Investigation of upstream and downstream events in CERK1-mediated signaling in plant immunity

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<tbody>
<tr>
<td>amiRNA</td>
<td>Artificial micro RNA</td>
</tr>
<tr>
<td>At</td>
<td>Arabidopsis thaliana</td>
</tr>
<tr>
<td>AtDORN1</td>
<td>Arabidopsis does not respond to nucleotides 1</td>
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<td>AtWAK1</td>
<td>Arabidopsis wall-associated kinase 1</td>
</tr>
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<td>Avr gene</td>
<td>Avirulence gene</td>
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<td>BRI1-associated kinase 1</td>
</tr>
<tr>
<td>BRI1</td>
<td>Brassinosteroid insensitive 1</td>
</tr>
<tr>
<td>BiFC</td>
<td>Bimolecular fluorescence complementation</td>
</tr>
<tr>
<td>cDNA</td>
<td>Complementary DNA</td>
</tr>
<tr>
<td>C(D)PK</td>
<td>Calcium-dependent protein kinase</td>
</tr>
<tr>
<td>CfU</td>
<td>Colony forming unit</td>
</tr>
<tr>
<td>DAMP</td>
<td>Damage-associated molecular pattern</td>
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<td>DNA</td>
<td>Deoxyribonucleic acid</td>
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<tr>
<td>ECD</td>
<td>Extracellular domain</td>
</tr>
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<td>EF1a</td>
<td>Elongation factor 1a</td>
</tr>
<tr>
<td>EFR</td>
<td>Elongation factor-Tu receptor</td>
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<tr>
<td>EGF</td>
<td>Epidermal growth factor</td>
</tr>
<tr>
<td>EF-TU</td>
<td>Elongation factor-Tu</td>
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<tr>
<td>ETI</td>
<td>Effector triggered immunity</td>
</tr>
<tr>
<td>ETS</td>
<td>Effector-triggered susceptibility</td>
</tr>
<tr>
<td>FLS2</td>
<td>Flagellin-sensing 2</td>
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<tr>
<td>FPLC</td>
<td>Fast protein liquid chromatography</td>
</tr>
<tr>
<td>FRK1</td>
<td>Flagellin responsive kinase 1</td>
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<tr>
<td>GFP</td>
<td>Green fluorescent protein</td>
</tr>
<tr>
<td>GlcNAc</td>
<td>N-acetylglucosamine</td>
</tr>
<tr>
<td>GNBP1</td>
<td>Gram-negative bacteria-derived binding protein 1</td>
</tr>
<tr>
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<td>Description</td>
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</tr>
<tr>
<td>HPLC</td>
<td>High performance liquid chromatography</td>
</tr>
<tr>
<td>HR</td>
<td>Hypersensitive response</td>
</tr>
<tr>
<td>HRP</td>
<td>Horseradish peroxidase</td>
</tr>
<tr>
<td>KD</td>
<td>Kinase domain</td>
</tr>
<tr>
<td>kDa</td>
<td>KiloDalton</td>
</tr>
<tr>
<td>LORE</td>
<td>Lipooligosaccharide-specific reduced elicitation</td>
</tr>
<tr>
<td>LPS</td>
<td>Lipooligosaccharide</td>
</tr>
<tr>
<td>LRR</td>
<td>Leucine rich repeat</td>
</tr>
<tr>
<td>LYS1</td>
<td>Lysozyme-like enzyme 1</td>
</tr>
<tr>
<td>LysM</td>
<td>Lysin motif</td>
</tr>
<tr>
<td>MAPK</td>
<td>Mitogen-activated protein kinase</td>
</tr>
<tr>
<td>MbSUS</td>
<td>Mating-based split ubiquitin system</td>
</tr>
<tr>
<td>MDP</td>
<td>Muramyl dipeptide</td>
</tr>
<tr>
<td>MIMP</td>
<td>Microbe-associated molecular patterns</td>
</tr>
<tr>
<td>MIND</td>
<td>Membrane-based interactome network database</td>
</tr>
<tr>
<td>MurNAc</td>
<td>N-Acetylmuramic acid</td>
</tr>
<tr>
<td>NASC</td>
<td>Nottingham Arabidopsis stock center</td>
</tr>
<tr>
<td>NOD</td>
<td>Nucleotide-binding oligomerization domain-containing protein</td>
</tr>
<tr>
<td>PAMP</td>
<td>Pathogen-associated molecular pattern</td>
</tr>
<tr>
<td>Peptidoglycan</td>
<td>PGN</td>
</tr>
<tr>
<td>PGRP</td>
<td>PGN-recognition protein</td>
</tr>
<tr>
<td>PR protein</td>
<td>Pathogenesis-related protein</td>
</tr>
<tr>
<td>PRR</td>
<td>Pattern recognition receptor</td>
</tr>
<tr>
<td>PTI</td>
<td>PAMP-triggered immunity</td>
</tr>
<tr>
<td>Pto</td>
<td><em>Pseudomonas syringae</em> pv. <em>tomato</em></td>
</tr>
<tr>
<td>R gene</td>
<td>Resistance gene</td>
</tr>
<tr>
<td>RFP</td>
<td>Red fluorescent protein</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
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<td>--------------</td>
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</tr>
<tr>
<td>RLK</td>
<td>Receptor-like kinases</td>
</tr>
<tr>
<td>RLP</td>
<td>Receptor-like protein</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive oxygen species</td>
</tr>
<tr>
<td>RT-qPCR</td>
<td>Reverse transcription quantitative PCR</td>
</tr>
<tr>
<td>SAR</td>
<td>Systemic acquired resistance</td>
</tr>
<tr>
<td>SOBIR1</td>
<td>Suppressor of bir1</td>
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<tr>
<td>T-DNA</td>
<td>Transfer DNA</td>
</tr>
<tr>
<td>TLR2</td>
<td>Toll-like receptor 2</td>
</tr>
<tr>
<td>WT</td>
<td>Wild type</td>
</tr>
<tr>
<td>YFP</td>
<td>Yellow fluorescent protein</td>
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1. Introduction

1.1 Origin and evolution of plant immunity concepts

1.1.1 Plant and microbe interactions

The advent of the earliest land plants around 480 million years ago is one of the most important evolutionary events in the history of life on Earth (Gehrig et al., 1996). During their establishment in terrestrial ecosystems, land plants have had to adapt to an environment that houses a large variety of microbes such as fungi, oomycetes, bacteria, and viruses. From then on, if not earlier, plants and microbes have had continual interaction, and this interplay has been influencing the evolution of both plants and microbes (Wang et al., 2010).

