S:PrpS₁-GFP (red bars) and H₂O control (blue bars), incubated with recombinant PrsS₁ Error bars represent ± St. Deviation. N=2

At t₀ the viability of Col0 - 35S:PrpS₁-GFP and H₂O control was around 80 %. After 24h the average viability of the control was 60±4 % and the viability of PrpS₁-GFP expressing protoplasts was 44±6 %. The viability after 48h was 36±20 % for the control and 18±14 % for the transfected protoplasts, which was 50 % less than the control.

The preliminary experiments indicate the S-specific reduction of the viability of PrpS₁-GFP expressing protoplasts upon PrsS₁ incubation. However, for more reliable and firmer conclusion, the experiments should be repeated several times.

The protoplasts were not difficult to produce but were extremely sensitive to any sort of stress from the environment and often the transformation rates were not high. So in order to
produce high quality protoplasts continuously, the protocol should be optimised further to suit the needs of our experiment (Yoo et al., 2007, Wu et al., 2009). The preliminary work has so far confirmed our hypothesis that PrpS1-GFP could be expressed and potentially functional in other tissue. However, further work would be required on this area to provide strong, conclusive evidence. Experiments for interaction between PrpS1-GFP expressed protoplasts and recombinant PrsS1 proteins should be conducted, such as: examination of actin structures (preliminary experiments pointed at the problem of overlapping red signal of chloroplasts in protoplasts and rhodamine-phalloidin stained actin), changes in the viability, involvement of the caspase-like activity, examination of PCD. Protoplast system could also be used to study interaction between PrpS and PrsS.

5.3 DISCUSSION

The studies presented here demonstrate that the PrpS proteins can be functional when expressed in A. thaliana. Experiments focused on the use of semi-in vitro system by growing transgenic A. thaliana pollen expressing PrpS-GFP and challenging it with incompatible stigmatic recombinant PrsS proteins. Several aspects were examined, which will be discussed here.

5.3.1 Alterations to the actin cytoskeleton

In this study we have established the presence of the punctate actin foci in A. thaliana pollen tube expressing PrpS-GFP after 3 h challenge with incompatible PrsS recombinant protein.
Actin cytoskeleton is of vital importance for normal pollen tube growth. During the SI process in *Papaver rhoeas*, it undergoes through rapid and dramatic alterations (Geitmann et al., 2000, Snowman et al., 2002, Poulter et al., 2008, Poulter et al., 2010a). First changes of actin depolymerisation were observed as early as 1-2 min (Snowman et al., 2002) and after 3 h stable punctate actin foci were observed (Geitmann et al., 2000, Snowman et al., 2002).

Results presented in this study show that controls challenged with compatible or heat denatured PrsS proteins resulted in the same display of bundled actin filaments as untreated BG16 pollen or as non-transgenic pollen. It was also demonstrated that PrsS proteins, that were added to the untransformed pollen had no effect on the actin cytoskeleton. Although some intermediate actin structures (filaments-foci) or no structures at all were also observed, their presence was significantly lower than the filaments. The restriction of the staining technique was mostly in uneven staining of the large heterogeneous pollen population. Although the sample was mixed when the staining (or permeabilization, fixative, or treatment) was added, the pollen settled to the bottom of the tube, and consequently each of the pollen grains and tubes were unevenly exposed to the media. Mixed pollen population was unavoidable, as we needed to collect enough pollen to compensate for its losses during the washing steps of the protocol. Great precaution was taken with the experimental procedure and with collecting the same stage flowers from healthy *A. thaliana* plants.

The cytoskeleton in *A. thaliana* pollen tubes was only disturbed upon the challenge of PrpS1 or PrpS3 expressing pollen with its cognate incompatible PrsS1 or PrsS3 recombinant protein, respectively. Foci were quantified in PrpS1 expressing BG16 pollen where they were present in majority of all analysed pollen. This suggests that functional interaction between biologically active PrsS protein and its cognate PrpS receptor was required for the disruption of the actin cytoskeleton in pollen. Therefore, the punctate actin foci that we observed in the
transgenic *A. thaliana* model upon the interaction between PrpS\(_1\)-GFP and PrsS\(_1\) represents an important landmark in exploring *Papaver* SI system in other species.

The next step with the actin cytoskeleton would be to characterise the temporal dynamics, composition and formation of the punctate actin foci in order to investigate how closely it resembles the well-characterised poppy SI response, where reorganisations are observed as early as 1-2 min post SI. Another aspect of SI induced actin cytoskeleton remodelling, that was not investigated in this study, were the ABPs involved in regulating these modifications, such as CAP and ADF, and were demonstrated and analysed in *Papaver* pollen (Poulter et al., 2010a). It would be also interesting to know whether there was a crosstalk between microtubules and microfilaments (Poulter et al., 2008) and also link between actin depolymerization and PCD (Thomas et al., 2006) should be explored further in incompatible *A. thaliana* pollen.

Our studies, conducted in transgenic *A. thaliana*, a model for *Papaver* SI, reported for the first time, that reorganisations of the actin cytoskeleton occured upon PrpS interaction with PrpS-GFP in *A. thaliana* and the appearance of characteristic punctate actin foci was confirmed. In a self compatible *A. thaliana*, that is a more closely related to Brassicaceae than Papaveraceae, this might seem surprising but it suggests, that just PrpS and PrsS are required for the SI response. However, the recent advances in the research on the SI induced actin reorganisations in different systems suggest, that there might be a parallel between the mechanistically different SI systems and that the SI systems could be more conserved then known so far, which would put our research of two evolutionary distant species into perspective (Liu et al., 2007c, Iwano et al., 2007, Wang et al., 2010, Poulter et al., 2011).
5.3.2. Decrease of viability involving DEVDase activity

Pollen viability was examined in PrpS<sub>1</sub> and PrpS<sub>3</sub> expressing *A. thaliana* pollen, and in non-transgenic pollen and *Papaver* S<sub>1</sub> and S<sub>3</sub> pollen that were used as a control, using Evans Blue dye due to the overlap of the signal from FDA stain and GFP in transgenic pollen tubes. However, many investigations on PCD processes in plants utilize Evans Blue dye as a tool to assess cell death levels, for example, the investigation of the kiss-of-death (KOD)-induced PCD or UV and H<sub>2</sub>O<sub>2</sub> induced PCD in *A. thaliana* (He et al., 2007b, Blanvillain et al., 2011). Evans Blue dye quenches the fluorescent GFP signal, therefore the imaging of Evans Blue staining and corresponding GFP fluorescence of the same pollen tube was not possible. The GFP expression would be very informative, especially initially, with the pollen from the heterozygous lines, where we could directly compare the GFP fluorescence in PrpS-GFP expressing pollen and how it is affected during the interaction between PrpS and PrsS. *Papaver* pollen viability decreased to 7 % as previously reported by (Jordan et al., 2000), while the viability of *A. thaliana* pollen decreased to 30 %, which was significant decrease. The compatible stigmatic PrsS recombinant proteins had no effect on the integrity of transgenic pollen membrane. We also established that only biologically active incompatible PrsS<sub>1</sub> (with BG16 pollen) or PrsS<sub>3</sub> (with HZ3 pollen) proteins can trigger the decrease in the membrane integrity as heat denatured version did not affect pollen viability, which remained same as in untreated pollen. So, the decrease in the membrane integrity of pollen and therefore loss of viability occurred in an S-specific manner upon the interaction between PrpS-GFP and PrsS.

We examined viability after 24 h incubation. Control pollen viability was around 50 %, while the viability of SI induced PrpS<sub>1</sub>-GFP or PrpS<sub>3</sub>-GFP pollen decreased for further 50
%. Lower viability in pollen from homozygous lines was expected earlier, considering rapid inhibition of pollen tubes and homozygous state of pollen used. There can be several possible explanations for this: *A. thaliana* pollen could retain membrane integration and therefore viability for longer period of time, suggesting that the “SI” in *A. thaliana* is acting slower. On the other hand, data, that 50 % of control pollen tubes remained viable for 24 h was not surprising as (Fan et al., 2001) also reported that *A. thaliana* pollen tubes exhibit clear cytoplasmic streaming after 24 h incubation and even 36 h, although not all pollen tubes were healthy. In conclusion, our data presented indicate that S-specific interaction between PrpS and PrsS results in cell death of transgenic *A. thaliana* pollen expressing PrpS1-GFP and PrpS3-GFP fusion proteins, respectively.

Thomas and Franklin-Tong, (2004) reported for the first time that SI interaction in *Papaver* triggers PCD. They pretreated incompatible pollen with DEVDase inhibitor and measured pollen tube length. They demonstrated that Ac-DEVD-CHO alleviates pollen tube inhibition that is consequence of SI response and that pollen tube growth was recovered with the use of the caspase-3 inhibitor. Our results also indicate, that the SI induced decrease in viability could provide a link with DEVDase activity in *A. thaliana* pollen. PrpS1-GFP or PrpS3-GFP expressing *A. thaliana* pollen was challenged with PrsS1 or PrsS3, respectively, and stimulated SI response, which resulted in significantly decreased pollen viability. When pollen was pretreated with the inhibitor of the caspase-3 activity, Ac-DEVD-CHO, the viability was significantly higher in BG16 and HZ3 *A. thaliana* pollen. The results confirmed that poppy PrpS can be functional in *A. thaliana* pollen at least “*in vitro*”.
5.3.3. PCD in *A. thaliana* transgenic pollen

The data presented in the Section 5.4. indicates, that *Arabidopsis* transgenic pollen expressing PrpS₁-GFP or PrpS₃-GFP fusion proteins triggered PCD upon SI interaction. We analysed SI-mediated PCD in transgenic *A. thaliana* pollen further by investigating the role of the caspase-3-like activity. Jordan et al., (2000) demonstrated DNA degradation in incompatible *Papaver* pollen and the complete evidence for the PCD was provided by (Thomas and Franklin-Tong, 2004) with the study utilising caspase-3 inhibitor Ac-DEVD-CHO. In their study they also demonstrated SI-specific release of cytochrome-c into cytosol from mitochondria and PARP cleavage activity. Bosch and Franklin-Tong, (2007) took investigations of PCD in *Papaver* further. They widened their investigation by examining the involvement of other caspase-like activities in *Papaver* SI. They also provided a temporal profile on the caspase-3-like activity by imaging live cell activity with the use of specific probe CR-(DEVD)₂. The caspase-like activities in their study were measured by utilising a specific fluorescent probes, attached on the tetrapeptide, specific for caspase cleavage. In addition to DEVDase activity, VEIDase, LEVDase and YVADase activities were also identified (Bosch and Franklin-Tong, 2007, Bosch et al., 2010).

We investigated PCD in *A. thaliana* pollen by measuring DEVDase activity upon cleavage of the AMC tagged tetrapeptide DEVD (Ac-DEVD-AMC) and identified caspase-3-like activity that was highly increased when the transgenic pollen expressing PrpS₁-GFP or PrpS₃-GFP was incubated with PrsS₁ or PrsS₃ recombinant proteins. This activity was alleviated when pollen was pretreated with caspase-3 inhibitor Ac-DEVD-CHO prior to the SI induction. However, the basal DEVDase activity was observed in the control pollen and
for firmer statistical data the experiment should be repeated with higher pollen protein concentration.

In order to confirm that the transgenic *A. thaliana* system, expressing *Papaver* PrpS proteins, has the same downstream elements as poppy pollen, analysis of other caspase-like activities that were identified in *Papaver* would be beneficial (Bosch and Franklin-Tong, 2007, Bosch et al., 2010). For the investigation of the detailed temporal profile of caspase-3-like activity in incompatible transgenic *A. thaliana* pollen, live cell imaging could be applied using probe CR(DEVD)$_2$.

Characterization and functional analysis of transgenic *A. thaliana*, presented in this chapter, has confirmed that *Papaver* S-locus determinants can be transformed into *A. thaliana* and that hallmark features of *Papaver* SI are exhibited in *A. thaliana* pollen expressing fusion proteins PrpS-GFP. The demonstrated hallmark features, that were observed upon *in vitro* SI challenge of *A. thaliana* pollen expressing PrpS-GFP and cognate stigmatic PrsS recombinant protein, were pollen tube growth inhibition, formation of punctate actin foci, decrease of viability and increase in DEVDase activity.

Until recently, SI induced alterations to actin cytoskeleton, production of ROS and PCD were characteristic for SI in *Papaver*, but recently they were reported from another SI system – S-RNase induced SI in *Pyrus pyrifolia* pollen (Liu et al., 2007c, Wang et al., 2010). The SI induced alterations to actin cytoskeleton were not reported so far in the *Brassica* pollen upon SI interaction, however the study conducted by (Iwano et al., 2007) reported the SI induced actin alterations in the stigmatic papillae cells in *Brassica*. These data indicates that downstream events of SI, such as the alterations to the actin cytoskeleton, ROS production
and PCD, might be more conserved in different species than known so far, despite the evolutionary difference between the species. A possible hypothesis is that downstream events observed in *Papaver* upon the incompatible interaction between PrpS and PrsS (e.g. Ca$^{2+}$ influx and signaling, alterations to actin cytoskeleton and PCD) are the conserved and “primitive” signalling mechanisms that were used in response to convey a message and form a quick inhibitory response in plant cells. Plant-pathogen interaction and SI are examples of such stressful situations where a response to prevent the further growth of the cell is required very quickly. So it is possible that these mechanisms were used initially in many Angiosperms and *Papaver* kept it in this “basic” form to prevent self-fertilization, while other species evolved and refined ways by adding other signalling components to achieve the same result. However, that basic mechanism might be still conserved in plants and might be utilized again upon the interaction between cognate PrpS-PrsS pair. It would be interesting to know more specifically if this receptor-ligand interaction could work in plant somatic cells as well or maybe even in mammalian cells and it would be also interesting to see with other components of receptor-ligand signalling (e.g. SCR-SRK in *Brassica*) if they also trigger this basic response leading to inhibition of cell growth and cell death in other species. If the poppy SI-like mechanism of self-pollen inhibition in *A. thaliana* could be demonstrated as well *in vivo*, it would be very useful tool for further studies related to SI. The *Papaver* SI-like system would be useful to initiate studies in crop species as it would represent a novel way to prevent self-fertilization, which is an obstacle for successful plant breeding.
CHAPTER 6

Functional analysis \textit{in vivo} of transgenic \textit{A. thaliana} expressing \textit{Papaver} \textit{S}-determinants, PrsS and PrpS
6.1 INTRODUCTION

Papaver female S-determinant PrsS gene and whole $S_I$-locus were introduced into A. thaliana using Agrobacterium transformation. The transformation of the $S_I$-locus was carried out by Barend de Graaf (line BG6) and transformation of PrsS$_1$ and PrsS$_3$ by Huawen Zou (lines HZ1 and HZ2). This chapter describes the characterization of transgenic A. thaliana lines expressing PrsS$_1$ (HZ1), PrsS$_3$ (HZ2) and $S_I$-locus as well as in vivo functional analysis using PrsS$_1$ and PrsS$_3$ expressing plants and pollen from transgenic A. thaliana expressing PrpS$_1$-GFP and PrpS$_3$-GFP, described in previous chapter. The constructs are presented in Table 6.1.

Table 6.1: Cartoon illustrates the constructs of the poppy SI determinants that were used for Agrobacterium mediated transformation into A. thaliana.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Resistance Marker</th>
<th>Name</th>
<th>Construct</th>
</tr>
</thead>
<tbody>
<tr>
<td>PrsS$_1$</td>
<td>Basta</td>
<td>HZ1</td>
<td><img src="#" alt="Diagram" /></td>
</tr>
<tr>
<td>PrsS$_3$</td>
<td>Basta</td>
<td>HZ2</td>
<td><img src="#" alt="Diagram" /></td>
</tr>
<tr>
<td>$S_I$-locus</td>
<td>Kan</td>
<td>BG6</td>
<td><img src="#" alt="Diagram" /></td>
</tr>
</tbody>
</table>

To achieve tissue specific expression of the female S-determinants PrsS$_1$ and PrsS$_3$ Stig1 promoter from N.tabacum was used and for expression of $S_I$-locus native poppy promoter was used. Kanamycin (Kan) resistance cassette was used as a selection marker with $S_I$-locus expressing transformants BG6 and Basta resistance was used for selection of the PrsS expressing A. thaliana.
In order to analyse PrsS expressing transgenic *A. thaliana* lines *in vivo* and semi *in vitro* approaches were used. *Papaver*-like SI was analysed *in vivo* in *A. thaliana* in order to determine whether *Papaver* S-genes can be fully functional in *A. thaliana*. The functionality of the inserted PrsS and PrpS in *A. thaliana* were tested *in vivo* by pollinations assays and crosses for seed set. For this, T3 generation of transgenic *A. thaliana* lines HZ1.1-3.1 and HZ1.8-2.1 flowers expressing PrsS1, and HZ2.6-1.1 and HZ2.13-5.1 expressing PrsS3 were used as females, and lines BG16B#25.1-1 and HZ3.1-15.1, expressing PrpS1 and PrpS3, respectively, were used as pollen donors. Scheme of crosses can be seen on Figure 6.1. Incompatible response was expected with PrsS1 expressing stigma that was pollinated with PrpS1-GFP expressing pollen, and PrsS3 expressing stigma, pollinated with PrpS3-GFP expressing pollen. Controls were comprised of compatible crosses, e.g. PrsS1 stigma pollinated with PrpS3 pollen or PrsS3 stigma pollinated with PrpS1 pollen. Pollen functionality was tested by using Col-0 stigma as female, and PrpS1 or PrpS3 expressing pollen as male, while the receptiveness of the PrsS1 or PrsS3 expressing stigmas was tested by pollinations with Col-0 pollen. Following the emasculation and pollination, stigmas were either collected in aniline blue for the analysis of pollen tube growth or left to develop on the plant for the analysis of the seed set.
When conducting emasculations, the stage of stigma was crucial. The stigma had to be mature with the anthers just below the upper part of gynoecium so that it was not in contact with the self-pollen. This was normally achieved in stage 12, which was described by Smyth et al., (1990), as the stage where petals level with stamen. (Nasrallah et al., 2002) also reported that transgenic A. thaliana, ecotype Colombia, exhibiting A.lyrata SRKb/SCRb, was exhibiting transient SI that varied with the age of flowers. Young flowers in stage 12 and older flowers in late stage 14 were self-compatible, whereas stigmas in stage 13 and early 14 exhibited SI.

For our assays, pollen free stigmas were collected at the stage 12, when stigmas were receptive but before the pollen grains were mature. For pollination assays, following emasculation, pistils were excised and placed on a layer of 0.5 % agar to mature over night. The following day, stigmas reached the optimal biological stage 13 (or early stage 14) and they were manually pollinated with hundreds of pollen grains obtained from the mature post-anthesis flowers from PrpS-GFP expressing plants (or controls) and collected in aniline blue
16 h after pollination. For the seed set analysis, pistils were left on the plant to develop the siliques, which were collected before shedding seeds.

6.2. RESULTS

6.2.1. Characterization of A. thaliana expressing Papaver PrsS determinants

The primary goal was to analyse the PrsS expression of transgenic A. thaliana lines expressing PrsS₁, PrsS₃ or PrsS₈ under the control of Stig1 promoter. It was expected that the promoter would direct specific expression of PrsS at high levels in stigmatic tissue. No reporter gene was tagged with the PrsS, so the analysis relied upon the segregation of the resistance marker Basta and on the PCR analysis.

Seedlings of Arabidopsis::PrsS were initially screened by spraying Basta and only the plants containing the PrsS transgene were expected to survive. For the segregation analysis, seedlings were grown in MS medium containing Basta (see Materials & Methods) and resistant and non-resistant seedlings were counted. Heterozygous plants were expected to segregate in Mendelian fashion, while homozygous plants were expected to be 100 % resistant on the Basta containing media. The aim of the analysis was to identify the homozygous PrsS-expressing A. thaliana lines with the highest expression of the transgene.

PCR screening for the presence of inserts was performed initially on the leaves of young plants, using gene specific primers, as described in the Materials and Methods. The plants that were positive for the insert were selected for the analysis of the transcript by RT-PCR.
Figure 6.2. presents the RT-PCR results on HZ1 line expressing PrsS1 and HZ2 line expressing PrsS3.

PrsS1 insert was detected in 55 progenies out of 65, which showed a ~6:1 segregation ratio. The PCR product size with designed primers was 225 bp. The RT-PCR analysis was performed on 33 progenies with the insert confirmed by PCR on genomic DNA (Figure 6.2.a). Transcription of the PrsS genes were lower compared to the GAPC used as standard, and in the line HZ1.2.3, no band was detected, which could mean that the plant was not expressing PrsS1 or that the expression was below the level of detection.

With the PCR genotyping of 38 plants for PrsS3 insert in transgenic A. thaliana, 33 were found to be expressing PrsS3 transgene which resulted in a ~ 6:1 segregation ratio. The
product size using primers was 196 bp, which is as expected. Flowers from all the plants were analysed by RT-PCR for the relative expression of PrsS3 (Figure 6.2.b).

The functional analysis was only performed on the highest expressing PrsS1 and PrsS3 expressing lines HZ1 and HZ2, so the further selection of 5 samples was made initially, by quantifying the intensity of the bands using ImageJ software (see Figure 6.3). cDNA samples of plants HZ1.1-1, HZ1.1-2, HZ1.1-3, HZ1.3-5 and HZ1.8-2 from PrsS1 expressing flowers, and HZ2.6-1, HZ2.6-2, HZ2.6-5, HZ2.13-5 and HZ2.13-8 from PrsS3 expressing flowers were further re-analysed on the DNA gel and compared to the GAPC control (Figure 6.3.a).

![Figure 6.3: (a) Top 5 highest expressing PrsS1 and PrsS3 plants, and the corresponding GAPC control (bottom). Legend: 11-HZ1.1-1, 12-HZ1.1-2, 13-HZ1.1-3, 14-HZ1.3-5, 15-HZ1.8-2, 16-HZ2.6-1, 17-HZ2.6-2, 18-HZ2.6-5, 19-HZ2.13-5, 20-HZ2.13-8. (b) Column chart presenting ratio between transgene signal of PrsS and GAPC signal. Quantification was performed using ImageJ software. The highest expressors outlined in red.](image)

In order to quantify the expression of the transgene, the peak band intensity was measured using ImageJ software for each sample and its GAPC control (Figure 6.3.b). From each transgenic line, two plants with the highest expression identified by the intensity ratio were used in subsequent functional experiments. The apparent highest expressing lines were:
HZ1.1.-3 and HZ1.8-2 expressing PrsS₁, and HZ2.6-1 and HZ2.13-5 expressing PrsS₃. The progeny of these plants were used in further experiments to determine whether the PrsS was functional when expressed in A. thaliana stigma and the family trees are presented in the Appendix II.

Basta selection by spraying was used initially and later the Basta herbicide was used as a selection with the MS medium. To confirm the homozygous lines by Basta resistance in the medium, only lines that were chosen as the highest expressors were selected (Figure 6.3.). Those were lines HZ1.1.-3 and HZ1.8-2 expressing PrsS₁, and HZ2.6-1 and HZ2.13-5 expressing PrsS₃. The results of the segregation analysis are presented in Tables 6.2 and 6.3.

**Table 6.2:** Segregation of Basta resistance in progenies of transgenic A. thaliana lines expressing *Papaver* stigma determinant PrsS₁. Statistical analysis was performed using Chi square analysis. HO – homozygous

<table>
<thead>
<tr>
<th>Seeds (PrsS₁)</th>
<th>Generation</th>
<th>Bastaᴿ</th>
<th>Bastaᴮ</th>
<th>N</th>
<th>Ratio</th>
<th>χ² (3:1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HZ1.1</td>
<td>T₁</td>
<td>16</td>
<td>2</td>
<td>1</td>
<td>8:1</td>
<td>1.85</td>
</tr>
<tr>
<td>HZ1.1-3</td>
<td>T₂</td>
<td>19</td>
<td>1</td>
<td>1</td>
<td>19:1</td>
<td>4.27</td>
</tr>
<tr>
<td>HZ1.1-3.1</td>
<td>T₃</td>
<td>63</td>
<td>0</td>
<td>2</td>
<td>HO</td>
<td>21</td>
</tr>
<tr>
<td>HZ1.1-3.1-x</td>
<td>T₄</td>
<td>76</td>
<td>0</td>
<td>2</td>
<td>HO</td>
<td>25</td>
</tr>
<tr>
<td>HZ1.8</td>
<td>T₁</td>
<td>6</td>
<td>5</td>
<td>1</td>
<td>1.2:1</td>
<td>2.45</td>
</tr>
<tr>
<td>HZ1.8-2</td>
<td>T₂</td>
<td>31</td>
<td>0</td>
<td>1</td>
<td>HO</td>
<td>10.3</td>
</tr>
<tr>
<td>HZ1.8-2.1</td>
<td>T₃</td>
<td>61</td>
<td>0</td>
<td>2</td>
<td>HO</td>
<td>20.3</td>
</tr>
<tr>
<td>HZ1.8-2.1-x</td>
<td>T₄</td>
<td>75</td>
<td>0</td>
<td>2</td>
<td>HO</td>
<td>25</td>
</tr>
</tbody>
</table>

**Table 6.3:** Segregation of Basta resistance in progenies of transgenic A. thaliana lines expressing *Papaver* stigma determinant PrsS₃. Statistical analysis was performed using Chi square analysis. HO - homozygous

<table>
<thead>
<tr>
<th>Line (PrsS₃)</th>
<th>Generation</th>
<th>Bastaᴿ</th>
<th>Bastaᴮ</th>
<th>N</th>
<th>Ratio</th>
<th>χ² (3:1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HZ2.6</td>
<td>T₁</td>
<td>15</td>
<td>6</td>
<td>1</td>
<td>2.5:1</td>
<td>0.14</td>
</tr>
<tr>
<td>HZ2.6-1</td>
<td>T₂</td>
<td>11</td>
<td>6</td>
<td>1</td>
<td>1.8:1</td>
<td>0.96</td>
</tr>
<tr>
<td>HZ2.6-1.1</td>
<td>T₃</td>
<td>47</td>
<td>12</td>
<td>2</td>
<td>3.9:1</td>
<td>0.68</td>
</tr>
<tr>
<td>HZ2.6-1.1-x</td>
<td>T₄</td>
<td>81</td>
<td>12</td>
<td>2</td>
<td>7:1</td>
<td>7.3</td>
</tr>
<tr>
<td>HZ2.13</td>
<td>T₁</td>
<td>15</td>
<td>4</td>
<td>1</td>
<td>3.8:1</td>
<td>0.16</td>
</tr>
<tr>
<td>HZ2.13-5</td>
<td>T₂</td>
<td>11</td>
<td>0</td>
<td>1</td>
<td>HO</td>
<td>3.67</td>
</tr>
<tr>
<td>HZ2.13-5.1</td>
<td>T₃</td>
<td>46</td>
<td>2</td>
<td>2</td>
<td>23:1</td>
<td>11.1</td>
</tr>
<tr>
<td>HZ2.13-5.1-x</td>
<td>T₄</td>
<td>59</td>
<td>0</td>
<td>2</td>
<td>HO</td>
<td>19.7</td>
</tr>
</tbody>
</table>
The expected 3:1 segregation ratio (Basta\textsuperscript{R}:Basta\textsuperscript{s}) based on the inheritance of Basta resistance was observed only in T\textsubscript{1} (HZ2.6) and T\textsubscript{3} (HZ2.6-1.1) generation of PrsS\textsubscript{3} expressing \textit{A. thaliana} (Table 6.3). In PrsS\textsubscript{1} expressing \textit{A. thaliana} (Table 6.2) it was not observed. Instead, the analysed HZ1.1 line exhibited distorted Mendelian 8:1 segregation ratio in T\textsubscript{1} generation (Table 6.2) and its progeny showed 19:1 ratio in T\textsubscript{2} generation. The line became homozygous for Basta in generation T\textsubscript{3}. Segregation ratio of 7:1 was also observed in T\textsubscript{4} generation of HZ2.6-1.1.

(Page and Grossniklaus, 2002) reported that 7:1 segregation ratio is expected for the recessive mutants obtained by seed mutagenesis. However, we did not mutagenise the seeds. Another possibility (Rudolph, 1966) could be that 7:1 segregation might result from the 15:1 ratio distorted due to the lethality dependent on PrsS expression, and in next generation of HZ1.1-3 we observed increased segregation ratio, meaning we could have the insertion at two loci.

The other line selected for PrsS\textsubscript{1} expression showed 1:1 segregation ratio in T\textsubscript{1}, but interestingly exhibited homozygous state already from generation T\textsubscript{2} onwards. HZ2.6 and HZ2.13 lines analysed showed the expected Mendelian segregation ratio 3:1 in T\textsubscript{1} generation. However, line HZ2.6 exhibits non-Mendelian ratios in subsequent generations: 2:1 ratio in generation T\textsubscript{2}, 4:1 ratio in T\textsubscript{3} and very distorted ratio 7:1 in generation T\textsubscript{4}. On the other hand, in the line HZ2.13 the segregation of resistance seems equally unstable, changing from generation to generation. It shows homozygous expression in T\textsubscript{2} and T\textsubscript{4} but ratio 23:1 in generation T\textsubscript{3}. \textit{A. thaliana} transgenic lines that were selected for further analysis based on the expression of the transcript analysed by RT-PCR were HZ1.1-3 and HZ1.8-2 expressing PrsS\textsubscript{1}, and lines expressing PrsS\textsubscript{3} were HZ2.6-1 and HZ2.13-5.
Since the RT-PCR showed that PrsS1 mRNA was specifically present in HZ1 flowers, we wished to confirm the PrsS1 protein presence using western blot, however, it was impossible to detect any PrsS1 signal using the 20-50 μg total protein extract and only PrsS1 recombinant protein, expressed in *E. coli*, resulted in a very strong band (data not shown). Considering the RT-PCR detection of the PrsS1 mRNA transcripts in transgenic *A. thaliana* flowers, the PrsS1 protein are expressed at extremely low levels that are below detection using western blot, or the protein is not properly expressed or folded. In any case, this could explain some of the results described later in section 6.2.4.

**6.2.2. *S1*-locus expressing *A. thaliana* line**

Another important construct that was generated by Barend de Graaf included the whole *S1*-locus from *Papaver* (both, PrpS1 and PrsS1 determinants at the same time) and was coupled with endogenous *Papaver* promotors for PrpS1 and PrsS1 (see Table 6.1). Segregation of the Kan resistance was investigated in the progeny of T1 generation of *A. thaliana* line BG6 expressing the whole *S1*-locus (Table 6.4).
Table 6.4.: Segregation of the kanamycin resistance in progenies of transgenic *A. thaliana* lines expressing *Papaver S1*-locus. Statistical analysis was performed using Chi square analysis.

<table>
<thead>
<tr>
<th>Line (S1-locus)</th>
<th>Generation</th>
<th>KanR</th>
<th>KanS</th>
<th>N</th>
<th>Ratio</th>
<th>$\chi^2$ (3:1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BG6.1</td>
<td>T1</td>
<td>47</td>
<td>20</td>
<td>1</td>
<td>2.35:1</td>
<td>0.84</td>
</tr>
<tr>
<td>BG6.2</td>
<td>T1</td>
<td>24</td>
<td>22</td>
<td>1</td>
<td>1.1:1</td>
<td>12.8</td>
</tr>
<tr>
<td>BG6.3</td>
<td>T1</td>
<td>0</td>
<td>50</td>
<td>1</td>
<td>0:1</td>
<td>150</td>
</tr>
<tr>
<td>BG6.4</td>
<td>T1</td>
<td>0</td>
<td>50</td>
<td>1</td>
<td>0:1</td>
<td>150</td>
</tr>
<tr>
<td>BG6.5</td>
<td>T1</td>
<td>71</td>
<td>25</td>
<td>1</td>
<td>3:1</td>
<td>0.056</td>
</tr>
<tr>
<td>BG6.6</td>
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<td>0</td>
<td>52</td>
<td>1</td>
<td>0:1</td>
<td>150</td>
</tr>
<tr>
<td>BG6.7</td>
<td>T1</td>
<td>27</td>
<td>11</td>
<td>1</td>
<td>2.5:1</td>
<td>0.32</td>
</tr>
<tr>
<td>BG6.8</td>
<td>T1</td>
<td>18</td>
<td>32</td>
<td>1</td>
<td>0.5:1</td>
<td>40.6</td>
</tr>
<tr>
<td>BG6.9</td>
<td>T1</td>
<td>19</td>
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<td>1</td>
<td>0.2:1</td>
<td>103.2</td>
</tr>
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<td>BG6.10</td>
<td>T1</td>
<td>4</td>
<td>21</td>
<td>1</td>
<td>0.2:1</td>
<td>46.4</td>
</tr>
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<td>34</td>
<td>1</td>
<td>0.3:1</td>
<td>61.3</td>
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<td>T1</td>
<td>2</td>
<td>48</td>
<td>1</td>
<td>0.04:1</td>
<td>134.4</td>
</tr>
<tr>
<td>BG6.13</td>
<td>T1</td>
<td>8</td>
<td>28</td>
<td>1</td>
<td>0.3:1</td>
<td>53.5</td>
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In total, 28 original primary transformants were analysed. Seven were Kan sensitive and did not survive the selection process. The other lines exhibited various segregation ratios: four exhibited the 3:1 segregation ratio (BG6.5, BG6.7, BG6.17 and BG6.19), five lines exhibited a ratio of 2:1 (BG6.1, BG6.7, BG6.16, BG6.22 and BG6.28), only five lines exhibited the expected segregation ratio of 1:1 (BG6.2, BG6.14, BG6.15, BG6.21 and BG6.26) and seven lines exhibited a ratio of less than 0.5:1 (BG6.9, BG6.10, BG6.11, BG6.12, BG6.13, BG6.20, BG6.28).
and BG6.27). As previously mentioned, the expected segregation ratio in fully functional BG6 line would be 1:1. If the plants were heterozygous then 50 % of pollen would be expected to express PrpS1 and 50 % of pollen would be wt and only the wt pollen would be allowed to grow through the stigma and successfully fertilize the ovules. The pollen expressing PrpS1 would be expected to be inhibited upon landing on the stigma expressing PrsS1. Therefore, the expectancy of a 1:1 segregation ratio, but it was found that only ~20 % of all BG6 plants exhibited this ratio.

Some plants expressing S1-locus were analysed by RT-PCR using PrpS1 and PrsS1 primers. Initially, only two samples (randomly chosen showing 3:1 and 0.2:1 ratio) were analysed using PrpS1 and PrsS1 primers in the reaction simultaneously (Figure 6.4.a). The PrsS1 band was detected upon analysis on the DNA gel but PrpS1 product was not visualised. The RT-PCR was repeated using more samples with different segregation ratios, including the expected 1:1, and the primers were used in separate reactions (e.g PrsS1 primers were used with an aliquot of RNA in one reaction, and another aliquot of RNA was used in the RT-PCR reaction using PrpS1 primers, see Figure 6.4.b) but no PrpS1 band was visualised and identified when analysing the PCR product by electrophoresis.
The RNA extraction and RT-PCR test was repeated again by colleague Javier-Andres Juarez-Diaz, using primers designed over different area on PrpS1 (PrpS1 a.n. primers), different set of pipettes and different work space but with the same result: PrpS1 mRNA could not be amplified from BG6 samples.