The interplay of plants and microbes has led to a wide range of relationships in which one or both of the organisms may have a beneficial, neutral or negative effect on the other partner. These diverse plant-microbe interactions have been broadly classified into three categories from the plant perspective: beneficial, detrimental or neutral interaction (Schenk et al., 2012).

Most plant-microbe interactions have neutral effects on plants. These “neutral” microbes often utilize plant derived organic compounds as energy sources, and thus play important roles in nutrient cycling and modification of environments. Many microbes are beneficial for plants and promote growth or suppress diseases via a number of mechanisms, including improved nutrient acquisition, production of growth regulators, and biosynthesis of pathogen-inhibiting compounds (Schenk et al., 2012). Microbes that are harmful to plants are much fewer, although more noticeable, such as plant pathogenic fungi, oomycetes, bacteria and viruses that can potentially cause diseases in plants. The outcome of these plant–microbe interactions are further influenced by wounding, herbivores and abiotic stress factors such as salinity, drought, temperature (Thaler et al., 1999; Cheong et al., 2002; Berg, 2009).

1.1.2 Immunity in animals and plants

To respond appropriately to diverse microbes, plants have evolved abilities to distinguish self from non-self and friends from enemies. With this distinction, plants can adapt to
their environment by either activating immune responses to defend against pathogens or initiating symbiosis signaling to facilitate the accommodation of symbionts.

The last common ancestor of plants and animals is thought to have lived approximately one billion years ago, which may explain dramatic differences in structures and lifestyles between plants and animals. The differences made biologists initially believe that plants and animals defend against microbial pathogens in separate and distinct ways (Ronald and Beutler, 2010). For example, vertebrates have developed two types of immunity, innate and adaptive immunity, to defend against microbial invaders (Medzhitov and Janeway, 2002; McGuinness et al., 2003). Adaptive immunity in vertebrate animals is mediated by a clonal system of T lymphocytes and B lymphocytes, and is characterized by the creation of antigen-specific receptors through somatic recombination in maturing lymphocytes, and this system does not exist in plants. Moreover, specialized cell types such as macrophages, neutrophils, and dendritic cells, which as parts of a circulatory blood system are the key players of the animal immune system, are not found in plants either. In contrast, each plant cell is autonomously capable of sensing the presence of microbial infection and of mounting defence responses (Nürnberger et al., 2004).

Despite the well-documented differences, increasing evidences over the past twenty years have revealed striking structural and strategic similarities in the immune systems of plants and animals (Nürnberger and Brunner, 2002b). For instance, both are able to respond to microbial signature molecules using analogous regulatory modules. Pathogen-associated molecular patterns (PAMP) that are similar to those activating innate immune responses in animals have been shown to trigger the activation of plant defence. Moreover, recognition complexes that are structurally related to animal PAMP receptors are now being discovered in plants (Nürnberger and Brunner, 2002a).

The remarkable similarities of the innate immune systems between animal and plant raise an intriguing issue: Did an ancient ancestor common to plants and animals evolve a basic innate immune system, one that began to differ in the two lineages once they split (divergent evolution)? Or did plants and animals evolve innate immunity independently but ended up with similar mechanisms (convergent evolution)? Comparative analyses of innate immune systems in plants and animals, based on
significantly broad data sets, may provide a basis to address this issue more precisely in the future.

1.1.3. History of plant immunity

Although plant diseases were mentioned as early as 750 BCE in the Hebrew Bible, it was not until 1853 that De Bary discovered that rust and smut fungi were causal agents of certain diseases in cereal crops (Agrios, 2005). The first description of a genetic relationship between host plants and their pathogens was in the early 1940s by Harold Flor, based on genetic investigation of flax and the flax rust fungus, and the investigation lead to the “gene-for-gene hypothesis” in 1946 (Flor, 1971). The hypothesis proposed that the ability of fungal pathogens to cause disease was controlled by two matching genes: an avirulence (Avr) gene from the pathogen and a matching resistance (R) gene from the host. This hypothesis was a theoretical breakthrough in plant pathology and led to practical advances in plant breeding.

Flor’s model presumed that specific sensors for microbial molecules, termed elicitors, or Avr gene products, were present in immunocompetent hosts. In the 1980s an intense hunt for the genes encoding these receptors and their corresponding elicitors, based on biochemical approaches, identified specific binding sites for elicitors in intact plant cells and on isolated plasma membranes, but failed to purify the expected receptors (Boller, 1995).

In the 1990s, an avalanche of genetic experiments led to the isolation of the first R genes and, subsequently, more R genes from multiple plant species (Martin et al., 1993). These discoveries established that diverse molecules and mechanisms govern the resistance phenotypes described by Flor, and that the resistance response was more complex than previously realized.

A philosophical divergence between geneticists, who isolated R genes, and biochemists, who identified general elicitors, lead to two schools of thought: microbial elicitor-induced plant non cultivar-specific defence and microbial avirulence factor-induced plant cultivar-specific defence. The divergence raised a debate whether the elicitor-induced defence responses were physiologically relevant to plant immunity and what the relationship
between the gene-for-gene resistance and the elicitor-induced defence would be. Since the first elicitor receptor (Gomez-Gomez and Boller, 2000) was identified, the debate has given way to the idea that two types of defence both contribute to plant immunity in an integrated fashion. A simple and elegant view of innate immunity in plant pathogen interactions is depicted by the so-called zigzag model, introduced by Jones and Dangl, providing an integrated model for plant disease resistance (Jones and Dangl, 2006).