Since the RT-PCR analysis could not be used in order to identify lines with the highest transgene expression, seed set analysis was performed on all the lines from the BG6 family expressing S1-locus

### 6.2.3. *Papaver* pollen growth in the presence of stigmatic extract of *A. thaliana* expressing PrsS1 and PrsS3

Previous chapter presenting “in vitro” results demonstrated that *Papaver* PrpS proteins were functional when expressed in *A. thaliana*. The functional SI response was achieved through the interaction between recombinant stigmatic PrsS proteins and PrpS-GFP receptor expressed in *A. thaliana*. As the stigmatic PrsS proteins were also transformed to *A. thaliana*, with targeted expression in the stigma, we wished to investigate whether stigmatic PrsS proteins were functional when expressed in *A. thaliana*. To *Papaver* S1S3 pollen
growing in petridishes we added the stigmatic protein extract derived from transgenic *A. thaliana* flowers expressing PrsS₁ and PrsS₃ proteins, respectively. Transgenic *A. thaliana* T₄ generation plant lines were characterized homozygous in previous section, and were used for the experiment. *A. thaliana* plants from lines HZ1.1-3.1-x and HZ1.8-2.1-x were used as a source for PrsS₁ expressing stigmas, and lines HZ2.6-1.1-x and HZ2.13-5.1-x for PrpS₃ expressing stigma.

Employing this semi-*in vitro* system, “incompatible” pollen (*Papaver* S₁ and S₃ pollen incubated with stigmatic HZ1 and HZ2 extracts) should be inhibited, while pollen tube length of the control pollen (*Papaver* S₂ and S₄ pollen incubated with HZ1 and HZ2 extracts) should not be affected by the PrsS₁ and PrsS₃ expressing flower extracts. HZ1 and HZ2 stigmatic extracts were added to *Papaver* PrpS₁/PrpS₃ and PrpS₂/PrpS₄ pollen. As a positive control we induced SI response in *Papaver* pollen with recombinant stigmatic PrsS₁ and PrsS₃. In order to affirm that the effect of the PrsS expressing *A. thaliana* stigmas was due to PrsS proteins and not to other soluble proteins present in *A. thaliana* stigmatic tissue, we also challenged *Papaver* PrpS₁/PrpS₃ pollen with *A. thaliana* untransformed Col-0 stigmatic extract and left them to grow in GM only (untreated, UT). Additional controls were the heat denatured (hd) stigmatic extracts from HZ1 and HZ2 flowers and from Col-0 flower, in order to ensure that the response was due to the biologically active PrsS proteins. The S-specificity control was comprised of *Papaver* PrpS₂/PrpS₄ pollen that was incubated with stigmatic extracts from HZ1 expressing PrsS₁ and HZ2 flowers expressing PrsS₃ (i.e. a compatible allelic combinations).

Pollen was left to incubate with proteins overnight and pollen tubes were imaged and measured after 20 h incubation. *Papaver* PrsS₁/PrsS₃ pollen was inhibited by the addition of *A. thaliana* HZ1 and HZ2 stigmatic extract (Figure 6.5.e), although not as strongly as pollen
challenged with incompatible recombinant PrsS₁ and PrsS₃ proteins (Figure 6.5.b). Moreover, some pollen tube tips were affected by the HZ1 and HZ2 stigmatic extracts and some developed extremely swollen tubes and bulbous tips (Figure 6.5.e). When pollen was incubated with wild type stigmatic extracts (Figure 6.5.c) or with heat denatured HZ1&HZ2 extract (Figure 6.5.f), pollen tubes were not inhibited. There were some pollen tube tips slightly swollen, however, to lesser extent than pollen incubated with the HZ1 and HZ2 stigmatic extracts. Long pollen tubes without swollen tips were observed when HZ1&HZ2 stigmatic extracts were incubated with *Papaver S₂S₄* pollen (Figure 6.5.i).

**Figure 6.5:** Images of *Papaver* pollen tubes incubated with stigmatic protein extracts from transgenic *A. thaliana* flowers: HZ1.1-3.1-x and HZ1.8-2.1-x flowers expressing PrsS₁, and HZ2.6-1.1-x and HZ2.13-5.1-x (a) UT *Papaver S₁S₁* pollen; (b) *Papaver S₁S₁* pollen incubated with recombinant PrsS₁ and PrsS₃; (c) *Papaver S₁S₁* pollen incubated with Col-0 stigmatic extract; (d) *Papaver S₁S₁* pollen incubated with hd Col-0 stigmatic extract; (e) *Papaver S₁S₁* pollen incubated with HZ1&HZ2 stigmatic extract; (f) *Papaver S₁S₁* pollen incubated with hd HZ1&HZ2 stigmatic extract; (g) *Papaver S₂S₄* pollen UT; (h) *Papaver S₂S₄* pollen with Col-0 stigmatic extract; (i) *Papaver S₂S₄* with HZ1&HZ2 stigmatic extract. Scale bar represents 100 μm.
In order to quantify the observed effects of *A. thaliana* stigmatic extracts on *Papaver* pollen we measured pollen tube length and tip diameter (see Figure 6.6). Figure 6.6.a presents the mean length of *Papaver* pollen tubes, incubated with *A. thaliana* stigmatic proteins.

![Figure 6.6: Quantification of *Papaver* pollen tube length and tip diameter upon incubation with *A. thaliana* stigmatic protein extracts (Col-0 – untransformed *A. thaliana* stigmatic protein extract, HZ1HZ2 – PrsS1 and PrsS3 expressing *A. thaliana* stigmatic protein extracts) and controls (a) average pollen tube length and (b) average tube tip diameter. S1S3 *Papaver* pollen untreated and recombinant stigmatic controls are presented with light grey bars, pollen incubated with *A. thaliana* stigmatic proteins and controls are presented with dark grey bars, while control S2S4 pollen is presented in teal bars. Red outline indicates incompatible interaction between pollen and pistil determinant. Error bars represent ± St.Deviation. UT – untreated, rS8 – recombinant PrsS8; hd – heat denatured. Some of these experiments were conducted by Javier-Andres Juarez-Diaz.](image)

Unchallenged PrsS1/PrsS3 *Papaver* pollen tube length was 958±60 μm, while length of pollen with recombinant PrsS1 and PrsS3 proteins added was 38±6 μm (P= 2.7x10^{-108},
N=80), which is only 4% of untreated control. The pollen tube length of *Papaver* incubated with stigmatic protein extract from PrsS₁ and PrsS₃ expressing *A. thaliana* flowers, was reduced significantly by 60% compared to the UT control (P=1.1x10⁻⁶¹, N=80) or to pollen incubated with non-transgenic stigmatic extract (945±69 μm, P=1.3x10⁻⁵⁶, N=80), reaching average length of 380±51 μm. The tip diameter of pollen incubated with stigmatic protein extract from HZ1 and HZ2 *A. thaliana* flowers (Figure 6.6.b) was significantly increased with 30±2 μm compared to 11±1 μm for untreated pollen (P=1.9x10⁻⁷⁴, N=80) or 15±1 μm for pollen incubated with Col-0 protein extract (P=7.0x10⁻⁵⁷, N=80). Untreated compatible poppy PrsS₂/PrsS₄ pollen reached the length of 925±56 μm, which was comparable to untreated PrsS₁/PrsS₃ pollen (P=0.25; N=60). The length of PrsS₂/PrsS₄ pollen treated with Col-0 stigmatic extract was 911±68 μm, which was not different to Col-0 treated PrsS₁/PrsS₃ pollen (P=0.35, N=60). However, the length of “compatible” poppy pollen incubated with HZ1 and HZ2 stigmatic extracts was 831±54 μm, which was significantly different compared to HZ1 and HZ2 incubation of PrsS₁/PrsS₃ pollen (P=1.5x10⁻³⁵, N=60). Quantification also revealed the significant difference between tips of compatible PrsS₂/PrsS₄ pollen incubated with HZ1 and HZ2 stigmatic extracts, with diameter of 16±1 μm and HZ1&HZ2 incubated S₁S₃ pollen (P=3.7x10⁻⁴²).

These data demonstrate PrsS₁ and PrsS₃ stigmatic proteins that were expressed in *A. thaliana* stigmas under stigma specific Stig1 promoter, had an inhibitory effect on PrsS₁/PrsS₃ *Papaver* pollen in an S-specific manner. However, the interaction causes not only *Papaver* pollen tube inhibition but also some other morphological changes, like tube swelling.
6.2.4 *Papaver SI in vivo in transgenic Arabidopsis thaliana*

6.2.4.1 Pollination assays in transgenic *A. thaliana expressing Papaver PrsS and PrpS*

Incompatible crosses and controls were examined with a microscope. Figure 6.7 shows analysis of the highest PrsS$_1$ expressing *A. thaliana* lines HZ1.1-3.1 and HZ1.8-2.1. Crosses for the aniline blue staining of the pollen tube growth were conducted on *A. thaliana* plants fourteen times in total. Pollination tests were initially conducted on T$_2$ plants, and after the highest expressing lines were identified on T$_3$ plants (see Figure 6.7).
Figure 6.7: Aniline blue staining of crosses with PrsS₁ expressing *A. thaliana* stigmas. (a) HZ1.1-3.1 x BG16#25.1-1 (PrsS₁ x PrpS₁), (b) HZ1.1-3.1 x HZ3.1-15.1 (PrsS₁ x PrsS₁), (c) HZ1.1-3.1 x HZ1.1-3.1 (PrsS₁ x PrsS₁); (d) HZ1.1-3.1 x Col-0 (PrsS₁ x Col-0); (e) HZ1.8-2.1 x BG16#25.1-1 (PrsS₁ x PrpS₁); (f) HZ1.8-2.1 x Col-0 (PrsS₁ x Col-0). Scale represents 500 μm.
Few crosses resulted in reduced pollen tube growth but generally the incompatible crosses resulted in long pollen tube growth through the style, which could indicate that the plants were heterozygous. More likely, the pollen tube growth could be due to the weak or ‘leaky’ SI response. The incompatible crosses HZ1.1-3.1 x BG16#25.1-1 and HZ1.8-2.1 x BG16#25.1-1, presented in Figures 6.7.a and 6.7.e, respectively, show long pollen tube growth, which was comparable to the control and compatible pollinations. Control crosses confirmed that HZ1.1-3.1 and HZ1.8-2.1 flowers were receptive and functional (Figures 6.7.c, d & f) but did not exhibit full SI.

Preliminary investigation also examined the pollination of PrsS3 expressing HZ2.13-5.1 and HZ2.6-1.1 flowers (see Figure 6.8) by PrpS3 expressing HZ3.1-15.1 and non-transgenic or compatible control.
Figure 6.8: Aniline blue staining of crosses with PrsS$_3$ expressing A. thaliana stigmas. (a) HZ2.13-5.1 x HZ3.1-15.1 (PrsS$_3$ x PrpS$_3$) (b) HZ2.13-5.1 x Col-0 (PrsS$_3$ x Col-0); (c) HZ2.6-5.1 x HZ3.1-15.1 (PrsS$_3$ x PrpS$_3$); (d) HZ2.6-5.1 x BG16#25.1-1 (PrsS$_3$ x PrpS$_1$).
Incompatible crosses of HZ2.13-5.1 x HZ3.1-5.1 and HZ2.6-5.1 x HZ3.1-15.1, presented on figure 6.8.a and 6.8.c, respectively, exhibited compatible phenotype as there were numerous pollen tubes growing down the style.

In order to characterize the pollinations in greater detail, more systematic and quantitative analysis was proposed. However, the PrsS₁ protein signal was not detected using western blot analysis on the PrsS₁ expressing stigmatic extracts, despite the RT-PCR analysis confirmed the presence of PrsS₁ mRNA. There could be several reasons that there was no protein detected by western blot. It is possible that PrsS₁ protein was not correctly translated, not correctly targeted or it was expressed at very low concentration that was below the level of detection by western blot analysis. Therefore the inhibition of PrpS₁-GFP pollen in vivo did not occur. However, the results of incubation of Papaver S₁S₃ pollen with HZ1 and HZ2 stigmatic protein extracts indicated, that HZ1 and HZ2 expressing PrsS₁ and PrsS₃, respectively could be functional. The total protein concentration within that experiments was 100 μg mL⁻¹ but we could not quantify what proportion (if any) of that concentration was represented by properly folded, active PrsS proteins. The inhibition of Papaver pollen was observed, but at the same time also pollen tube swelling which could indicate that the PrsS proteins were possibly not properly folded or at very low concentration. There was also possibility that the insertion of the PrsS by transformation in A. thaliana caused some other transient stigmatic phenotype that could not be observed visually but affected the total stigmatic protein concentration. And finally the stochiometry of the SI response, i.e. number of molecules of receptor and ligand required for the SI response to be triggered, also remains unknown.
6.2.4.2 Seed set analysis in transgenic *A. thaliana* expressing *Papaver* PrsS and PrpS

Seed set analysis was conducted in parallel on transgenic *A. thaliana* lines to test *Papaver* SI *in vivo*. HZ1.1-3.1 and HZ1.8-2.1 plants expressing PrsS₁ and HZ2.6-1.1 and HZ2.13-5.1 plants, expressing PrsS₃ were emasculated and pollinated with selected pollen donor to obtain: incompatible cross, reciprocal cross or other allele specific controls (see Materials and Methods). Siliques were left on the plant to develop seeds before they were collected and measured with the seed set counted. However, not all the crosses were successful, hence the discrepancy in the number of fertilized siliques.

Figure 6.9.a presents the seed set from all the individual incompatible crosses (Figure 6.9.a, red outline, yellow bars for PrsS₁ x PrpS₁ and light green bars for PrsS₃ x PrpS₃) and controls. Certain incompatible crosses resulted in smaller silique size and lower seed set (HZ1.8-2.1 x BG16A#19.8-3, HZ2.6-1.1 x HZ3.1-15.1 and HZ2.6-1.1 x HZ3.1-15.1-1), however, the majority of incompatible crosses resulted in the normal seed set, compared to compatible reciprocal crosses. Crosses were performed by Steve Price.

In order to analyze the female expressors (PrsS₁ expressing lines HZ1.1-3.1 or HZ1.8-2.1 and PrsS₃ expressing lines HZ2.6-1.1 or HZ2.13-5.1) as pollen acceptors and pollen donors, the silique length and seed set data of incompatible pollinations with the same PrsS expressing line was joined (Figure 6.9.b, red outline) and its reciprocal control for individual PrsS expressing lines (see Figure 6.9.b).

The incompatible crosses with PrsS₁ expressing *A. thaliana* stigma HZ1.1-3.1 and PrpS₁-GFP expressing BG16 pollen resulted in 29.1±12.9 seeds and silique length 16.3±2.1 mm (n=14), and were 43 % compared to its reciprocal cross with 41.6±16.2 seeds and silique
length 17.0±2.4 mm (n=11) (see Figure 6.3.3.b, yellow bars). The seed set of this incompatible cross (PrsS₁ x PrpS₁) was also 17 % when compared to the cross with Col-0 (HZ1.1-3.1 x Col-0), which resulted in 34 seeds (n=1) but 18 % higher when compared to the compatible cross HZ1.1-3.1 x HZ3 with 23.9±8.6 seeds and silique length 14.1±1.9 mm (n=5). PrsS₁ expressing HZ1.8.2-1 plants crossed with incompatible PrpS₁-GFP expressing BG16 had on average 19 % higher seed set with 34.5±14.3 seeds and silique length 15.1±2.7 mm (n=13) when compared with HZ1.1-3.1, and as well 30 % higher seed set than its reciprocal control BG16 x HZ1.8.2-1 with 24.7±12.1 seeds and silique length 13.1±3.8 mm (n=5). When incompatible HZ1.8-2.1 cross (PrsS₁ x PrpS₁) was compared to the compatible control HZ1.8-2.1 x HZ3, the compatible seed set was 18 % lower with 28.3±11.4 seeds and silique length 12.8±2.8 mm (n=5). These data indicates that incompatible cross was not completely functional, indicating that transgenic A. thaliana might not be expressing PrsS at high enough levels.
Figure 6.9: (a) Seed set of all the incompatible crosses and controls; yellow bars, red outline: incompatible crosses PrsS₁ x PrpS₁, spring green bars, red outline: incompatible crosses PrsS₃ x PrpS₁, dark blue bars: reciprocal crosses: PrpS₁ x PrsS₁ and PrpS₃ x PrsS₃, light blue bars compatible reciprocal crosses: PrpS₁ x PrsS₃ and PrpS₃ x PrsS₁, dark green bars compatible controls: PrsS₁ x PrpS₃ and PrsS₃ x PrpS₁, grey bars: various control crosses including Col-0. Error bars ± St. Error. (b) Joined data of a same PrsS₁ expressing plant line with incompatible crosses (red outline on bars), its reciprocal crosses and compatible controls. Yellow bars: pooled HZ1.1-3.1 data, orange bars: pooled HZ1.8-2.1 data, spring green bars: joined HZ2.6-1.1 data and dark green bars: joined HZ2.13-5.1 data. Grey bars represent average siliquae length. Error bars ± Standard Deviation.

PrsS₃ expressing flowers HZ2.6-1.1 that were crossed with incompatible PrpS₃-GFP expressing HZ3 pollen had an average seed set of 14.0±9.0 seeds and siliquae length 13.2±2.6
mm (n=11), which is ~130 % lower than its reciprocal crosses HZ3 x HZ2.6-1.1 with 31.9±14.0 seeds and siliques length 18±3.3 mm (n=10). The seed set of compatible control HZ2.6-1.1 x BG16 was 36 % higher with 19.1±9.7 seeds and siliques length of 13.8±2.7 mm (n=6). Another set of incompatible crosses with PrsS3 expressing flower HZ2.13-5.1 x HZ3 was 6 % lower with 23.2±12.1 and siliques length 14.8±2.8 mm (n=21), compared to its reciprocal cross HZ3 x HZ2.13-5.1 with 24.6±11.4 seeds and siliques length 14.6±2.4 mm (n=8) and 25 % lower compared to the compatible cross HZ2.12-5.1 x BG16 with 29.1±16 seeds and siliques length of 12±4 mm (n=7).

Taken together, the analysed seed set data indicate that the incompatible PrpS combinations might have an effect on the PrsS1 expressing line HZ1.1-3.1 and PrsS3 expressing line HZ2.6-1.1. However, the incompatible pollinations should not result in any seed set, so the result observed could be due to leaky phenotype, very weak expression as discussed above, or does not work at all.

6.2.4.3 Seed set and siliques length of the F1 progeny from incompatible crosses

Seeds from incompatible crosses of *A. thaliana* expressing *Papaver PrsS* and *PrpS-GFP* genes were tested for germination and seed set after self-pollination in F1 generation. 30 seeds from each of the following crosses were selected: HZ1.1-3.1 x BG16#25.1-1 (PrsS1 x PrpS1), HZ1.8-2.1 x BG16#25.1-1 (PrsS1 x PrpS1), reciprocal cross BG16#25.1-1 x HZ1.8-2.1 (PrpS1 x PrsS1), HZ2.6-1.1 x HZ3.1-15.1-1 (PrsS3 x PrpS3) and HZ2.13-5.1 x HZ3.1-15.1 (PrsS3 x PrpS3), and germinated on MS media with double selection Kan and Basta. If the transfer of the genetic material occurred then the resulting seeds would be resistant to
Basta that was inherited through maternal side, and to Kan that was inherited through the paternal side by pollination. Because the parents used were homozygous, all the seeds were expected to survive the selection. If the incompatible cross would have been successful then no seeds would have been expected, however, seeds resulted from all the crosses so it would as well be possible that seeds would be sterile in the case of incompatible cross.

The Mendelian segregation (live to dead ratio) of F1 seedlings from cross HZ1.1-3.1 x BG16#25.1-1 (PrsS1 x PrpS1) was 23:7 (3:1). The F1 seedlings resulting from incompatible cross HZ1.8-2.1 x BG16#25.1-1 (PrsS1 x PrpS1) exhibited segregation 5:25 (0.2:1) with the first repeat and 7:23 (or 0.3:1) with the second repeat on double selection media. F1 from reciprocal cross BG16#25.1-1 x HZ1.8-2.1 (PrpS1 x PrsS1) resulted in 2:1 segregation ratio. Segregation ratios of F1 seedlings in case of incompatible crosses with PrsS3-expressing stigmas were as follows: HZ2.6-1.1 x HZ3.1-15.1 (PrsS3 x PrpS3) resulted in one surviving seedling and exhibited a segregation ratio of 1:30 (or 0.03:1), and HZ2.13-5.1 x HZ3.1-15.1 (PrsS3 x PrpS3) exhibited a 26:4 ratio (or 6.5:1) with first repeat, and a 14:16 ratio (or ~ 1:1) with the second repeat. A. thaliana non-transformed control germinated only on MS media without any selections and did not germinate on double selection media. These data indicates that both determinants (PrsS and PrpS) are present in the F1 seeds, therefore it would be expected that self-fertilization is prevented in those plants if both PrsS and PrpS are functional in A. thaliana.

Surviving F1 seedlings were planted in soil and left to self-fertilize following which the seed set was counted and silique length was measured. From each F1 line 5 plants were chosen and 10 siliques measured on each plant, except in the case of the first repeat of HZ1.8-2.1 x BG16#25.1-1 (2 plants) and HZ2.6-1.1 x HZ3.1-15.1
(1 plant only) as they were the only plants surviving after selection on plate and growth in pots. Seed set of F₁ plants derived from incompatible crosses is presented on Figure 6.10.

Self-fertilized A. thaliana Col-0 plants had 54±7.4 seeds (n=50 siliques measured in total). F₁ plants resulting from HZ1.1-3.1 x BG16#25.1-1 that exhibited 3:1 segregation ratio had the seed set of 37.8±11.9, which is significantly lower compared to Col-0 seed set (n=50; P=1.4*10⁻¹²). Seed set of F₁ plants resulting from cross HZ1.8-2.1 x BG16#25.1-1 (PrsS₁ x PrpS₁) with the segregation of 0.2:1 was 17.9±5.3 (n=20, P=7.3*10⁻³⁰ compared to Col-0) for the first repeat and 21.8±5.4 (n=50, P=4.9*10⁻⁴⁴ compared to Col-0) for the second repeat, both repeats were significantly lower compared to Col-0 self seed set. Reciprocal cross BG16#25.1-1 x HZ1.8-2.1 (PrsS₁ x PrpS₁) with 2:1 segregation ratio gave 47.8±9 seeds, which is significantly higher compared to seed set resulting from incompatible cross (n=50; P=3.1*10⁻⁴ compared to Col-0, P=3.8*10⁻²¹ compared to its incompatible cross, repeat 1 and P=1.5*10⁻³¹ compared to its incompatible cross, repeat 2).
Figure 6.10: Silique length (grey bars) and seed set of surviving F1 seedlings. Green bars PrsS3 expressing stigmas used as females, yellow bars PrSs1 expressing stigmas used as females in original cross, teal bar, reciprocal cross with BG16#25.1-1 used as a female and light blue bar Col-0 control. n=10 siliques; N=5 plants (except in case of the first repeat of HZ1.8-2.1 x BG16#25.1-1 (N=2 plants) and HZ2.6-1.1 x HZ3.1-15.1 (N=1 plant). Error bars represent ± St. error.

Seed set from F1 plants derived from HZ2.6-1.1 x HZ3.1-15.1 (PrsS3 x PrSs3) cross was 15.9±4.2 seeds and is significantly lower compared to Col-0 self-seed set, which was 54±7.4 (n=10, P=1.2*10^{-16}). On the other hand, seed set from F1 plants that were derived from HZ2.13-5.1 x HZ3.1-15.1 cross was 46.4±9.7 seeds for repeat 1, (n=50) and 48.3±8.4 seeds for the repeat 2 (n=50). Both these values were not significantly different from the Col-0. F1 with the lowest seed set number were from plant lines deriving from crosses HZ1.8-2.1 x BG16#25.1-1 and HZ2.6-1.1 x HZ3.1-15.1.

Taken together, these data indicate that incompatible crosses HZ1.8-2.1 x BG16#25.1-1 (PrsS1 x PrpS1) and HZ2.6-1.1 x HZ3.1-15.1 (PrsS3 x PrpS3) displayed the strongest SI phenotype. However, the F1 progeny resulting from these crosses exhibited distorted Mendelian segregation based on the seed germination on Kan and Basta MS media. Maternal PrSs expressing line HZ1.8-2.1 was homozygous, while the HZ2.6-1.1 line was
exhibiting ~4:1 segregation. The distorted and very low segregation could be due to reduced transmission through the male or female gametes and insertional inactivation of a gene expressed postmeiotically that is essential for the male and female gametophytic development or to epigenetic gene silencing and DNA methylation (Howden et al., 1998, Finnegan, 2004, Muskens et al., 2000).

In order to identify whether the PrsS was expressing correctly and to analyse the possibility of the reduced transmission through male or female gametes, the F₁ seeds should have been plated in a double selection medium (Basta and Kan) as well as a single selection media (Basta or Kan).

The in vivo results provided in this and previous sections lead to conclusion that PrsS was not functional in transgenic A. thaliana lines, due to low expression of the transgene and possible other effects that depend on the insertion of the transgene.

6.2.5. Investigating the function of the whole Papaver S₁ locus in transgenic A. thaliana

Transgenic A. thaliana line BG.6 expressing whole S₁-locus (described in previous section with segregation analysis) was systematically analysed in generation T₂. The in vivo analysis measured silique length and seed set. Results, presented in Figure 6.11 (a –seed set and b – silique length) show that majority of lines exhibit highly significantly reduced seed set, compared to non-transformed control.

A. thaliana non-transformed control had 58.6±7.9 seeds (n=30) and 13 lines out of 18 (72 %) exhibit lower seed set. The lines with the highest seed set reduction also exhibited decreased
silique length (11 lines, line BG6.14 and BG6.15 exhibited lower seed set but not lower silique length).

The plant line with the most affected seed set was BG6.5 (Figure 6.11.a), which had 23±5.7 seeds (n=30, P=1.5*10^-32 compared to Col-0) and it exhibited a 3:1 Mendelian segregation ratio for Kan insert. The lines that had seed set not significantly different from Col-0 were BG6.16 (exhibiting 2:1 segregation), BG6.17 (exhibiting 3:1 segregation), BG6.19 (exhibiting 3:1 segregation) and BG6.20 (exhibiting segregation <1:1). These implicate that the seed set number was not consistent with the segregation ratios, indication that there were
post-translational modifications, errors in gamete transmission or epigenetic gene silencing occurring.

In order to investigate the plant phenotype further, seed set of some plants was analysed in generation T3 after selfing. Plants were selected based on their low or fertility expression in generation T2. Those plants were BG6.1 with 44.1±9.0 seeds (n=30, P=1.2*10^-09 compared to Col-0; ~3:1 segregation ratio), BG6.2 that had 33.1±13.2 seeds (n=30, P=6.7*10^-14 compared to Col-0; 1:1 segregation ratio), BG6.5 with 23±5.7 seeds (3:1 segregation ratio). BG6.9, which had 50.8±6.4 seeds (n=30, P=2.6*10^-05 compared to Col-0; 0.2:1 segregation ratio) was chosen as a test control.

Figure 6.12 presents analysis of seed set on T3 generation of transgenic plants expressing S1-locus after self-pollination. Within each sub-line, 10 siliques were measured from 3 different plants.

![Figure 6.12](image)

**Figure 6.12:** Seed set analysis on the T3 generation of S1-locus expressing *A. thaliana* BG6 plants (blue bars) and non-transgenic Col-0 control (green bar). Error bars represent ± standard deviation.

If the reduced seed set was due to the functionality of the S1-locus, then it would be expected than in the next generation after selfing, plant fertility would be severely affected. The highest degree of seed set reduction or possibly non-fertile seeds, was expected in the case of
BG6.5 plants as they exhibited lowest seed set in generation T2. As the seeds were heterozygous in T2 generation with 3:1 segregation ratio, only the non-transgenic pollen would be able to pollinate S1 expressing stigmas and such seeds would not germinate on the MS with Kan selection marker. Therefore, the seed set in generation T2 was not expected to survive another round of selection if the SI system would be strongly exhibited.

However, results on figure 6.12 indicate that the plants are displaying various degrees of seed set reduction compared to non-transgenic plants. If we take BG6.5 plants for example, in generation T3 we counted 31.9±9.4 seeds in plant sub-line BG6.5-1 (n=30; P=8.3*10^{-14}), 33.9±11.9 seeds in BG6.5.10 (n=30; P=9.4*10^{-11}) and 36.9±8.8 seeds in BG6.5-11 (n=30; P=7.2*10^{-11}). In fact, they actually exhibited higher seed set number than the parental plant BG6.5 in generation T2. Lower seed set than parental line was observed in sub-line BG6.1, where 30.7±5.5 seeds were counted in BG6.1-1 plants (n=30, P=9.6*10^{-18}) and 30.4±7.6 seeds in BG6.1-4 (n=20, P=7.1*10^{-14}). Unusual pattern can also be observed with BG6.2 plants, where plant BG6.2-22 exhibit the highest seed set number 53.2±4.1 (n=30; P=0.13), which is not significant from the Col-0, although the BG6.2 line was the only one exhibiting expected segregation of the inserted whole S1-locus, 1:1. The other BG6.2 sub lines are highly significantly different from Col-0 and from its sister line (BG6.2-2 and BG6.2-10).

The sub-line BG6.9 also shows reduced seed set compared to Col-0, although the fertility reduction was comparable between the two sub-lines. BG6.9-1 had 38±4.9 seeds (n=30; P=1.6*10^{-12}) while BG6.9-3 had 47.3±8.2 seeds (n=20; P=1*10^{-03}). So, this data could indicate that the S1-locus might be functional when transformed into A. thaliana, however, some further detailed analysis for the presence of PrpS1 and PrsS1 transgene and transcript is required in order to confirm that.
In order to analyse SI on transgenic *A. thaliana* expressing $S_I$-locus further, a preliminary analysis of BG6 self-pollination was attempted and representative images presented on figure 6.13. For representation, only two lines were chosen: BG6.2-2, which exhibited reduced seed set and line BG6.22-2, whose seed set was not significantly different from non-transgenic.

**Figure 6.13.**: Aniline blue staining analysis of self pollination and control pollination of non-transgenic Col-0 on BG6 flowers (a) BG6.2-2 x BG6.2-2 – brightfield image; (b) BG6.2-2 x BG6.2-2; (c) BG6.2-2 x Col-0; (d) BG6.2-22 x BG6.2-22; (e) BG6.2-22 x Col-0. Scale represents 100 μm.
Figure 6.13.b shows that the inhibition of self-pollen occurred to some degree, but was not 100% (see figure 6.13.a). The rejection is not due to the non-receptive BG6.2-2 stigma, as can be seen on figure 6.13.c, as the pollination with Col-0 leads to development of long pollen tubes. On the other hand, self pollination of BG6.2-22, that exhibited normal seed set also exhibit ‘compatible’ self-pollination (Figure 6.13.d), which is comparable to the pollination with Col-0 pollen (Figure 6.13.e).

Due to the small size of analysed samples no firm conclusions could be made. The reduction of the seed production and decreased self-pollination in some BG6 lines, could indicate that the locus is functional. In order to analyse $S_1$-locus expressing A. thaliana lines (and therefore functionality of Papaver SI in A. thaliana) further, a more detailed analysis would be required, including systematic RT-PCR analysis and pollination assays.

### 6.3 DISCUSSION

Preliminary in vivo analysis of transgenic A. thaliana expressing stigmatic PrsS1 (HZ1) and PrsS3 (HZ2), pollen PrpS1 (BG16) and PrpS3 (HZ3) or $S_1$-locus (BG6) revealed that incompatible phenotype was not so easy to obtain. Some lines and crosses exhibited a reduced pollen tube growth, especially at the beginning of investigations. However, with time and experience it became obvious that age of plant and stage of stigma were crucial for this analysis. If plants were too old and producing more than 50% of their siliques, then their stigmas, despite the right biological stage, were not fully receptive, and the pollen was not germinating well, even on the receptive stigmas. (Nasrallah et al., 2002) illustrates the right biological stage of stigma and also demonstrates that maturity of pollen is very
important. They collected stigmas from stage 12, stage 13, early stage 14 (14E) and late stage 14 (14L) buds of transgenic *A. thaliana*, ecotype Colombia, expressing SRKb/SCRb. Stigmas were then pollinated with self or cross pollen. The results show that stigmas in the stage 12 or stage 14L allowed all the pollen to grow, regardless whether it was self (and therefore should be incompatible) or cross (compatible) pollen. However, in stage 13 and 14E, when stigmas reached their optimal biological stage, self-pollen was rejected and it’s growth inhibited (Nasrallah et al., 2002). As briefly described at the beginning of the result chapter, we emasculated buds in the stage 12, excised the stigmas and place them in 0.5 % agar where they were left to mature for 20-24 h before pollination. According to (Smyth et al., 1990), stage 12 lasts for 12 h in total so our pollinations were done in stage 13 (which lasts for 6 h) or early stage 14 (stage 14 lasts 18 h), when stigmas were not only receptive but should as well have maximum expression of PrsS. This technique differs from the other labs where they emasculated buds on plants and immediately pollinated them (Nasrallah et al., 2002). As *A. thaliana* pollen germinates well on stigma, the stigmas were collected stigmas for pollination assays after 2 h (Nasrallah et al., 2002). Despite the fast in vivo pollen germination, that takes ~15 minutes in *A. thaliana* (Boavida et al., 2005), we left pollen to germinate on stigma for 16 h, which was the optimal identified time for germination of *A. thaliana* flowers in vitro (Boavida and McCormick, 2007). This time-point was used for pollination assays in transgenic *A. thaliana* expressing *Papaver* genes as we did not know if, and what role the temporal expression of *Papaver* genes plays in *A. thaliana* in vivo pollen tube growth.

Nasrallah et al., (2002) demonstrated, that SI phenotype in transgenic *A. thaliana* expressing *Brassica S*-determinants was developmentally regulated. The temporal analysis on the expression of the *Papaver* female *S*-determinant, PrsS, was initially detected two days

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preanthesis, increased at one day preanthesis and remained high for several days after flower opening (Foote et al., 1994). In *Papaver*, inhibition of incompatible pollen occurs very quickly on the stigmatic papillae, within minutes of pollination (Franklin-Tong et al., 2002), so the same was expected to occur in *A. thaliana* expressing PrsS or whole *S1*-locus. *In vitro* pollen tube inhibition and viability assay in transgenic *A. thaliana* expressing *Papaver* pollen determinant PrpS demonstrated that the significant decrease of pollen viability occurred as quick as 2 h after incompatible pollen challenge (data not shown), and incompatible pollen tubes were inhibited and very short, comparable to the inhibited *Papaver* pollen tubes. Very informative were also the results of actin analysis where PrpS-expressing pollen was challenged with incompatible recombinant PrsS proteins. Actin foci were present mostly in *A. thaliana* pollen grains or grains with very short tubes, indicating that inhibition occurred very rapidly upon the SI interaction as no pollen tubes were observed, which was comparable to the *Papaver* SI response. Therefore it was expected that *in vivo* analysis of pollination of PrsS-expressing *A. thaliana* stigmas with its cognate PrpS-GFP pollen expressing *A. thaliana* would confirm the rapid inhibition of pollen on the stigma surface and show no or very little penetration of pollen through the stigma surface. The results obtained *in vivo* showed no inhibition of the incompatible pollen tube growth through PrsS expressing stigmas and such incompatible pollinations resulted in fertile seed set, indicating that there is no or very low expression of PrsS. Although the mRNA for PrsS was detected using RT-PCR on stage 13 flowers, the transcript levels of PrsS were lower compared to PrpS, and the PrsS protein was not detected using western blotting. Very low expression of PrsS could be explained by translation of low levels of mRNA into protein that was expressed below critical threshold in flowers; however, it was also possible that protein was not translated properly.
**In vivo** pollination assay and crosses for the seed set had a low reproducibility rate, as it was very difficult to assure the same conditions. It was impossible to cross-pollinate flowers with the same amount of pollen each time, and number of pollen was of great importance to ensure reproducibility and accuracy for the seed set data. To ensure that comparable crosses or repeats were pollinated with the same pollen mix of pollen from several different flowers was used. Pistils were also very susceptible to any sort handling damage that occurred mostly during the emasculation stage and this was the reason why there are less repeats of certain crosses.