1.2 The plant immune system

Plants, unlike animals, are sessile organisms and cannot escape from environmental threats such as biotic and abiotic stresses. Instead, plants have evolved a wide range of mechanisms to cope with these biotic and abiotic stresses for survival. Biotic stresses result from surrounding living organisms such as microbes, parasites, herbivores and other plants. Of these biotic stress factors, phytopathogenic microbes frequently and seriously threat food security and thus are overwhelmingly focused on by scientists.

1.2.1 The three-layer-defence in plants

Resistances of plants to phytopathogens have been described as three broad layers relevant to pathogens’ foliar infection strategies and plant counter-defensive responses (Senthil-Kumar and Mysore, 2013). Upon landing on the surface of a plant, a potential microbial pathogen initially encounters the first plant defence layer that restricts pathogen entry. This step usually involves preformed defence, such as a wax layers, rigid cell walls, cuticular lipids (Reina-Pinto and Yephremov, 2009), antimicrobial enzymes or secondary metabolites (Ahuja et al., 2012; Bednarek, 2012), as well as inducible defence, such as stomatal closure (Melotto et al., 2006). Pathogens that overcome this surface defence layer encounter a second layer of defence in the apoplastic region. This step also involves preformed defence, such as the presence of antimicrobial compounds, coupled with overall apoplastic physiological incompatibility and induced defence, such as phytoalexin production (Dixon et al., 2002; Ham et al., 2007; Hann and Rathjen, 2007; Huckelhoven, 2007). The pathogens that successfully break through the first two layers of defence must finally face a third layer of defence at and inside the plant cell. This defence layer involves two inducible interconnected
Mechanisms: pathogen-associated molecular pattern (PAMP)-triggered immunity (PTI) and effector-triggered immunity (ETI). Both PTI and ETI are induced by pathogen-generated stimuli and regulate the protective responses of plants against pathogens in synergistic and antagonistic ways (Jones and Dangl, 2006).

1.2.2 Plant PTI: PAMPs and pattern-recognition receptors

Activation of plant PTI depends on the perception of invariant structures of mostly pathogen surfaces, formerly known as “general elicitors” (Boller, 1995). These invariant pathogen structures are named as pathogen-associated molecular patterns (PAMPs) (Medzhitov and Janeway, 1997). PAMPs are highly conserved molecular patterns across a wide range of pathogens but are not present in the host and thus enable hosts to distinguish self and non-self (Medzhitov and Janeway, 2002; Akira et al., 2006). PAMPs include (poly)peptides, glycoproteins, lipids, and oligosaccharides, which are essential for microbial fitness and survival (Nürnberger et al., 2004). Classical PAMPs are exemplified by eubacterial flagellin, elongation factor-Tu (EF-Tu), peptidoglycans (PGNs), oomycete glucans, and fungal chitin (Sharp et al., 1984; Cheong et al., 1991; Felix et al., 1993; Felix et al., 1999; Zipfel et al., 2004; Zipfel et al., 2006; Gust et al., 2007). As PAMP molecules are found not only in pathogenic microbes but also non-pathogenic microorganisms (Ausubel, 2005), in the recent literature more correct terms have therefore been introduced, such as MAMPs (microbe-associated molecular patterns) (Ausubel, 2005) or MIMPs (microbe-induced molecular patterns) (Mackey and McFall, 2006). While these terms have their merits and justification, using the ‘historical’ term PAMPs allows for better understanding among researchers in animal and plant immunity (Nürnberger and Kemmerling, 2009), and therefore in this thesis I will follow this suggestion.

In addition to microbial PAMPs, the structural barriers of plant tissues can be degraded by the lytic enzymes of plant pathogens, releasing products such as plant cell wall fragments (Darvill and Albersheim, 1984), cutin monomers (Kauss et al., 1999) and peptides (Boller, 2005; Huffaker et al., 2006), all of which can function as endogenous elicitors called damage-associated molecular patterns (DAMPs). These DAMPs
characteristically emerge in the apoplast and serve as danger signals to induce innate immunity similar to PAMPs (Henry et al., 2012).

Plant PTI relies on the perception of PAMPs by plant cell surface-localized pattern recognition receptors (PRRs). All known plant PRRs are plasma membrane-localized receptor-like kinases (RLKs) or receptor-like proteins (RLPs) with modular functional domains. RLKs contain an extracellular domain (ECD), a single-pass transmembrane (TM) domain, and an intracellular kinase domain (KD), whereas RLPs include an ECD and a TM but lack the intracellular KD (Monaghan and Zipfel, 2012). Typically, small epitopes within PAMPs provide ligands for PRR receptors. Various receptor types mediate pattern recognition that differ in extracellular ligand-binding domains, membrane embedding and intracellular cytoplasmic domains (Böhm et al., 2014). Structural biochemistry has recently highlighted ligand induced immune receptor complex formation as a common principle in pattern binding and cytoplasmic signal transduction (Böhm et al., 2014).

The first major class of PRRs is leucine-rich repeat receptor kinases/proteins (LRR-RLKs/RLPs). LRR-RLKs contain a leucine-rich repeat (LRR) ectodomain, which can bind to proteinaceous immunogenic patterns (Zhang and Thomma, 2013). For example, flagellin-sensing 2 (FLS2) recognizes a 22-amino acid epitope (flg22) conserved in bacterial flagellins (Zipfel et al., 2004) and EF-Tu receptor (EFR) perceives the first 18 amino acids (elf18) of bacterial EF-Tu (Zipfel et al., 2006). These LRR-RLKs form heterodimeric complexes with the brassinosteroid-insensitive 1 (BRI1)-associated kinase 1 (BAK1), a LRR-RLK, in a ligand dependent manner. Likewise, LRR-RLP receptors that lack intracellular kinase domains also form heteromeric complexes with different LRR-RK-type co-receptors such as BAK1 or suppressor of bir1 (SOBIR1), in a ligand-dependent manner as well (Jehle et al., 2013; Zhang et al., 2013; Zhang et al., 2014).