With pollination assays, emasculated stigmas in early stage 13 were excised and placed in Elisa plate with 0.5 % agar, where they were left to mature, pollinated and finally collected in aniline blue containing tube. If there was any degree of pollen rejection and inhibition on incompatible crosses, then it was possible that inhibited pollen fell off the stigma or was washed away when stigma was placed in an eppendorf tube with aniline blue. Masters student Stephanie Smith, who continued with the pollination experiments, solved this issue with a temporarily “compartmentalised” microscope slide where stigmas were loaded before staining. With this technique, the visualisation of stigmas included total pollen used for pollination as in incompatible situation none got washed away or was lost as it fell of the stigma. However, no incompatible pollinations were observed due to low or no expression of PrsS in transgenic *A. thaliana*.

Intriguing results were obtained when PrsS₁ and PrsS₃ expressing *A. thaliana* stigmas were used to incubate *Papaver* S₁ and S₃ pollen. Pollen inhibition was observed in an S-specific manner and in addition also pollen tube tip swelling. This indicates that transgenic PrsS expressing *A. thaliana* stigmas are expressing PrsS protein, but most likely in low
concentrations as many stigmas were used for the stigmatic extract. It could also be possible that proteins are not properly folded or some other components of the stigmatic extract caused the *Papaver* pollen tube tip swelling, although the later explanation seems unlikely as non-transgenic Col-0 stigmatic extract was used as a control and no such dramatic pollen tube tip swelling was observed with it. Also, if the protein is not folded correctly we would not be expecting to observe such a response, as it was demonstrated *in vitro* (See Chapter 5) that heat denatured recombinant PrsS proteins had no effect on growth of pollen tubes.

Following successful *in vitro* analysis on PrpS expressing *A. thaliana* pollen, it was anticipated that *in vivo* analysis, although preliminary, might indicate stronger level of inhibition in incompatible pollinations. Most likely, the problem lies within the expression levels of PrsS in stigma. For successful incompatible response, the expression level of stigmatic gene is crucial and it was also demonstrated in the case of transgenic *A. thaliana* expressing *Brassica* SRK genes in the model of transgenic SI *A. thaliana* (Bi et al., 2000, Nasrallah et al., 2002, Boggs et al., 2009b). In transgenic *A. thaliana* expressing *Brassica* SI they solved the problem by transforming the SI system from more closely related SI *Arabidopsis lyrata* and *Capsella grandiflora* (Nasrallah et al., 2002, Boggs et al., 2009b, Nasrallah et al., 2007). In this light, the expression of the stigmatic genes in *A. thaliana* seems more problematic than expression of pollen genes. However, the proteins that are being utilised in *Papaver*-like transgenic *A. thaliana* SI model and *Brassica*-like transgenic model are completely different. In *Papaver*, PrsS are small globular, secreted stigmatic proteins, while in *Brassica* the stigmatic S-receptor kinase, SRK, is a single-pass transmembrane protein kinase. Structurally the proteins are very different, so the problem of
successful expression of foreign stigmatic genes in *A. thaliana* probably lies with the incorrect processing of the stigmatic determinants or in a weak expression.

The stigmas used for the *in vivo* experiments were homozygous but even if diploid stigmas would be heterozygous for PrsS, this would mean that they still contain one copy of the *PrsS* gene. When such flower is crossed with pollen that is homozygous for PrpS, the outcome of an incompatible cross should be 100% pollen inhibited (*PrsS* expressing stigma x *PrpS* pollen), if we assume, that pollen inhibition occurs on the stigmatic surface. The seeds of such cross (F1 progeny) would therefore be able to survive the Basta and Kan selection. So if incompatible crosses between PrsS and PrpS expressing *A. thaliana* lines (HZ1 x BG16 or HZ2 x HZ3) would be successful, then there would be no seed produced as the pollen should be inhibited on the surface of the stigma or at least very early as it would grow through the style. However, our crosses between PrsS and PrpS determinant of the cognate S-haplotype resulted in pollen tubes growing thorough the style and fertilizing the ovaries. The analysis of F1 generation seedlings arriving from some of the incompatible crosses presented in this section showed various segregation phenotype and seed set. The segregation was analysed on a double selection media, using both, Kan and Basta. Only the seedlings expressing both, PrsS and PrpS, could survive and it was therefore expected that surviving seedlings would have reduced seed set as a result of simultaneous expression of both, PrsS and PrpS. This was true for most of the analysed F1 lines. Significantly different seed set from Col-0 plants were observed in F1 plants deriving from cross between HZ1.1-3.1 x BG16#25.1-1, HZ1.8-2.1 x BG16#25.1-1 and HZ2.6-1.1 x HZ3.1-15.1 but not from two crosses of HZ2.13-5.1 x HZ3.1-15.1, whose seed set was not significantly different from Col-0. For more complete analysis the presence of the genes should be analysed by RT-PCR and the seeds from F1
progeny should be grown on individual selection media in order to give us more information about the nature of parental expression. It would also be important to perform Southern analysis in order to investigate number of inserts present as well as study the protein expression by Western analysis and immunolocalization. The lowest seed set was observed in F1 derived from HZ1.8-2.1 x BG16#25.1-1 (17.9±5 seeds) and HZ2.6-1.1 x HZ3.1-15.1 (15.9±4.2 seeds), which could indicate that these two have the highest expression of PrsS1 or PrsS3, respectively, and those lines would therefore be the most interesting for the future in vivo analysis of Papaver SI in transgenic A. thaliana.

Nevertheless, the in vivo data suggest that we ended with a “leaky” PrsS phenotype and despite all the above possibilities, the PrsS expression is very low (probably below the level required for inhibition) or the protein is not being properly folded. In order to obtain a more complete picture of PrsS expression in A. thaliana, the repeat of transformation with highly active stigma-specific promoter Stig1 from N.tabacum would be proposed or transformation with another stigma-specific promoter, such as SLG promoter from Brassica (Thorsness et al., 1993, Goldman et al., 1994).

Preliminary analysis of A. thaliana BG6 lines expressing whole $S_I$-locus revealed that majority of lines exhibited a significantly reduced seed set after self-pollination, while seed set of some lines was not affected after self-pollination (BG6.16, BG6.17, BG6.19 and BG6.20). If the transgenic BG6 A. thaliana were homozygous, then the seed set would not be expected. Therefore the poppy SI (using the $S_I$-locus) was probably leaky in A. thaliana. Most probably the problem lies within the protein expression levels as the protein translation is coupled with the endogenous promoter (the PrpS$_1$ in BG16 is coupled with a strong
ntp303 promoter, hence the high and strong expression). However, the RT-PCR analysis that was so far conducted on a limited number of BG6 samples, resulted in only PrsS mRNA present, and not PrpS; so the pollen seed set could also indicate that pollinations are due to the Col-0 phenotype of pollen (heterozygous) and that the reduced seed set occurred for different reasons. The place of insertion on the genome and the number of inserted copies, were also unknown and this might well be the contributing factor to the reduced fertility. Transgenic BG6 lines expressing whole $S_I$-locus require further detailed analysis. The RT-PCR analysis should be systematically carried on all the lines from BG6 plants, screening the PrsS and PrpS expression, as we cannot be certain whether the reduction of seeds was due to the functional SI.

The transgenic $A. thaliana$ model for Brassica SI did not express the whole $S$-locus, but the functional SRK-SCR gene pairs (Nasrallah et al., 2004). If the whole Papaver $S$-locus could be transferred to and be functional in a such an evolutionary distant species as $A. thaliana$, as was demonstrated for pollen determinant PrpS, then this could provide an information into evolution of SI. It would demonstrate a possible signalling mechanism being conserved in $A. thaliana$ that would probably open up a further debate not only about the evolution SI but also about the evolution of other recognition systems, such as plant-pathogen. Furthermore, this system could be used for plant breeding in a production of F1 hybrids where self-compatibility poses a problem that needs to be overcome. Furthermore, this Papaver-like SI system could potentially be also linked to different promoters for expression in somatic plant cells or even mammalian cells in order to use it as a potential inducible switch to trigger PCD, which would be of great interest with mammalian cancer cells. And most importantly, the $A. thaliana$ model exhibiting Papaver-like SI would represent a helpful and valuable tool
in studying the SI events on a detailed molecular level, as it would enable the crosses with various other lines of interest or analysis by mutations of various events, from the details of interaction between PrsS and PrpS to PCD.
CHAPTER 7

GENERAL DISCUSSION
The studies presented in this thesis cover the analysis of the initial events of SI, the interaction between the stigmatic S-determinant, PrsS and pollen S-determinant, PrpS, the preliminary investigation into the involvement of the ubiquitin-proteasomal pathway during SI, and *in vitro* and *in vivo* analysis of transgenic *A. thaliana*, expressing *Papaver* S-determinants.

*Papaver* SI model in Figure 7.1. summarizes the SI events including the S-specific interaction between PrpS and PrsS (presented in Chapter 4, published in Wheeler et al., 2009) and the model of UbP degradation pathway connected to DEVDase activity (presented in Chapter 3). The involvement of UbP pathway in *Papaver* SI is discussed in details in Chapter 3 and not dealt here.

![Figure 7.1: Schematic model of the SI response in *Papaver*. Upon the interaction of secreted stigmatic PrsS proteins and pollen transmembrane protein PrpS, a downstream signalling network is triggered, starting with an increase in K⁺ and [Ca²⁺], who is signalling to the inhibition of pyrophosphatase activity and pollen tube tip growth, activation of MAPK, depolymerization of actin and microtubule cytoskeleton and appearance of punctate actin foci, activation of ROS and NO and activation of caspase-like activities, leading to the PCD of the pollen tube. The model presents DEVDase activity that most likely acts in parallel with proteasomal degradation. DEVDases are presented as yellow circles within the proteasome, as identified by Hatsugai et al., (2009).](image)
7.1. *Papaver S*-determinant interactions

SI in *Papaver* is achieved when incompatible pollen grains lands on the stigma where it is recognized and discriminated from the genetically different pollen, and selectively inhibited. The stigmatic secreted PrsS proteins mediate this recognition and rejection, and act as a signalling ligand for the pollen transmembrane protein, PrpS (Foote et al., 1994, Wheeler et al., 2009). The discovery of PrpS as the pollen determinant of the SI response in *Papaver* represented a breakthrough in the SI research and the long sought missing link required for the recognition and rejection of the incompatible pollen.

The results presented in this thesis demonstrate the S-specific binding between PrpS and PrsS using the far western blot and the peptides designed to mimic the extracellular domain. The extracellular domain was predicted to be positioned between TM 2 and 3 using a variety of prediction programmes and was proposed to be the interacting region for the secreted ligand PrsS (Wheeler et al., 2009). The binding was further indicated when the peptides designed over the extracellular domain, alleviated the incompatible pollen inhibition when added to the *in vitro* bioassay (Wheeler et al., 2009). The far western method is well established and widely used for determining protein interaction sites in receptor-ligand interaction as well as the use of the synthetic peptides (Volkmer et al., 2011). Far western approach was used to confirm many protein-protein interactions, for example, to demonstrate the interaction in plant-pathogen system between rice Pi-ta receptor and host AVR-Pita protein (Jia et al., 2000) or to identify the S-specific interaction between stigmatic microsomal fraction and recombinant SCR in *Brassica* (Kemp and Doughty, 2007).
The PrpS represents a novel transmembrane protein and shares no homology to any other protein, so the exact mechanism of the perception is not yet known. PrpS could be part of a greater multimeric complex where it could play a role of the $S$-specific signal perception protein but not the downstream transduction. An example of such complex is CLV complex where LRR RLKs CLV1 and CLV2 dimerize in order to form a receptor complex (Clark et al., 1995, Jeong et al., 1999). CLV1 binds the CLV3 derived CLE ligand through its extracellular domain and functions in stem cell specification (Clark et al., 1995, Clark, 2001, Stahl and Simon, 2005, Ogawa et al., 2008). CLV1 and CLV3 are expressed only in the central zone of SAM (Clark et al., 1997, Fletcher et al., 1999). Below the central zone a control zone is located where another protein, implicated in SAM is expressed. Transcription factor WUSCHEL (WUS) forms a feedback network with CLV (Schoof et al., 2000). WUS upregulates CLV3 when the number of stem cells is low, while CLV3 binds to CLV1 and downregulates WUS when the stem cells are in abundance (Reddy and Meyerowitz, 2005).

If PrpS forms a part of a larger protein complex, it is yet unknown. However, the $S$-protein binding protein (SBP) was previously identified that apparently binds to the PrpS, but not in the $S$-specific manner (Hearn et al., 1996). The connection between PrpS and SBP is not yet analysed as no protein or gene has yet been identified for SBP. Alternatively, the PrpS could perceive and transduct a signal on its own. Some analogies might be drawn between the *Drosophila* protein Flower (Yao et al., 2009) or the mammalian CRAC calcium channel (Luik et al., 2008) and PrpS. Protein Flower is TM protein with 4 predicted TM domains and is involved in the endocytosis of the synaptic vesicles. It displays properties of $\text{Ca}^{2+}$ channel: negatively charged aminoacids in the TM domains, which play a key role in the ion selectivity. Similar to Flower, the 4 TM protein CRAC/Orai also carries negatively charged aminoacids in the TM domains and multimerizes to form a channel. In this process each
monomer contributes one TM domain with the negatively charged aminoacid to the pore (Yao et al., 2009, Prakriya et al., 2006). Flower proteins also forms multimers and is thus able to form a calcium channel by which it controls Ca\textsuperscript{2+} influx (Yao et al., 2009). Protein alignment of PrpS proteins revealed conserved glutamic acid residue on position 75. This region is mostly predicted to be extracellular in PrpS\textsubscript{1} and PrpS\textsubscript{8}, and intracellular in PrpS\textsubscript{3}, although some prediction programmes also gave inverted prediction, but none predicted a TM domain. Concluding from the analysis conducted, this region is involved in the PrpS interaction (Wheeler et al., 2009). Interestingly, this glutamic acid residue is also conserved with Flower protein, when PrpS and Flower are aligned, although in Flower the region is predicted TM. The negatively charged aminoacids appear to be crucial in the TM domains for the formation of the voltage gated calcium channels. Alignment of PrpS proteins revealed in total 7 conserved negatively charged aminoacids: three glutamic acids (E75, E129 and E168), and three aspartic acid (D39, D90, and D123) with some being the candidates for the potential pore formation. Additional motif that is well known to be involved in the protein oligomerization and formation of the membrane channel is the GxxxG motif (Dawson et al., 2002, Senes et al., 2004, McClain et al., 2003). Helicobacter pylori vacuolating toxin can form an anion-selective membrane channel as it contains three tandem GxxxG motifs, characteristic for TM dimerization and membrane channel formation (McClain et al., 2003). The PrpS alignment revealed two conserved tandem GxxxG motifs in positions 126 and 130, with the first one also conserved in Flower. This region is also not predicted to be part of TM domain but the long cytoplasmic tail. More detailed analysis of the residues suggests fourth TM domain, so the motif is most likely part of the fourth TM domain. The GxxxG motif in position 126 also contains conserved glutamic acid residue.
Taken together, PrpS S-specifically recognizes the PrsS proteins and it most likely acts as a receptor. One of the first downstream events following the interaction is the increase in the cytosolic calcium (Franklin-Tong et al., 1993, Franklin-Tong et al., 2002), and it was also demonstrated that the SI activates ligand-gated nonspecific cation conductance (Wu et al., 2011), so the PrpS might form a channel, composed of PrpS multimers or a channel complex where it would form a subunit. This is something that is currently being investigated in the Franklin-Tong’s lab.

7.2 Functional analysis of *Papaver rhoeas* self-incompatibility in transgenic *Arabidopsis thaliana*

We initiated studies to see whether the *Papaver* SI system might be functionally transferable to other plant species and studies in this thesis describe the successful interspecific transfer of the *Papaver* pollen determinants of SI, PrpS, to SC *A. thaliana*. Several transgenic *A. thaliana* lines were created, expressing the PrpS fused to the GFP: PrpS1-GFP and PrpS3-GFP, and stigmatic S-determinants PrsS1 and PrsS3 under the control of pollen and stigma specific promoters, respectively, and lines expressing whole S1 locus or PrpS1, under the control of the native promoter and constitutive 35S promoter, respectively. Our results “in vitro” revealed an S-specific inhibition of the PrpS-GFP expressing transgenic *A. thaliana* pollen tube growth by recombinant PrsS, and key hallmark features for *Papaver* SI: loss of pollen viability, appearance of actin foci and increase in DEVDase activity. These data suggest that PrpS expressed in evolutionary very distantly related species *A. thaliana*, is functional and that only PrpS and PrsS are required for a fully functional SI response in *A. thaliana* pollen, where all the required components for the pollen tube inhibition are present.
However, preliminary *in vivo* assay demonstrated that PrsS exhibits weaker expression in *A. thaliana* flowers. Although the full SI response was not demonstrated *in vivo*, we observed significant inhibition of *Papaver* pollen with protein extract from PrsS expressing *A. thaliana* stigmas, leading to the conclusion of low expression of PrsS in transgenic *A. thaliana*.

Our study represents the first such study of transforming *A. thaliana* with the GSI determinants. Transformation of *A. thaliana* by determinants of *Brassica* SI is already developed by Nassrallah lab and studies on *A. thaliana* expressing functional intact or chimeric SRK/SCR pairs are underway to identify the mechanism of SI and to elucidate components of pollen-pistil signalling (Tantikanjana et al., 2010, Nasrallah et al., 2002, Rea et al., 2010). *SRK-SCR* receptor-ligand gene pair used for the studies of *Brassica* SI were isolated from the SI crucifers *Arabidopsis lyrata* or *Capsella grandiflora*, and the SI analysed by performing crosses between them (Nasrallah et al., 2002, Boggs et al., 2009b, Rea et al., 2010, Nasrallah et al., 2004). Although *Brassica* model of SI in *A. thaliana* took several years to develop and optimize, it now represents invaluable system for the studies of the mechanism of SI and pollen-pistil interactions, as well as evolutionary aspects of SI (Nasrallah et al., 2002, Nasrallah et al., 2004, Rea et al., 2010). The transgenic *A. thaliana* expressing SRK-SCR gene pair was used for example for the analysis of the residues important for the SRK specificity and therefore the SI response. Site-directed mutagenesis approach was used on the polymorphic aminoacid residues in the extracellular domain of SRK and six were identified as crucial for ligand-specific SI activation (Boggs et al., 2009a). Although in *Papaver* we have identified the extracellular domain of PrpS1, required for the SI response, and 15 aa domain that is crucial for the interaction with PrsS1, it was so far impossible to narrow down the S-specificity to the individual amino acids. The transgenic *A.
thaliana pollen expressing PrpS is a model for Papaver-like SI that now enables the possibility to investigate the specific residues within PrpS, determining the specificity of the ligand binding interaction or protein multimerization.

Papaver belongs to the basal order in the lineage Ranunculales, diverged very early in the history of eudicots (Allen and Hiscock, 2008). The evolutionary distance between Ranunculales and Brassicales, lineage where belongs A. thaliana, is estimated to 140 million years (Bell et al., 2010). This suggests that Papaver SI response pre-dates other SSI mechanism as well as other GSI and it may represent the ancestral state of SI. It is also possible that SSI system evolved from the SC species originating from the GSI ancestors (Allen and Hiscock, 2008). Although A. thaliana operated Brassica-type of SI and it diverged from A. lyrata ~10 million years ago (Hu et al., 2011), the transfer of the Papaver pollen S-determinant was functional. A. thaliana pollen had the cellular machinery required to elicit a S-specific pollen rejection upon the PrpS-PrsS interaction. This could indicate that an ancestral SI mechanism, operating in Papaver might still be conserved in other species, although they evolved more complex mechanisms of SI (for review see McClure et al., (2011), Tantikanjana et al., (2010)) but might have kept basic components and recruited this mechanism for different purposes. The recent advances of the research on the SI induced actin reorganisations in different SI systems suggest, that there could be a link between the mechanistically different SI systems and the SI could be more conserved than known so far (Liu et al., 2007c, Iwano et al., 2007, Wang et al., 2010, Poulter et al., 2010a).

The preliminary analysis to investigate whether the Papaver-like SI response could be induced in any plant tissue was tested using A. thaliana leaf mesophyll protoplasts, and the results so far confirmed the functionality of the PrpS-PrsS system.
Ca$^{2+}$ is mediating downstream events of SI, observed in incompatible *Papaver* pollen tube, however, it is also a well known and conserved secondary messenger in other plant species, mammals and also bacteria, relaying the signals, perceived by the receptors on the cell surface to target molecules in the cytosol (Shemarova and Nesterov, 2005, Ma and Berkowitz, 2007). It is very important, for example, for the signal transduction during plant-pathogen interactions, as Ca$^{2+}$ increase is one of the first steps during plant immune response to the PTI and ETI, signalling to the PCD. Considering many similarities between SI and HR as well as animal histocompatibility, such as aforementioned Ca$^{2+}$ signalling to phosphorylation of target proteins, alterations to the cytoskeleton, activation of ROS and induction of the DEVDase activities leading to PCD, might indicate that these targets are ancient and conserved, and it would be interesting to investigate the evolutionary connection between them.
CHAPTER 8

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## APPENDIX I

Clustal W alignment of the PrpS1, PrpS3, PrpS8 and Flower protein (CG6151).

Identical amino acids are marked with “*”, conserved substitutions with “:” and semi-conserved substitutions with “.". Underlined region of PrpS1 is the predicted extracellular domain; highlighted in yellow are the conserved aminoacid residues, potentially involved in channel formation; highlighted in red are the glycines forming GxxxG motif also potentially involved in the channel formation. Clustal W source:

http://www.ebi.ac.uk/Tools/clustalw2/index.html

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<td>PrpS3</td>
<td>MPRNHIAYVFVFVFVTFLGVAFLVRINKSALTWKLATAFVVVLIVLAVGGGVSILM 60</td>
</tr>
<tr>
<td>CG6151</td>
<td>-MSFAEKITGGLARPWQDPIGPQPHLYLKVSSRLGTVAAPFA1LPGLWNVS-IIITLS 58</td>
</tr>
<tr>
<td>PrpS1</td>
<td>CVKLLGLVLHRLS-----FSEDQKWVVAFGTAAICDVLLVPK                 115</td>
</tr>
<tr>
<td>PrpS8</td>
<td>YLKLLGWVLQHLT-----VTENGKVVAOTATYFLVFATIVTPITSICFLSSIMICV 115</td>
</tr>
<tr>
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<tr>
<td>CG6151</td>
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</tr>
<tr>
<td>PrpS1</td>
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<td>PLNFPVGVHSSHRGVEV 193</td>
</tr>
<tr>
<td>PrpS3</td>
<td>S-AAILVGNNASPCPNEA- 191</td>
</tr>
<tr>
<td>CG6151</td>
<td>N-NAQPSFTGAVTGDSNV 194</td>
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a) Family tree of the PrpS<sub>1</sub>-GFP expressing <i>A.thaliana</i> line BG16. The underlined lines were analysed by RT-PCR and the lines outlined in red were identified as the highest expressing lines.
b) Family tree of the PrpS3-GFP expressing *A.thaliana* line HZ3. The underlined lines were analysed by RT-PCR and the lines outlined in red were identified as the highest expressing lines.
c) Family tree of the PrsS₁ expressing *A. thaliana* line HZ1. The underlined lines were analysed by RT-PCR and the lines outlined in red were identified as the highest expressing lines.

![Family tree of the PrsS₁ expressing A. thaliana line HZ1](image)
d) Family tree of the PrsS₃ expressing *A.thaliana* line HZ2. The underlined lines were analysed by RT-PCR and the lines outlined in red were identified as the highest expressing lines.

![Family tree of the PrsS₃ expressing A.thaliana line HZ2.](image)

**Plant generation**

**Insert in Arabidopsis:**

*Stig1::PrsS₃*

(Basta resistance)

<table>
<thead>
<tr>
<th>Plant generation</th>
<th>A.t.HZ2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>T₀</strong></td>
<td>A.t.HZ2</td>
</tr>
<tr>
<td><strong>T₁</strong></td>
<td>HZ2. [1-20]</td>
</tr>
</tbody>
</table>
| **T₂**           | HZ2.1 [1-9]*  
|                  | HZ2.2 [1-9]*  
|                  | HZ2.3 [1-9]*  
|                  | HZ2.4 [1-9]*  
|                  | HZ2.5 [1-10]  
|                  | HZ2.6 [1-10]  
|                  | HZ2.7 [1-10]  
|                  | HZ2.10 [1-10] |
|                  | **self**    |
| **T₃**           | HZ2.6.1 [1-13] |
|                  | **self**    |
| **T₄**           | HZ2.6.1.1 [1-22]  
|                  | **self**    |
| **T₅**           | HZ2.6.1.1.1 [1-9]  
|                  | **self**    |

[1-20]

[1-13]

[1-18]

[1-22]

[1-17]

[1-9]

[1-10]
APPENDIX III

Published papers


My contribution: Natalie Poulter carried out most of the studies. I carried out some of the experiments in collaboration doing my final year project for University of Ljubljana, Slovenia. I helped with caspase assays presented in the paper, tube length studies as well as western blot (Figure 3). I proof-read the manuscript.
Microtubules Are a Target for Self-Incompatibility Signaling in *Papaver* Pollen

Natalie S. Poulter, Sabina Vatovec, and Veronica E. Franklin-Tong*

School of Biosciences, University of Birmingham, Edgbaston, Birmingham B15 2TT, United Kingdom

Perception and integration of signals into responses is of crucial importance to cells. Both the actin and microtubule cytoskeleton are known to play a role in mediating diverse stimulus responses. Self-incompatibility (SI) is an important mechanism to prevent self-fertilization. SI in *Papaver rhoeas* triggers a Ca\(^{2+}\)-dependent signaling network to trigger programmed cell death (PCD), providing a neat way to inhibit and destroy incompatible pollen. We previously established that SI stimulates F-actin depolymerization and that altering actin dynamics can push pollen tubes into PCD. Very little is known about the role of microtubules in pollen tubes. Here, we investigated whether the pollen tube microtubule cytoskeleton is a target for the SI signals. We show that SI triggers very rapid apparent depolymerization of cortical microtubules, which, unlike actin, does not reorganize later. Actin depolymerization can trigger microtubule depolymerization but not vice versa. Moreover, although disruption of microtubule dynamics alone does not trigger PCD, alleviation of SI-induced PCD by taxol implicates a role for microtubule depolymerization in mediating PCD. Together, our data provide good evidence that SI signals target the microtubule cytoskeleton and suggest that signal integration between microfilaments and microtubules is required for triggering of PCD.

The plant cytoskeleton comprises actin microfilaments and tubulin microtubules that are highly dynamic through their interaction with various actin-binding proteins and microtubule-associated proteins (Erhardt and Shaw, 2006; Hussey et al., 2006). Both actin microfilaments and cortical microtubules play a key role in determining cell shape and growth, and recent work has provided valuable insights (Smith and Oppenheimer, 2005). There is now considerable evidence that the plant actin cytoskeleton plays a key role in modulating signal-response coupling, with many examples of actin mediating various biotic and abiotic responses (Staiger, 2000). Cortical microtubules are also involved in signal-response coupling. It has been shown that abiotic stimuli, such as gravity (Himmelspach et al., 1999), hormones (Shibaoka, 1994), freezing (Bartolo and Carter, 1991), and salt stress (Shoji et al., 2006), result in the reorientation or depolymerization of microtubules. Biotic interactions resulting in microtubule alterations also exist. Plant interactions with pathogenic fungi and symbiotic interactions with mycorrhizal fungi and rhizobia are known to stimulate microtubule reorganization (for review, see Wasteynes and Galway, 2003; Takemoto and Hardham, 2004).

Self-incompatibility (SI) is a genetically controlled system to prevent self-fertilization in flowering plants. A multi-allelic *S*-locus is responsible for specifying *S*-specific pollen rejection to allow discrimination between incompatible and compatible pollen. Interaction of pollen *S* - and pistil *S*-determinants that have matching alleles allows "self" (incompatible) pollen to be recognized and rejected, while compatible pollen is allowed to grow and set seed. In this way, SI provides an important mechanism to prevent inbreeding through specific recognition and rejection of incompatible pollen. Several different SI systems exist; they have quite distinct molecular and genetic control; thus, different mechanisms are involved in SI in different species (for review, see Takayama and Isogai, 2005; McClure and Franklin-Tong, 2006).

In *Papaver rhoeas*, the pistil part of the *S*-locus encodes small, approximately 15-kD proteins that act as signaling ligands named *S* proteins (Foote et al., 1994). Their interaction with incompatible pollen triggers *S*-specific increases of cytosolic-free calcium concentration ([Ca\(^{2+}\); Franklin-Tong et al., 1993). The SI-induced Ca\(^{2+}\)-dependent signaling network comprises several intracellular events in incompatible pollen, indicating quite complex networks of interconnected events involved in the SI response. Ca\(^{2+}\)-dependent phosphorylation of a cytosolic pollen soluble inorganic pyrophosphatase (sPPase), Pr-p26.1 (de Graaf et al., 2006), inhibits its sPPase activity. As sPPases are important enzymes for driving biosynthesis, they are crucial for cell growth, so SI, by targeting this enzyme, results in pollen tube inhibition. SI also triggers reorganization and depolymerization of the F-actin cytoskeleton (Geitmann et al., 2000; Snowman et al., 2002). As the actin cytoskeleton is required for pollen tube growth (Gibbon et al., 1999), this represents another

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1 This work was supported by the Biotechnology and Biological Sciences Research Council (to V.E.F.-T. and a studentship to N.S.P.).

S.V. worked as an undergraduate project student from the University of Ljubljana, Slovenia.

* Corresponding author; e-mail v.e.franklin-tong@bham.ac.uk.

** The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (www.plantphysiol.org) is: Veronica E. Franklin-Tong (v.e.franklin-tong@bham.ac.uk).

www.plantphysiol.org/cgi/doi/10.1104/pp.107.107052
mechanism to inhibit incompatible pollen tube growth. SI also triggers programmed cell death (PCD), involving several caspase-like activities (Thomas and Franklin-Tong, 2004; Bosch and Franklin-Tong, 2007). PCD is a conserved mechanism to get rid of unwanted cells and is used to sculpt tissues during development as well as in response to abiotic stress and pathogens (van Doorn and Woltering, 2005). SI activates PCD specifically in incompatible pollen, thereby preventing self-fertilization. Recent investigations revealed that alterations in actin dynamics can push pollen tubes into PCD (Thomas et al., 2006), and an SI-activated mitogen-activated protein kinase (Rudd et al., 1996) is implicated in signaling to PCD (Li et al., 2007). These data suggest that in *Papaver*, these components contribute to an integrated SI signaling network to achieve inhibition and death of incompatible pollen.

While the actin cytoskeleton is well established as being essential for tip growth in plant cells (Gibbon et al., 1999; Staiger, 2000), the role of the microtubule cytoskeleton is more variable, depending to some extent on the cell type. In some tip-growing plant cells, microtubule-disrupting drugs have no effect on tip growth; in others, they result in inhibition of growth, multiple growth initiation sites, or loss of directionality in root hairs (Bibikova et al., 1999) and pollen tubes (Anderhag et al., 2000; Gossot and Geitmann, 2007). However, it is well established that microtubules do not play an obvious role in regulating angiosperm pollen tube growth rate (Heslop-Harrison et al., 1988; Aström et al., 1995; Raudaskoski et al., 2001). Apart from data showing that they help organize the generative cell (GC) and vegetative nucleus (Raudaskoski et al., 2001; Laitiäinen et al., 2002), relatively little is known about their function (Cai and Cresti, 2006). As the actin cytoskeleton is known to play a role in SI, we speculate that microtubules are a target for SI-induced signaling. Here, we report that SI stimulates rapid and massive apparent microtubule depolymerization, demonstrating that the pollen microtubule cytoskeleton is an early target for SI signals. Our data implicate signal integration between the microfilament and microtubule cytoskeleton and suggest a role for microtubules in SI-induced PCD.

### RESULTS

**Microtubule Cytoskeleton Organization in Growing *Papaver* Pollen Tubes**

The microtubule cytoskeleton organization in normally growing *P. rhoes* pollen tubes, using immunolocalization and probing with α-tubulin (Fig. 1), has previously been described (Gossot and Geitmann, 2007). The microtubule arrangement is very similar to that described previously (Aström et al., 1995; Gossot and Geitmann, 2007). The tip region is relatively microtubule-free; behind this region are arrays of short, longitudinally organized microtubule bundles (Fig. 1A). Further back, in the shank region, there are longer, more regularly organized longitudinal microtubule bundles (Fig. 1, A and B), which are mainly cortical (Fig. 1, B and C). Pollen tubes have a vegetative nucleus and a GC, which has a distinctive population of spindle-shaped GC microtubules (Fig. 1B).

**SI Triggers Microtubule Depolymerization**

To establish whether microtubules are a target for SI signaling, we examined the microtubule cytoskeleton using immunolocalization at various time points after incompatible SI induction (Fig. 2). Typical microtubule and microfilament organization was seen in control pollen tubes (Fig. 2, A and B). The microtubule cytoskeleton was rapidly altered after SI induction. As early as 1 min after SI, cortical microtubule bundles were virtually undetectable in incompatible pollen tubes; much weaker staining suggested that they had depolymerized (Fig. 2C). The GC spindle-shaped microtubules remained relatively intact at this time point (Fig. 2D). F-actin also dramatically reorganized by 1 min and accumulated in the tip, where it is not normally detected; many of the filament bundles had disappeared (Fig. 2E). At 3 min, the cortical microtubule bundles were virtually undetectable (Fig. 2F), and F-actin appeared disintegrated (Fig. 2G). At 30 min, cortical microtubules remained depolymerized (Fig. 2H), the GC spindle-shaped microtubules were still evident but disintegrating (Fig. 2I), and F-actin was aggregating (Fig. 2J). These data demonstrate that SI induces very rapid alterations to the cortical microtubule organization.

![Figure 1. Microtubule organization in untreated *Papaver* pollen tubes.](image-url)
bule cytoskeleton of incompatible pollen tubes, which appeared to be depolymerized. The spindle-shaped microtubules were more or less intact. At 1 min after SI, F-actin is in the apical region; many F-actin bundles have disappeared. F, At 3 min after SI, cortical microtubules are undetectable; spindle-shaped microtubules show signs of disintegration. G, At 3 min after SI, F-actin has formed small punctate foci. H, At 30 min after SI induction, cortical microtubules are undetectable. I, At 30 min after SI induction, GC microtubules are further degraded. J, At 30 min after SI induction, F-actin comprises larger punctate foci. Microtubules were detected using immunolocalization with anti-α-tubulin; F-actin was colocalized using rhodamine-phalloidin. Images are full projections of confocal sections. Scale bar = 10 μm.