Lysin motif (LysM)-type immune receptor, a second major class of PRRs, mediates recognition of N-acetylglucosamine (GlcNAc)-containing carbohydrate ligands. For example, bacterial PGN recognition in Arabidopsis involves two LysM-RLPs (AtLYM1/3) and one LysM-RLK (Chitin elicitor receptor-like kinase 1, CERK1) (Gust et al., 2007).
Similarly, the chitin perception in *Arabidopsis* also involves one LysM-RLK (CERK1) and one kinase-inactive LysM-RLK (LYK5) (Miya et al., 2007; Liu et al., 2012b; Cao et al., 2014) and in rice similar perception systems for PGN and chitin have been described (Kaku et al., 2006; Liu et al., 2012a; Hayafune et al., 2014).

Additional PRRs include an epidermal growth factor (EGF)-like domain-containing receptor kinase AtWAK1 (Arabidopsis wall-associated kinase 1) (Brutus et al., 2010), a DAMP receptor implicated in resistance to fungal infection, and a lectin receptor kinase I.9/AtDORN1 (Arabidopsis Does not Respond to Nucleotides 1) that binds extracellular ATP likely released during wounding, herbivory or microbial infection (Choi et al., 2014). Recently, the bulb-type lectin S-domain-1 receptor–like kinase LORE (lipooligosaccharide-specific reduced elicitation) was identified as a potential receptor to recognize the lipid A moiety of lipooligosaccharides (LPS) (Ranf et al., 2015).

With the expansion of plant post-genomic tools, many more PRRs are likely to be identified in the future.

Although it is not fully understood, accumulating evidences support the idea that plant PRRs likely appear as central components of multiprotein complexes in the plasma membrane, which links the detection of PAMPs to the activation of downstream signaling components under tight control by regulatory proteins such as phosphatases or E3 ligases (Böhm et al., 2014; Macho and Zipfel, 2014).

### 1.2.3 Plant ETI

As basal immunity, PTI is sufficient for plants to defend against non-adapted pathogens. In response, some would-be pathogens have successfully evolved the capability to avoid PTI by secreting a collection of effectors into the host apoplast or directly inside the host cell to interfere with PTI (Hogenhout et al., 2009; Bozkurt et al., 2012; Feng and Zhou, 2012). These effectors suppress PTI either upstream of PAMP recognition by targeting PRRs directly (de Jonge et al., 2010; Mentlak et al., 2012), or downstream by blocking signaling events (Zhang et al., 2007a; Göhre et al., 2008; Navarro et al., 2008), and finally result in the effector-triggered susceptibility (ETS). To counteract ETS, some plants have further evolved additional resistance (R) proteins that specifically recognize
the pathogen-secreted effectors, establishing a second layer of immunity known as the effector-triggered immunity (ETI) (Jones and Dangl, 2006).

Although the PTI-ETI nomenclature unarguably has its merits in illustrating how co-evolution of microbes and their hosts has shaped the plant immune system, the distinction between PAMPs and effectors, between PAMP receptors and resistance proteins, and therefore, also between PTI and ETI, cannot strictly be maintained. Some examples support a continuum between PTI and ETI, and therefore it has been suggested that plant immune activation is generally determined by immune receptors or PRRs that mediate pattern recognition (Thomma et al., 2011; Böhm et al., 2014).

1.2.4 Signal transduction in plant immunity

Perception of PAMPs (during PTI) or effectors (during ETI) rapidly activates a set of common defence responses, which are usually sufficient to defend against most non-adapted pathogens. These often include a rapid influx of calcium ions from external stores (Lee et al., 2001; Jeworutzki et al., 2010), a burst of active oxygen species (Apel and Hirt, 2004), activation of mitogen-activated protein kinases (MAPKs) and calcium (Ca$^{2+}$)-dependent protein kinases (CDPKs) (Tena et al., 2011), activation of WRKY transcription factors (Asai et al., 2002), reprogramming of gene expression and transcription of defence-related genes (Zipfel et al., 2004), deposition of callosic cell wall appositions at sites of attempted infection (Lazarovits and Higgins, 1976), phytoalexin accumulation (Nürnberger, 1999; Zipfel et al., 2006; Chinchilla et al., 2007) and plant hormonal responses (Denance et al., 2013). These responses also play important roles in defence against adapted pathogens (Bittel and Robatzek, 2007) and might lead to ETI-associated cell death (hypersensitive response, HR) (Beers and McDowell, 2001), often followed by the establishment of systemic acquired resistance (SAR) which is a long lasting and broad-spectrum resistance mechanism (Durrant and Dong, 2004).

One of the important early responses following PAMP perception is the activation of CDPKs, resulting in an expression of defence-related genes (Tena et al., 2011). Perception of PAMPs induces an immediate Ca$^{2+}$ influx that contributes to immune signaling through the activation of Ca$^{2+}$-binding proteins such as CDPKs (Romeis and
Herde, 2014). Several CDPKs are involved in plant immune signaling, mostly as positive regulators (Boudsocq and Sheen, 2013) and rarely as negative regulators (Monaghan et al., 2014). For example, CPK5, CPK4, CPK6, and CPK11 participate in immunity as early transcriptional regulators (Boudsocq and Sheen, 2013). In contrast, CPK28 negatively regulates immune signaling through controlling BIK1 turnover (Monaghan et al., 2014).

Although PTI and ETI share many signaling components, it has been proposed that immune responses in ETI occur more quickly, are more prolonged, and are more robust than those in PTI, suggesting that PTI is a weak variant of ETI (Thomma et al., 2011).

One of the big gaps in our understanding of plant immunity is in the signaling pathways that operate immediately downstream of PRR and R protein activation. Genetic screens have had very limited success in identifying signaling components, and thus the components of these pathways remain mostly elusive (Dodds and Rathjen, 2010).