Figure 3. SI does not trigger tubulin degradation. Western blot of extracts from untreated pollen tubes (UT) and extracts from SI-induced pollen tubes at 15, 30, and 60 min after SI, probed with anti-α-tubulin antibody clone B-5-1-2. Overall α-tubulin levels (equal loading of samples; arrowhead) did not significantly change.
Actin Alterations

Microtubule Depolymerization Is Not Required for Microtubule Depolymerization

Actin Stabilization Prevents or Delays SI-Induced Microtubule Depolymerization

To investigate further whether actin depolymerization plays a role in the SI-induced apparent microtubule depolymerization, we stabilized F-actin using jasplakinolide (Jasp) and then induced SI. We reasoned that if actin depolymerization was important for microtubule depolymerization, stabilizing actin should prevent or delay this event. Untreated pollen tubes showed normal microtubule configurations (Fig. 5A); 30 min treatment with 0.5 \(\mu M\) Jasp, which causes bulbous tips due to actin stabilization/reorganization (Thomas et al., 2006), stimulated reorganization, but not depolymerization, of microtubules (Fig. 5B). After SI, microtubules were rapidly depolymerized by 1 to 3 min (see Fig. 2, C and F). Ten minutes after SI, microtubules were completely depolymerized (Fig. 5C), but with a pretreatment of 0.5 \(\mu M\) Jasp 30 min prior to SI induction, at 10 min post-SI significant remnants of microtubules remained (Fig. 5D). Thus, Jasp-mediated stabilization of F-actin alleviated or delayed SI-induced microtubule depolymerization, providing further evidence consistent with the notion that F-actin depolymerization signals to microtubule depolymerization during SI.

Microtubule Depolymerization Is Not Required for Actin Alterations

Because actin depolymerization results in microtubule depolymerization, this suggested cross talk between actin and tubulin. As the response was rapid, we wondered whether microtubules might signal to actin. We therefore examined the effect of microtubule depolymerization on the pollen tube actin cytoskeleton, using oryzalin to artificially depolymerize tubulin. The relatively high concentrations used were to ensure that the SI effect of rapid depolymerization within a couple of minutes was mimicked as closely as possible. After 5 min treatment with 10 \(\mu M\) oryzalin, no cortical microtubules were evident (Fig. 6A); there was no detectable effect on the actin cytoskeleton (Fig. 6B). Even after 30 min treatment with oryzalin, when cortical microtubules were undetectable (Fig. 6C), F-actin organization appeared normal (Fig. 6D). To confirm that oryzalin did not affect actin, we measured pollen tubes, as actin depolymerization inhibits pollen tube growth (Gibbon et al., 1999). For pollen tubes treated with 10 \(\mu M\) oryzalin for 60 min, mean lengths were 293 ± 10 \(\mu m\), compared with 314 ± 11 \(\mu m\) for untreated controls (n = 3 independent experiments). These values were not significantly different from each other (P = 0.156, nonsignificant), establishing that oryzalin had no effect on actin. Our data demonstrate that microtubule depolymerization does not stimulate actin depolymerization in pollen tubes, confirming data from Gossot and Geitmann (2007). Because these high levels of oryzalin do not affect actin or growth, we can be reasonably sure that possible side-effects are not an issue. This suggests there is one-way cross talk from actin to tubulin cytoskeleton, but not vice versa.

We also investigated whether stabilizing microtubules with taxol might affect actin reorganization. Taxol inhibits microtubule dynamics, causing stabilization of microtubules (Blagosklonny and Fojo, 1999), and is effective in plant cells (Baskin et al., 1994; Collings et al., 1998). Taxol does not dramatically affect microtubule organization, but some bundling is generally observed (Collings et al., 1998). Taxol, as expected, did not stimulate any major alterations to the organization of either the microtubule (Fig. 7, A and B) or actin microfilament (Fig. 7, C and D) cytoskeleton of pollen tubes, but the microtubule bundles were slightly larger and brighter, suggesting stabilization. The concentrations of taxol used are in line with other studies (see, e.g. Collings et al., 1998). To confirm that

Figure 4. Actin depolymerization triggers changes in microtubule organization and apparent depolymerization. A to D, Typical untreated pollen tube cytoskeleton organization. A and B, F-actin; C and D, cortical microtubules. E to H, Pollen tubes treated with 1 \(\mu M\) LatB for 5 min. F-actin in the apical (E) and shank (F) region is fragmented. G and H, Cortical microtubules in the apical and mid-region are short and disorganized (G), while cortical microtubules in the shank region are relatively undisturbed (H). I to L, Pollen tubes treated with 1 \(\mu M\) LatB for 30 min. F-actin in the apical (I) and shank (J) region is extensively fragmented, indicating depolymerization. K, Microtubules in the apical and mid-region are virtually undetectable. L, Microtubules are virtually undetectable, with a few fragmented bundles remaining. Microtubules were detected using anti-a-tubulin; F-actin was colocalized using rhodamine-phalloidin. Images are full projections. Scale bars = 10 \(\mu m\).
taxol had no effect on the actin cytoskeleton, we measured pollen tubes after treatment with 5 and 10 μM taxol for 1 h. The mean pollen tube lengths were 194.6 ± 9.8 μm and 194.0 ± 9.1 μm, respectively, compared to 190.0 ± 8.1 μm for untreated controls (n = 3). Thus, taxol had no significant effect on pollen tube growth (P = 0.717, 0.744, respectively, nonsignificant), consistent with taxol not having an effect on the actin cytoskeleton. To investigate whether stabilizing microtubules affected the ability of actin to depolymerize, we pretreated pollen tubes with 5 μM taxol for 30 min and then added 1 μM LatB for 30 min (Fig. 7, E–H). Although the organization of the microtubules in the apical region was disturbed (which is expected, as LatB inhibits pollen tube growth), the shank microtubules appeared relatively normal (Fig. 7, E and F), but F-actin depolymerized as normal (Fig. 7, G and H). Thus, microtubule depolymerization, although it accompanies actin depolymerization, is not required for actin depolymerization in pollen tubes. This confirms our data suggesting one-way signaling from actin to tubulin cytoskeleton.

Disruption of Microtubule Dynamics Does Not Trigger PCD

We previously demonstrated that actin depolymerization or stabilization can trigger PCD in pollen tubes (Thomas et al., 2006). Because SI also stimulated apparent microtubule depolymerization, we wondered whether microtubule depolymerization might also signal to PCD. We investigated this using oryzalin to depolymerize, or taxol to stabilize, pollen tube microtubules. Pollen tubes were treated with 10 μM oryzalin or 5 μM taxol and extracts tested for caspase-3-like activity using Ac-DEVD-AMC, a caspase-3 substrate, which we have used previously (Bosch and Franklin-Tong, 2007; Li et al., 2007). Untreated pollen tube extracts exhibited low DEVDase activity. The DEVDase activities in oryzalin- and taxol-treated pollen tube extracts were not significantly different from the untreated controls (P = 0.9581 and 0.6286, respectively; n = 4). Thus, microtubule depolymerization or stabilization alone clearly does not trigger PCD in *Papaver* pollen.

SI-Induced PCD Requires Depolymerization of Microtubules to Progress

Although changes in microtubule dynamics alone are not sufficient to signal to PCD, we wondered whether tubulin depolymerization might be required in conjunction with actin depolymerization to allow progression into SI-induced PCD. As microtubule depolymerization accompanies actin depolymerization, this was an important point to establish. We investigated whether pollen tubes with stabilized microtubules prior to SI-induced actin depolymerization affected entry into PCD. Pollen tubes were pretreated with 5 μM taxol, SI was induced, and extracts were

![Figure 5](image_url)

**Figure 5.** Actin stabilization by Jasp alleviates or delays SI-induced apparent microtubule depolymerization (A). Typical untreated pollen tube microtubule organization. B, Microtubule organization 30 min after Jasp treatment. C, Microtubules were completely depolymerized 10 min after SI. D, Microtubules were detectable after 30 min Jasp pretreatment followed by 10 min SI induction. Microtubules were detected with anti-α-tubulin. Images are full projections. Scale bar = 10 μm.

![Figure 6](image_url)

**Figure 6.** Microtubule depolymerization does not trigger alterations to the actin cytoskeleton. A and B, A 5-min treatment with 10 μM oryzalin. A, Apparent complete depolymerization of cortical microtubules. B, No apparent effect on F-actin organization. C and D, A 30-min treatment with 10 μM oryzalin. C, Cortical microtubules are apparently completely depolymerized. D, No apparent effect on F-actin organization. Microtubules were detected with anti-α-tubulin; F-actin was colocalized using rhodamine-phalloidin. Images are full projections. Scale bar = 10 μm.
assayed for DEVDase/caspase-3-like activity. Untreated pollen tube extracts exhibited low DEVDase activity, while SI induced high DEVDase activity (72.5% higher than untreated samples), which was significantly different from the controls (P < 0.001, ***; n = 10). In pollen tubes pretreated with taxol prior to SI induction, the level of DEVDase activity was significantly reduced; 41% lower compared to SI alone (P = 0.0256, *; n = 10). The reduction in DEVDase activity by taxol firmly implicates that microtubule depolymerization plays a role in mediating SI-induced PCD in addition to actin depolymerization. Moreover, when pollen tubes were pretreated with oryzalin for 30 min prior to SI induction, there was no significant difference in the DEVDase activity compared with SI-induced samples (P = 0.7079; n = 5). Together with the results from the taxol treatment, this is consistent with the idea that microtubule depolymerization is involved in SI-induced PCD, but suggests that an optimal threshold level of caspase activation is already achieved by SI-induced actin depolymerization.

In summary, our data provide good evidence that SI targets the microtubule cytoskeleton and implicates signal integration between microfilament and microtubule cytoskeleton. They reveal that SI-induced microtubule disruption is very different from that of actin. Altering microtubule dynamics did not stimulate F-actin depolymerization, suggesting one-way signaling from actin to microtubules. While actin microfilament depolymerization is sufficient to trigger PCD in pollen tubes via activation of a caspase-3-like/DEVDase activity, microtubule depolymerization alone is not. However, stabilization of microtubules reduced SI-induced caspase-like activity, suggesting that microtubule depolymerization, although on its own is insufficient to trigger PCD, is not just a consequence of SI signaling but is required for SI-induced PCD to progress.

**DISCUSSION**

**Temporal Dynamics of the SI-Mediated Microtubule Alterations**

Here, we show that in *Papaver*, although like other angiosperm pollen tubes, microtubules do not play an obvious role in regulating pollen tube growth rate (Heslop-Harrison et al., 1988; Raudaskoski et al., 2001), they are clearly responding to SI signals. Moreover, as our data demonstrate that the cortical microtubule cytoskeleton is a very early target for SI signals, it suggests that these alterations are not just a consequence of events but are likely to play a role in mediating SI. SI induces very rapid alterations to the cortical microtubule cytoskeleton, which are apparently depolymerized within approximately 1 min. Although both microtubule and microfilament SI-induced responses are very rapid, they are quite distinct responses. In contrast to F-actin, which also depolymerizes very rapidly, the microtubules remain depolymerized, while F-actin reorganizes and aggregates into punctate foci later.

One problem with fixation and such rapid responses is that it is difficult to establish exactly how rapid these changes to the cytoskeleton are and how they interrelate. Our data, and those of Gossot and Geitmann (2007) using LatB to artificially trigger actin depolymerization, show consequent apparent microtubule depolymerization, suggesting that SI-induced actin depolymerization triggers microtubule depolymerization. As stabilizing actin using Jasp prevents complete microtubule depolymerization, this further suggests a causal link. However, because of their rapidity, it is difficult to ascertain the order of these events definitively using fixation and immunolocalization. Live-cell imaging of microtubule- and microfilament-localized GFP fusion proteins would help establish the timing and nature of cytoskeletal organization and dynamics.
This would aid elucidation of the relationship between the actin and microtubule networks, especially during these early, rapid responses; we will address this in future studies.

Signal-Mediated Cortical Microtubule Reorganization/Depolymerization

Because cortical microtubules are intimately associated with the plasma membrane, where numerous receptors reside, they are implicated as targets of signaling networks (Gilroy and Trewavas, 2001; Wasteneys and Galway, 2003). Our data contribute to the evidence for this, demonstrating that the *Papaver* pollen tube microtubules are an early target of the SI-signaling network. Here, we have shown that a specific recombinant protein stimulus, involved in a biologically relevant phenomenon, has a very distinctive effect on pollen tube microtubules. The SI-induced apparent microtubule depolymerization response is extremely rapid and dramatic, far more so than any physiological response previously reported in a plant cell, to our knowledge.

Microtubule reorganization and/or apparent depolymerization occurs in response to specific abiotic stimuli (Bartolo and Carter, 1991; Himmelspach et al., 1999; Shoji et al., 2006). Several examples of the microtubule cytoskeleton alterations in response to biotic stimuli, such as infection by pathogenic fungi or symbiotic interactions with mycorrhiza or rhizobia, exist (for review, see Takemoto and Hardham, 2004). These interactions generally involve reorganization and/or that the GC-associated microtubule population are specifically targeted to the cortical microtubules (for review, see Takemoto and Hardham, 2004). Our data provide evidence for signaling to the microtubule cytoskeleton from another physiologically relevant system.

Microtubule Depolymerization Plays a Functional Role in SI-Mediated PCD

We previously showed that stabilizing F-actin using Jasp partially alleviates SI-induced PCD (Thomas et al., 2006) to about the same extent as taxol in this study. Although we did not know it at the time, actin depolymerization also stimulates microtubule depolymerization. Thus, our finding that stabilization of actin by Jasp also partially stabilizes microtubules implicates a role for microtubule depolymerization in mediating PCD. We provide a simple model outlining our understanding of the cytoskeletal events triggered by SI in order to clarify the relationship between microfilaments and microtubules (Fig. 8). Although microtubules are rapidly depolymerized by SI induction, microtubule depolymerization alone does not trigger PCD in pollen tubes. This is in contrast to actin depolymerization, which plays a key role in initiating PCD in pollen (Thomas et al., 2006). Despite this, stabilization of microtubules using taxol alleviates SI-induced PCD, suggesting that microtubules play a role in mediating PCD. Microtubule depolymerization, which we and others (Gossot and Geitmann, 2007) have shown occurs as a consequence of actin depolymerization, is effectively reduced by taxol. As we show here that taxol does not inhibit pollen tube growth, SI-induced actin depolymerization should progress normally in the presence of taxol. Thus, normal levels of SI-induced caspase induction should be triggered in the presence of taxol if microtubules play no role and are depolymerized merely as a consequence of SI-induced actin depolymerization. However, taxol alleviates PCD, this clearly demonstrates that preventing microtubule depolymerization is important for progression of PCD (Fig. 8). This strongly suggests that the microtubules are not just onlookers, but that they play a role in mediating caspase activation.

Microtubule reorganization triggered by pathogen infection hints at a possible microtubule involvement in PCD in plant cells. Our data are consistent with a model whereby microtubules, in concert with actin, somehow play a functional role in integrating signals involved in regulating PCD. However, a direct connection between microtubule reorganization and triggering of PCD remains to be elucidated.

Notably, the GC spindle-shaped microtubules were not dramatically affected by SI and remained relatively intact for a considerable time; these microtubules showed signs of disintegration but were still apparent at 60 min post-SI. This suggests that either the SI signals are specifically targeted to the cortical microtubules and/or that the GC-associated microtubule population is protected. Thus, it is the cortical microtubule population that is primarily affected and participates in this response. Interestingly, the GC appears to be a target for caspase-3-like/DEVase activity 2 to 3 h after SI induction (Bosch and Franklin-Tong, 2007).

Evidence for Cross Talk between Actin and Tubulin

It is evident from our data that there is cross talk between microfilaments and microtubules in pollen tubes during SI. We have shown that SI triggers both actin depolymerization (Snowman et al., 2002) and apparent microtubule depolymerization. Moreover, depolymerizing actin with LatB triggers microtubule depolymerization, while depolymerizing microtubules with oryzalin has no effect on actin organization, as also previously shown by Gossot and Geitmann (2007). This suggests the actin depolymerization triggers microtu-
bule depolymerization, but not vice versa, providing evidence for one-way signaling between these two cytoskeletal components in pollen tubes. As actin stabilization by Jasp delays or prevents microtubule depolymerization, this further suggests that actin influences microtubule polymerization status (Fig. 8).

Microtubules and actin microfilaments are often closely associated; in animal and yeast cells, there is no question that actin microfilament and microtubule cytoskeletons interact, and there is substantial evidence that this is also the case in plant cells. For example, transverse cortical microtubules and microfilaments in diffusely elongating cells can influence each other’s organization (Collings and Allen, 2000). Drug-induced microtubule disassembly in Characean internodal cells (Foissner and Wasteneys, 2000) and root hairs (Tominaga et al., 1997) exacerbate the effects of actin-targeted drugs, suggesting that microtubule dynamics can influence actin dynamics. In fern cells (Kadota and Wada, 1992; Collings et al., 2006) and pollen tubes (Gossot and Geitmann, 2007), actin-depolymerizing drugs affect cortical microtubules. Thus, there is good evidence for signaling and interplay between microtubules and microfilaments, but the direction of the signaling varies. In SI, both actin depolymerization (Thomas et al., 2006) and microtubule depolymerization play a role in PCD, providing evidence for an integrated signaling network between these components.

Emerging data are beginning to provide some clues about how interactions between actin and tubulin are achieved. Identification of proteins bridging these interactions has confirmed functional interactions between microtubules and microfilaments in animals and fungi (for review, see Goode et al., 2000). In plants, proteins that interact with both microtubules and actin microfilaments are beginning to be identified (Igarashi et al., 2000; Preuss et al., 2004; Huang et al., 2007), providing the first firm evidence for how these two dynamic cytoskeletal components are linked in plant cells. There is clearly much remaining to be explored in the future, and the SI-induced responses reported here appear to represent an excellent model system in which to examine interactions between microtubules and microfilaments.