1.3 Chitinases in plant immunity

As mentioned above, recognition of PAMPs leads to a transcriptional reprogramming within the plant cell, resulting in expression of immunity-related proteins. In fact, a group of plant-coded proteins induced by different stresses, named pathogenesis-related (PR) proteins, has been assigned an important role in plant defence against pathogens. This group of proteins has been defined as ‘proteins encoded by the host plant but induced only in pathological or related situations’ (Antoniw et al., 1980), and are historically classified into 17 PR protein families on the basis of their characteristics and biological activity (van Loon et al., 2006). Four PR protein families are classified as chitinases and are thought to play a role in defence through their hydrolytic activities towards major components of fungal cell walls (van Loon et al., 2006).

1.3.1 Chitinases

Chitinases (EC 3.2.1.14) catalyze the hydrolysis of the β-1,4-glycosidic bonds between the GlcNAc residues of chitin (Collinge et al., 1993). Chitin, the second most abundant
biopolymer on the planet (Shahidi and Abuzaytoun, 2005), is found in the outer skeleton of insects, fungi, yeasts, algae, crabs, shrimps, and lobsters, and in the internal structures of other invertebrates (Hamid et al., 2013). Interestingly, chitinases are present not only in chitin-containing organisms, like fungi and insects, but also appear in organisms that do not contain chitin, such as bacteria, higher plants and animals, and play important physiological and ecological roles (Nagpure et al., 2014).

Apart from chitin, chitinases can also hydrolyze chitin derivatives. For example, one of the chitinase substrates is chitosan, the deacetylated form of chitin (Tanabe et al., 2000). Likewise, lipochitooligosaccharides produced by rhizobia and functioning as nodulation factors during root nodule formation (Mergaert et al., 1997), essentially consist of an acylated chitin oligomeric backbone with various substitutions at the (non)reducing-terminal and/or nonterminal residues (den Hartog et al., 2003), and are also hydrolyzed by chitinases (Goormachtig et al., 1998). Moreover, bacterial peptidoglycan, a polymer of β-1,4-linked GlcNAC and N-acetylmuramic acid (MurNAc) residues, can be cleaved by plant chitinases (Collinge et al., 1993). Plant cell wall glycoproteins containing GlcNAC are considered to be the endogenous substrate for plant chitinases (Dyachok et al., 2002).

As far as the amino acid similarity of chitinases from various organisms is considered, 5 classes of chitinases have been proposed (Passarinho and de Vries, 2002), and have been categorized into 2 families, which include families 18 and 19 of glycosyl hydrolases (GH) based on the catalytic domain of chitinases (Henrissat, 1991). The GH18 chitinases include class III and V plant chitinase members (Takenaka et al., 2009). GH19 family is exclusively composed of chitinases of classes I, II, and IV (Santos et al., 2008). The chitinases of the 2 different families do not share amino acid sequence similarity, and have completely different 3-dimensional (3D) structures and molecular mechanisms. Therefore, they are likely to have evolved from different ancestors (Tyler et al., 2010).

1.3.2 Role of chitinases in plant defence
Bacterial and fungal chitinases often function in nutrition processes or in morphogenesis of the cell wall (Cohen-Kupiec and Chet, 1998). In contrast, the plant and animal chitinases mainly play a role in the host defence against pathogen attack (Kasprzewska, 2003), which can be confirmed by that fact that four PR protein families are plant chitinases (von Dach, 2006).

Plant chitinases have been shown to contribute to defence against fungi in two distinct manners. Exochitinases reside in the apoplast and are not considered directly detrimental to fungal growth, but are presumed to release fungal chitin oligosaccharides, which in turn trigger downstream defence responses (Kombrink et al., 2011). These responses include the production of endochitinases that accumulate in the vacuole and are released to directly inhibit the growth of the fungal hyphae during the infection process, when hyphae penetrate and affect cell integrity (Collinge et al., 1993).

In order to avoid plant chitinase-mediated defence, plant pathogens have evolved several counteracting strategies, including modification of carbohydrate chains (Ride and Barber, 1990), cell wall remodeling (Fujikawa et al., 2009), and secretion of effectors to inhibit chitinase activity (Marshall et al., 2011), scavenge chitin (van den Burg et al., 2006), or compete for receptor binding (de Jonge et al., 2010; Sanchez-Vallet et al., 2013).

In addition to biotic stress, chitinases are also involved in plant abiotic stress responses such as osmotic, salt, cold, wounding, and heavy metal stresses (Grover, 2012).

1.4 Peptidoglycan and its receptors as example for a PAMP-PRR pair

1.4.1 Peptidoglycan

Most eubacteria are entirely surrounded by a strong cell wall, usually namely a sacculus, to maintain the bacterial shape and counteract the internal pressure of the bacterial cell (Weidel and Pelzer, 1964). Besides its protective function and shape formation, the cell
wall also provides an interface for interaction with the surrounding environment and potential hosts.

The most important component of the bacterial cell wall conferring strength and rigidity is the heteropolymeric macromolecule peptidoglycan (PGN), firm glycan chains that are interlinked either directly or via short peptide bridges (Glauner et al., 1988). PGN is composed of three main building blocks: glycan strand, peptide subunit or stem peptide, and the inter-peptide bridge. The glycan strand is composed of alternating β (1,4) linked GlcNAc and MurNAc, which is cross-linked by short peptides attached by an amide linkage to the lactyl group of MurNAc.

The most common sequence of the stem peptide is L-Ala-D-iso-Glu-mDAP/L-Lys-D-Ala-D-Ala, with a dibasic amino acid (mostly either meso-diaminopimelic acid (mDAP) or L-Lys) residing at position three, hence distinguishing Dap-type from Lys-type PGN. DAP-type PGN is present in most Gram-negative bacteria as well as in Bacilli and Mycobacteria compared to the Lys-type in most Gram-positives (Schleifer and Kandler, 1972). Gram-positive and Gram-negative bacteria differ not only in the type of PGN they harbor, but also in the PGN amount. Gram-positive cell walls contain a thick PGN saccus of up to forty layers (20 - 80 nm), which is embedded with teichoic and lipoteichoid acids and proteins. Instead, Gram-negative cell walls mostly consist of a single layered PGN macromolecule (1-7 nm) and an additional membrane, the outer membrane (Cabeen and Jacobs-Wagner, 2005).

The PGN of bacteria is steadily remodeled and degraded during cell growth and differentiation as well as in response to changing environmental conditions. Up to 50% of the PGN is shed from the cell wall in one generation by cell wall turnover (Park and Uehara, 2008; Reith and Mayer, 2011). Thus it was proposed that particular PGN fragments can be exploited as signaling molecules for bacteria themselves, reflecting the status of their cell wall (Bertsche et al., 2015).

1.4.2 PGN perception systems
As signaling molecules, PGNs can be sensed by not only bacteria themselves but also by their host. PGNs have the features that are shared among all PAMPs. For instance, PGNs, as major building blocks of the bacterial cell walls, are highly conserved and exposed at the bacterial cell surface, and hence they constitute excellent targets for recognition by the innate immune system of potential host organisms that contain no PGNs. In fact, PGN has been known for a long time to promote innate immune responses in vertebrates, insects and plants (Dziarski and Gupta, 2005a; Gust et al., 2007; Erbs et al., 2008). PGN can be recognized in mammals by two intracellular receptors: Nucleotide-binding oligomerization domain-containing protein 1 (NOD1) and NOD2, which both contain an N-terminal caspase recruitment domain (CARD), a C-terminal LRR domain for ligand sensing, and a central nucleotide oligomerisation domain (NOD) (Tanabe et al., 2004; Inohara et al., 2005). NOD2 detects Lys-type PGN and muramyl dipeptide (MDP) as the minimal recognition structure, whereas NOD1 preferentially senses the mDAP-containing muramyl tripeptide found mostly in Gram-negative PGN (Chamaillard et al., 2003; Girardin et al., 2003). In addition, PGN can also be perceived by so called PGN-recognition proteins (PGRPs, PGLYRPs), highly conserved pattern-recognition receptors of the innate immune system present in insects and mammalian cells (Lu et al., 2006; Guan and Mariuzza, 2007; Dziarski and Gupta, 2010; Kurata, 2014). Mammalian PGRPs bind bacterial PGN and might even kill Gram-positive bacteria by directly interacting with PGN, thereby interfering with PGN maturation (Cho et al., 2007). Some structurally known PGRPs are able to bind PGN, but unable to hydrolyze it due to a lack of zinc-coordinating residues in the active site (Reiser et al., 2004; Guan et al., 2005). It was proposed that these non-lytic PGRPs may inhibit bacterial growth by enclosing parts of the PGN layer and thereby preventing further crosslinking (Bertsche et al., 2015). In contrast, there are also PGN-hydrolysing PGRPs in insects and mammals that have at least one C-terminal PGRP domain, which is homologous to bacteriophage and bacterial type 2 amidases such as the Drosophila PGRP-LB (Kim et al., 2003). Although it is not yet clear what role exactly PGN-hydrolysing PGRPs have during infection, it has been proposed that degradation of PGN might contribute to boosting the innate immune system and
coordinating the inflammation response by multiplication of PGN molecules recognized by NOD2 (Schaffler et al., 2014).

In addition to intracellular receptors of PGNs, mammals have also extracellular PGN receptors like membrane-bound Toll-like receptor 2 (TLR2) (Dziarski and Gupta, 2005b). The role of TLR2 as a specific PGN receptor has been heavily debated (Muller-Anstett et al., 2010). Recently, it has been shown that the NOD2 mediated activation of dendritic cells with polymeric PGN is dependent on a TLR2 co-stimulation. In other words, signaling via both receptors, NOD2 and TLR2, leads to a more potent activation of the immune system compared to stimulation of each receptor alone (Schaffler et al., 2014).

In plants, LysM proteins have been shown recently to act as receptors for PGN. The lysin motif (LysM), typically from 44 to 65 amino acids in length, is an ancient and ubiquitous protein domain (Pfam PF01476) found in all living organisms except for Archaea (Bateman and Bycroft, 2000). The LysM has a βααβ secondary structure with the two α-helices packing onto the same side of an antiparallel β-sheet (Zhang et al., 2007b). In prokaryotes, LysMs were found in bacterial lysins or chitinases, which hydrolyze glycosidic bonds of PGN or chitin (Buist et al., 2008). Thus, the GlcNAc moiety is generally thought to be the LysM ligand. Indeed, the plant LysM proteins characterized are implicated in recognition of GlcNAc-containing substrates such as chitin and PGN. For example, in Arabidopsis PGN can be recognized by a tripartite recognition system, composed of the two membrane-tethered LysM receptor proteins LYM1 and LYM3, and one transmembrane LysM receptor kinase CERK1 (Willmann et al., 2011). All three proteins are indispensable for PGN sensitivity and immunity to bacterial infection (Willmann et al., 2011). Similarly, the PGN perception system in rice is made of the two LysM receptor proteins LYP4 and LYP6 (Liu et al., 2012a). LYP4 and LYP6 were shown to physically interact with OsCERK1, which is also required for PGN perception (Kouzai et al., 2014).

1.5 From microbial complex structures to PAMPs

Plants are able to detect non-self via sensing microbial-derived PAMPs by PRRs. The efficient activation of PRRs relies on the form, amount, concentration, and size of PAMP ligands. It is generally little known how microbial patterns are released from complex
extracellular structures such as microbial cell walls. Two possible scenarios as to how soluble PAMP fragments might be generated from macromolecular assemblies of microbes are described below.