**MATERIALS AND METHODS**

**Pollen Treatments**

Pollen of *Papaver rhoes* was germinated and grown in vitro in liquid germination medium [0.01% H$_3$BO$_3$, 0.01% KNO$_3$, 0.01% Mg(NO$_3$)$_2$.6H$_2$O, 0.036% CaCl$_2$, 2H$_2$O, and 13.5% Suc] as described previously (Snowman et al., 2002) at 25°C. Pollen was grown for 1 h before any treatments were applied. For SI treatments, recombinant proteins were produced by cloning the nucleotide sequences specifying the mature peptide of the S$_1$, S$_2$, and S$_3$ alleles of the S gene (pPRS100, pPRS300, and pPRS800) into the expression vector pMS119 as described previously (Foote et al., 1994). Expression and purification of the proteins was performed as described by Kakeda et al. (1998). SI was induced by adding recombinant S proteins (final concentration 10 μg ml$^{-1}$) to pollen that had been grown for 1 h in vitro (Snowman et al., 2002).

For the cytoskeleton drug treatments, 1 μM LatB, 0.5 μM Jasp (Calbiochem), 5 or 10 μM taxol, or 10 μM oryzalin (Sigma-Aldrich) was added to pollen tubes grown for 1 h. Controls comprised addition of dimethyl sulfoxide at a final concentration of 0.1% (v/v). For the drug-SI experiments, pollen tubes were subjected to a consecutive treatment of the relevant drug for 30 min, followed by the addition of incompatible S proteins for 5 h.

**Immunolocalization**

Pollen tubes were prefixed using the cross-linker 3-maleimidobenzoic acid N'-hydroxysuccinimide ester (MBS; 400 μl; Pierce) for 6 min at 20°C, followed by 2% formaldehyde (1 h, 4°C), as described by Thomas et al. (2006); we used 2% formaldehyde as a compromise. Actin preservation was indistinguishable from what we previously obtained using 4% formaldehyde following MBS (Geitmann et al., 2000). MBS has been reported to stop cytoplasmic streaming within seconds (Ketelaar and Emons, 2001). The treatment times indicated in the text are the time point after treatment that MBS was added. Cells were washed in actin-stabilizing buffer (100 mM PIPES, pH 6.8, 1 mM MgCl$_2$, 1 mM CaCl$_2$, 75 mM KCl) then in MES buffer (15 mM MES, pH 5.0), then incubated in 0.05% cellulose, 0.05% macerozyme in MES buffer containing 0.1 mM phenylmethylsulfonyl fluoride and 1% bovine serum albumin for 10 min. Washes in MES, then Triton-buffered saline (TBS), were followed by permeabilization in

![Figure 8. Model for integration of cytoskeletal events triggered by SI. SI triggers actin depolymerization, which is sufficient to trigger caspase activation and PCD (Thomas et al., 2006). LatB causes actin depolymerization, caspase activation, and PCD (Thomas et al., 2006). Treatment with Jasp after SI induction alleviated the extent of PCD (Thomas et al., 2006). Thus, partial prevention of actin depolymerization gives some protection from PCD. SI also triggers microtubule depolymerization (this study). Use of LatB showed that actin depolymerization also triggers microtubule depolymerization (Gossot and Geitmann, 2007; this study). This suggests that during SI, microtubule depolymerization is a consequence of actin depolymerization. Use of oryzalin showed that microtubule depolymerization on its own is not sufficient to trigger caspase activation and PCD (this study). This raises the question of whether microtubule depolymerization is actually required for PCD, or whether it is just a consequence of actin depolymerization. Use of taxol, which alleviated the extent of PCD, showed that preventing microtubule depolymerization is somehow involved in regulating PCD (this study). This implicates a functional role for both actin and tubulin in signaling to PCD.](image-url)
were extracted by sonication (23 mouse alkaline phosphatase secondary antibody and detected using alkaline phospho-todilution overnight at 4°C. They were washed in TBS, then incubated for 1.5 h at room temperature in anti-mouse fluorescein isothiocyanate antibody (1:300 dilution). Following TBS washes, rhodamine-phalloidin (66 nm) was added. Pollen tubes were mounted with 5 μL of Vectashield (Vector Laboratories). Images were collected using a Bio-Rad Radiance 2000 laser-scanning system (50-W argon laser: 488-nm line; and 1.5-mW HeNe laser, 543 nm) with a 60× plan-Neofluar 1.4 NA oil objective (Nikon). z-Series of 0.5-μm optical slices were captured. Images were analyzed using ImageJ and archived as TIF files.

**Protein Extraction and Western Blotting**

SI was induced and pollen tubes collected by centrifugation in HEPES buffer (50 mM HEPES, pH 7.4, 10 mM NaCl, 0.1% CHAPS, 10 mM dithiothreitol, 1× EDTA, 10% glycerol) and samples snap-frozen in liquid N2. Proteins were extracted by sonication (2 × 10 s, 10 amps) and analyzed using SDS-PAGE and western blotting. Samples were measured using the Bio-Rad protein assay; equal amounts were loaded and checked by Phosphoimaging of gels. Blots were probed with a 1:4,000 dilution of the monoclonal anti-α-tubulin antibody clone B-5-1-2 (Sigma-Aldrich), then probed with an anti-mouse alkaline phosphatase secondary antibody and detected using alkaline phosphatase.

**Pollen Tube Length Measurements**

Pollen tubes were grown for 1 h, then samples were treated as specified in the text, and pollen tubes fixed in 2% formaldehyde for 1 h, washed in TBS, and mounted on glass slides. Thus, before treatment, all mean pollen tube lengths were similar. Fixed pollen tubes were imaged using a Nikon Eclipse TE-300 microscope attached to a SenSys camera, using a Quips PathVision image analysis system (Applied Imaging International). Final pollen tube lengths were measured (40 tubes for each of three independent treatments) using IPLab software. Lengths indicated are total lengths of the pollen tubes (i.e. 1 h pretreatment time plus treatment time with the relevant drug). Statistical analysis comprised a 1 test analysis.

**Caspase Assays**

PCD was assessed using a fluorogenic caspase-3/7-amino-4-trifluoromethyl coumarin substrate, Ac-DEVD-AMC, to measure caspase-like activity. Assays containing 10 μg of protein extract at 1 μg/mL-1 and 50 μM substrate were performed in caspase extraction buffer, pH 5.0. Release of fluorophore by cleavage was measured (excitation 380 nm, emission 460 nm) using a FLUOstar OPTIMA reader (BMG Labtechnologies) at 27°C for 5 h. Background relative fluorescent unit readings for control samples were subtracted from test samples. All assays were performed on at least four independent samples, each measured in duplicate. P values were calculated using a two-way ANOVA.

**ACKNOWLEDGMENTS**

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**LITERATURE CITED**


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0.1% Triton X-100/TBS for 10 min, and blocking in TBS/1% bovine serum albumin for 30 min.

Samples were incubated with anti-α-tubulin antibody (clone B-5-1-2; Sigma-Aldrich; 1:1,000 dilution) overnight at 4°C. They were washed in TBS, then incubated for 1.5 h at room temperature in anti-mouse fluorescein isothiocyanate antibody (1:300 dilution). Following TBS washes, rhodamine-phalloidin (66 nm) was added. Pollen tubes were mounted with 5 μL of Vectashield (Vector Laboratories). Images were collected using a Bio-Rad Radiance 2000 laser-scanning system (50-W argon laser: 488-nm line; and 1.5-mW HeNe laser, 543 nm) with a 60× plan-Neofluar 1.4 NA oil objective (Nikon). z-Series of 0.5-μm optical slices were captured. Images were analyzed using ImageJ and archived as TIF files.
Self-Incompatibility Signals to Microtubules


**My contribution:** I analysed the predicted transmembrane structure of the PrpS protein using various prediction programmes (supplementary Figure 2) and demonstrated, using western ligand-blotting, that a predicted extracellular domain interacts in an S-specific manner with the stigmatic PrsS protein (Figure 2e). I proofread the manuscript.
Identification of the pollen self-incompatibility determinant in *Papaver rhoeas*

Michael J. Wheeler¹, Barend H. J. de Graaf¹, Natalie Hadjiosif¹, Ruth M. Perry¹, Natalie S. Poulter¹, Kim Osman¹, Sabina Vatovec¹, Andrea Harper¹, F. Christopher H. Franklin¹ & Vernonica E. Franklin-Tong¹

Higher plants produce seed through pollination, using specific interactions between pollen and pistil. Self-incompatibility is an important mechanism used in many species to prevent inbreeding; it is controlled by a multi-allelic S locus². ‘Self’ (incompatible) pollen is discriminated from ‘non-self’ (compatible) pollen by interaction of pollen and pistil S locus components, and is subsequently inhibited. In *Papaver rhoeas*, the pistil S locus product is a the pollen component of the S locus on a cosmid clone comprising a 42-kilobase (kb) region at the S₁ locus.

Nucleotide sequencing and analysis identified a novel putative open reading frame (ORF) 457 base pairs (bp) from the S₁ pistil gene (Fig. 1a). Expression analysis using polymerase chain reaction with reverse transcription (RT–PCR) revealed that the ORF was specifically transcribed in pollen (Fig. 1b), appearing during anther

**My contribution:** This paper was a review. Noni Franklin-Tong, Maurice Bosch and Natalie Poulter mainly wrote the manuscript. I proof-read the manuscript.
Initiation of Programmed Cell Death in Self-Incompatibility: Role for Cytoskeleton Modifications and Several Caspase-Like Activities

Maurice Bosch\textsuperscript{a,b}, Natalie S. Poulter\textsuperscript{a}, Sabina Vatovec\textsuperscript{a} and Vernonica E. Franklin-Tong\textsuperscript{a,1}

\textsuperscript{a} School of Biosciences, University of Birmingham, Edgbaston, Birmingham. B15 2TT, UK
\textsuperscript{b} Present address: Institute of Biological, Environmental and Rural Sciences (IBERS), Aberystwyth University, Plas Gogerddan, Aberystwyth SY23 3EB, UK

ABSTRACT Programmed cell death (PCD) is an important and universal process regulating precise death of unwanted cells in eukaryotes. In plants, the existence of PCD has been firmly established for about a decade, and many components shown to be involved in apoptosis/PCD in mammalian systems are found in plant cells undergoing PCD. Here, we review work from our lab demonstrating the involvement of PCD in the self-incompatibility response in \textit{Papaver rhoeas} pollen. This utilization of PCD as a consequence of a specific pollen–pistil interaction provides a very neat way to destroy unwanted ‘self’, but not ‘non-self’ pollen. We discuss recent data providing evidence for SI-induced activation of several caspase-like activities and suggest that an acidification of the cytosol may be a key turning point in the activation of caspase-like proteases executing PCD. We also review data showing the involvement of the actin and microtubule cytoskeletons as well as that of a MAPK in signalling to caspase-mediated PCD. Potential links between these various components in signalling to PCD are discussed. Together, this begins to build a picture of PCD in a single cell system, triggered by a receptor–ligand interaction.

My contribution: This paper was a review containing some original data. I contributed original data demonstrating S-specific PrpS-PrsS binding using western ligand blotting (Figure 3). I also organized data into Figure 3 and wrote the figure legend. I did the proof-reading of the manuscript.
The pollen S-determinant in Papaver: comparisons with known plant receptors and protein ligand partners

Michael J. Wheeler†, Sabina Vatovec and Veronica E. Franklin-Tong*

School of Biosciences, University of Birmingham, Edgbaston, Birmingham B15 2TT, UK
† Present address: Warwick HRI, Wellesbourne, Warwick CV35 9EF, UK.
* To whom correspondence should be addressed: E-mail: v.e.franklin-tong@bham.ac.uk

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Abstract

Cell–cell communication is vital to multicellular organisms and much of it is controlled by the interactions of secreted protein ligands (or other molecules) with cell surface receptors. In plants, receptor–ligand interactions are known to control phenomena as diverse as floral abscission, shoot apical meristem maintenance, wound response, and self-incompatibility (SI). SI, in which ‘self’ (incompatible) pollen is rejected, is a classic cell–cell recognition system. Genetic control of SI is maintained by an S-locus, in which male (pollen) and female (pistil) S-determinants are encoded. In Papaver rhoeas, PrsS proteins encoded by the pistil S-determinant interact with incompatible pollen to effect inhibition of pollen growth via a Ca$^{2+}$-dependent signalling network, resulting in programmed cell death of ‘self’ pollen. Recent studies are described here that identified and characterized the pollen S-determinant of SI in P. rhoeas. Cloning of three alleles of a highly polymorphic pollen-expressed gene, PrpS, which is linked to pistil-expressed PrsS revealed that PrpS encodes a novel ~20 kDa transmembrane protein. Use of antisense oligodeoxynucleotides provided data showing that PrpS functions in SI and is the pollen S-determinant. Identification of PrpS represents a milestone in the SI field. The nature of PrpS suggests that it belongs to a novel class of ‘receptor’ proteins. This opens up new questions about plant ‘receptor’–ligand pairs, and PrpS-PrsS have been examined in the light of what is known about other receptors and their protein–ligand pairs in plants.

Key words: Cell–cell recognition, Papaver rhoeas, pollen S-determinant, pollen tube inhibition, PrpS, receptor, self-incompatibility, self-recognition.

The *Papaver* self-incompatibility pollen S-determinant, PrpS, functions in *Arabidopsis thaliana*.

* - these authors contributed equally to this work

**My contribution:**

Barend de Graaf initiated the project, contributed the original idea for the manuscript, produced *A.thaliana* transgenic line expressing PrpS1-GFP, carried out the experiments presented on Figure 1 and wrote the manuscript. Javier Juarez-Diaz helped me with the RT-PCR experiments (Figure 1 e), helped with preparation of samples for actin confocal imaging (Figure 3 a-j) and helped with the discussions for the manuscript. Huawen Zou produced *A.thaliana* transgenic line expressing PrpS3-GFP. Katie Wilkins carried out imaging of pollen on the confocal microscope (Figure 3 a-j) and Lijun Chai prepared samples and analysed the actin cytoskeleton of the PrpS3 expressing pollen (Figure 3 l). Kreepa Kooblall and Tom Forbes were the project students under my supervision. Kreepa Kooblall started the initial pollen tube inhibition studies on transgenic *A.thaliana* pollen expressing PrpS1-GFP, while Tom Forbes started the initial actin cytoskeleton analysis in transgenic *A.thaliana* expressing PrpS1-GFP. Chris Franklin contributed with counsel and discussions during the project. Noni Franklin-Tong managed the project, provided the guidance, wrote the paper and is the corresponding author.

I carried out the RT-PCR and segregation analysis of the transgenic *A.thaliana* lines (Figure 1 e), I carried out the experiment and statistical analysis of pollen tube inhibition of PrpS1 and PrpS3 expressing pollen (Figure 2) and pollen viability of PrpS1 and PrpS3 expressing
pollen (Figure 4 a-e). I produced samples for the confocal imaging of the actin cytoskeleton (Figure 3 a-j) and carried out the epifluorescence imaging, analysis and statistics of the actin cytoskeleton of the PrpS1 expressing pollen (Figure 3 k). I proof-read the manuscript.
The *Papaver* Self-Incompatibility Pollen S-Determinant, *PrpS*, Functions in *Arabidopsis thaliana*

Barend H.J. de Graaf,1,2,5 Sabina Vatovec,1,5 Javier Andrés Juárez-Díaz,1,5 Lijun Chai,1 Kreepa Kooblall,1,3 Katie A. Wilkins,1 Huawen Zou,1,4 Thomas Forbes,1 F. Christopher H. Franklin,1,6 and Vernonica E. Franklin-Tong1,6,*

1School of Biosciences, University of Birmingham, Edgbaston, Birmingham B15 2TT, UK

Summary

Many angiosperms use specific interactions between pollen and pistil proteins as “self” recognition and/or rejection mechanisms to prevent self-fertilization. Self-incompatibility (SI) is encoded by a multiallelic *S* locus, comprising pollen and pistil *S*-determinants [1, 2]. In *Papaver rhoeas*, cognate pistil and pollen *S*-determinants, PrpS, a pollen-expressed transmembrane protein, and PrsS, a pistil-expressed secreted protein [3, 4], interact to trigger a Ca$^{2+}$-dependent signaling network [5–10], resulting in inhibition of pollen tube growth, cytoskeletal alterations [11–13], and programmed cell death (PCD) [14, 15] in incompatible pollen. We introduced the *PrpS* gene into *Arabidopsis thaliana*, a self-compatible model plant. Exposing transgenic *A. thaliana* pollen to recombinant *Papaver* PrsS protein triggered remarkably similar responses to those observed in incompatible *Papaver* pollen: *S*-specific inhibition and hallmark features of *Papaver* SI [11–15]. Our findings demonstrate that *Papaver* PrpS is functional in a species with no SI system that diverged ~140 million years ago [16]. This suggests that the *Papaver* SI system uses cellular targets that are, perhaps, common to all eudicots and that endogenous signaling components can be recruited to elicit a response that most likely never operated in this species. This will be of interest to biologists interested in the evolution of signaling networks in higher plants.

Expression of Prps-GFP Is Sufficient to Induce S-Specific Inhibition of *AtPpS3*

To determine whether PrpS was functionally adapted the in vitro self-incompatibility system used for *Papaver* SI [3]. Transgenic *AtPpS3* was grown in vitro and recombinant proteins added. If PrpS functions in a self-incompatible model plant. Exposing transgenic *A. thaliana* pollen to recombinant *Papaver* PrsS protein triggered remarkably similar responses to those observed in incompatible *Papaver* pollen: *S*-specific inhibition and hallmark features of *Papaver* SI [11–15]. Our findings demonstrate that *Papaver* PrpS is functional in a species with no SI system that diverged ~140 million years ago [16]. This suggests that the *Papaver* SI system uses cellular targets that are, perhaps, common to all eudicots and that endogenous signaling components can be recruited to elicit a response that most likely never operated in this species. This will be of interest to biologists interested in the evolution of signaling networks in higher plants.