1.5.1 Generation of PAMPs through spontaneous release

During bacterial growth large amounts of PGN building blocks are steadily shed into the extracytoplasmic space. For example, *B. subtilis* releases about 50% of its PGN in one generation during growth (Goodell and Schwarz, 1985). It also been found that flagellin monomers are shed into the supernatant of *P. aeruginosa* cultures (Bardoel et al., 2011). Likewise, LPS is shed into liquid culture when *E. coli* grows *in vitro* (Mackowiak, 1984). Upon infection, these building blocks are shed into space surrounding cells and can serve as PAMPs for the activation of PTI signaling. For instance, culture supernatants from *Bacillus sp.* were shown to cause immune responses through activation of the NOD1 signaling cascade in response to mDAP containing cell wall peptides (Fujimoto and Fukase, 2011). *P. aeruginosa aprA* mutants induced an activation of TLR5 signaling via shedding monomeric flagellin into their environment (Bardoel et al., 2011). Similarly, fungal cells also shed chitin into the environment when growing (Bueter et al., 2013).

1.5.2 Generation of PAMPs through host hydrolytic enzymes

Upon infection, hosts often secrete hydrolytic enzymes to harm the invader and to either deliberately or incidentally produce soluble PAMPs. Eukaryotic PGN recogniton proteins (PGRPs), which are conserved from insects to mammals, bind PGN and function in antibacterial immunity (Bertsche et al., 2015). Non-enzymatic PGN sensors such as PGRP-SA depend on fragmented PGN for perception and immune stimulation, which is delivered by the PGN-hydrolytically active Gram-negative bacteria-derived binding protein 1 (GNBP1) (Wang et al., 2006). Apart from PGRPs, other PGN-hydrolytic enzyme activities such as lysozymes have been reported to functions in generating soluble PGN fragments a ligands for PRRs (Callewaert and Michiels, 2010).

Similarly, plants have also been reported to produce PAMPs through secreted hydrolytic enzymes. For instance, upon infection of fugal pathogens plant chitinases are often
induced to target fungal cell walls, releasing chitin as PAMPs (Sanchez-Vallet et al., 2015). Soluble PGN fragments have previously been shown to stimulate plant immune responses in *Arabidopsis* (Gust et al., 2007; Erbs et al., 2008). However, it is unknown how PGN PAMPs are generated, and if so, which host protein is involved in PGN hydrolysis in *Arabidopsis*.

### 1.6 Aims of the thesis

Overwhelming focus has been put on PTI studies since the innate immunity concept was first described in 1997 (Medzhitov and Janeway, 1997), but there are still many big gaps in our understanding of PTI, such as the molecular events pre- and post-recognition via the plasma-membrane located PRR complexes. For instance, although PGN and LYM1/LYM3/CERK1 have been identified as one PAMP-PRR pair in *Arabidopsis* (Gust et al., 2007; Willmann et al., 2011), upstream and downstream events of PGN perception remain largely elusive.

This thesis therefore focuses on the investigation of upstream and downstream molecular events of CERK1-mediated perception in plants.

It was generally proposed that soluble PAMPs derived from complex microbial cell walls serve as ligands for host PRRs and subsequent immune activation in plants. For example, macromolecular chitin was presumed to be hydrolyzed by exo-chitinases and released chitin oligomers can then act as PAMPs in PTI (Kombrink et al., 2011). However, it is unknown whether, and if so, how PGN is hydrolyzed prior to PGN-perception in *Arabidopsis*. The first aim of this thesis was thus to identify potential PGN hydrolases in *Arabidopsis* and afterwards characterize this enzyme(s), and finally to examine how the enzyme(s) might contribute to PGN recognition and thus to PGN-triggered immunity in plants.

CERK1 mediates both PGN- and chitin-induced immune responses (Miya et al., 2007; Willmann et al., 2011). But it is still unknown how the plant cell accomplishes the differentiation between bacterial PGN and fungal chitin by employing the same CERK1 receptor. Therefore, the second aim of the thesis was to decipher downstream signaling
components of CERK1, which might be the basis for differentiation of these two PAMP-mediated signaling pathways.
2. Materials and Methods

2.1 Materials

2.1.1 Plants

2.1.1.1 Arabidopsis thaliana

The experiments in this study were conducted using the Arabidopsis thaliana ecotypes Columbia-0 (Col-0), or Landsberg erecta (Ler) and transgenic lines generated in these ecotypes. The knock-out or knockdown lines used in this study, if not stated otherwise, were obtained from the Nottingham Arabidopsis Stock Centre (NASC) or received from other labs, and are listed in Table 1.

Table 1. Arabidopsis mutant and transgenic lines used in this study.

<table>
<thead>
<tr>
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<th>Gene</th>
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<th>stock</th>
<th>reference</th>
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<td>At5g24090</td>
<td>LYS1/CHIA</td>
<td>lys1-kd/chia-kd (amiRNA)</td>
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<tr>
<td>At3g21630</td>
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<td>cerk1-2 (T-DNA)</td>
<td>GABI_096F09</td>
<td>(Miya et al., 2007)</td>
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<tr>
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<td>lym3-1(T-DNA)</td>
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<tr>
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<td>(Willmann et al., 2011)</td>
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</table>
2.1.1.2 *Nicotiana benthamiana* and *Oryza sativa*

The proteins were transiently expressed in *Nicotiana benthamiana* to study protein interaction and protein activity. The pH assay was conducted using *Oryza sativa* and *Arabidopsis thaliana* cell cultures.

2.1.2 Microbes

Table 2. Bacterial strains used in this work

<table>
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<th>Strain</th>
<th>Genotype</th>
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<td>supE44 ΔlacU169 (Φ80 lacZM15) hsdR17 recA1 endA1 gyrA96 thi-1 relA1</td>
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<tr>
<td></td>
<td>TOP10</td>
<td>mcrA, delta (mrr-hsdRMS-mcrBC), phi 80 delta lac delta M15, delta lacX74,deoR, recA1, araD139 delta (ara, leu),7697, galU, galK, lambda-, rpsL,endA1, mupG</td>
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<tr>
<td></td>
<td>C58C1</td>
<td>T-DNA’ vir’ rif’, carb’</td>
</tr>
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</table>

2.1.3 Media and buffer
The media used in this study were sterilized by autoclaving for 20 minutes at 121°C and are summarized in Table 3. All buffers were prepared according to (Molecular cloning, 3rd edition, Sambrook and Russell). Antibiotics were added, if necessary, to the sterilized medium in appropriate final concentrations as listed in Table 4.

### Table 3. Media used in this study

<table>
<thead>
<tr>
<th>Media</th>
<th>Ingredients (1 liter)</th>
<th>Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>Luria-Bertani broth (LB)</td>
<td>10 g Bacto-Tryptone, 5 g NaCl, 5 g Yeast extract (YE)</td>
<td><em>Escherichia coli</em></td>
</tr>
<tr>
<td>Kings Medium B (King's B)</td>
<td>20 g glycerol, 40 g Proteose Pepton 3, after autoclaving addition of 0.1% (v/v) MgSO4 and KH2PO4</td>
<td><em>Pseudomonas syringae</em></td>
</tr>
<tr>
<td>½ Murashige-Skoog Medium (½ MS)</td>
<td>2.2 g MS (Duchefa), pH 5.7 (KOH)</td>
<td><em>Arabidopsis thaliana</em></td>
</tr>
</tbody>
</table>

### Table 4. Antibiotics used in this study

<table>
<thead>
<tr>
<th>Antibiotics</th>
<th>Working concentration (µg/ml)</th>
<th>solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbenicillin</td>
<td>50-100</td>
<td>Water</td>
</tr>
<tr>
<td>Cycloheximid</td>
<td>50</td>
<td>Water</td>
</tr>
<tr>
<td>Kanamycin</td>
<td>50</td>
<td>Water</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>50</td>
<td>Methanol</td>
</tr>
<tr>
<td>Spectinomycin</td>
<td>100</td>
<td>Water</td>
</tr>
<tr>
<td>Tetracycin</td>
<td>50</td>
<td>Ethanol</td>
</tr>
<tr>
<td>Ampicillin</td>
<td>50-100</td>
<td>Water</td>
</tr>
<tr>
<td>Gentamycin</td>
<td>25</td>
<td>Water</td>
</tr>
<tr>
<td>Hygromycin</td>
<td>50</td>
<td>Water</td>
</tr>
</tbody>
</table>

### 2.1.4 Vectors

Table 5. Vectors in this study
Materials and Methods

<table>
<thead>
<tr>
<th>Vector</th>
<th>Characteristics</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>pDONR201</td>
<td>Ori Puc, rnrB, T2, rnrB,T1, attP1, attP2, ccdB,Cm(^{r}), Kan(^{r})</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pDONR207</td>
<td>Ori Puc, rnrB, T2, rnrB,T1, attP1, attP2, ccdB,Cm(^{r}), Gent(^{r})</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pK7FWG2.0</td>
<td>P35S, T35S, eGFP, attR1, attR2, ccdB, Cm(^{r}), Kan(^{r})</td>
<td>VIB</td>
</tr>
<tr>
<td>pDEST17</td>
<td>PT7, RBS, His(_{6})-tag, attR1, attR2, ccdB, Cm(^{r}), PT7, bla, Promotor, Amp(^{r}), pBR322 origin, ROP, orf</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>miR319a pBSK</td>
<td>B reverse, T3 promotor, miR319a, T7 promotor, A forward, Amp(^{r})</td>
<td>Weigelworld.org</td>
</tr>
<tr>
<td>(pRS300)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pCR8/GW/TOPO</td>
<td>Entry vector for the Gateway system</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pGWB5/14/17</td>
<td>Binary gateway destination vector</td>
<td>(Nakagawa et al., 2007)</td>
</tr>
</tbody>
</table>

2.1.5 Enzymes and antibodies

Restriction enzymes, ligase and DNA modification enzymes were purchased from Fermentas (St. Leon-Rot) and New England Biolabs (Beverly, USA). Mutanolysin was purchased from Sigma-Aldrich. Antibodies were purchased from the companies Sigma-Aldrich (Taufkirchen), New England Biolabs (Beverly, USA) and Acris Antibodies GmbH (Herford) and are listed in Table 6. The antibody against tobacco class III chitinases from rabbit and the antibody against YFP from rabbit were kind gifts from Dr. Frédéric Brunner and Dr. Sara Mazzotta, respectively. The antibody against AtCERK1 was kindly provided by Dr. Gary Stacey (University of Missouri, USA).

Table 6. Antibodies used in this study

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Host</th>
<th>Dilution</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary antibodies</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>α-p42/44 MAPK</td>
<td>rabbit</td>
<td>1:3000</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>α-HA</td>
<td>rabbit</td>
<td>1:3000</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>α-MYC</td>
<td>mouse</td>
<td>1:3000</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>α-GFP</td>
<td>goat</td>
<td>1:3000</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>Secondary</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>α-mouse IgG HRP</td>
<td>rabbit</td>
<td>1:10000</td>
<td>Sigma-Aldrich</td>
</tr>
</tbody>
</table>
antibodies | conjugated | species | dilution | supplier  
--- | --- | --- | --- | ---  
α-goat IgG HRP conjugated | rabbit | 1:10000 | Sigma-Aldrich  
α-rabbit IgG HRP conjugated | goat | 1:10000 | Sigma-Aldrich  
Anti-goat IgG-AP | rabbit | 1:3000 | Sigma  
Anti-rat IgG-AP | goat | 1:3000 | Sigma  
Anti-mouse IgG-AP | rabbit | 1:3000 | Sigma  
Anti-rabbit IgG-AP | goat | 1:3000 | Sigma

### 2.1.6 Chemicals and solutions

Laboratory grade chemicals and reagents were purchased from Sigma-Aldrich (Taufkirchen), Carl Roth (Karlsruhe), Merck (Darmstadt), Qiagen (Hilden), Invitrogen (Karlsruhe), Duchefa (Haarlem, Niederlande), Molecular Probes (Leiden, Niederlande), Fluka (Buchs, Schweiz) and BD (Sparks, USA), unless noted otherwise in the text. All buffers and solutions were prepared, if not stated otherwise, with milli-Q water. Sterilization was conducted by autoclaving or filter sterilization.

### 2.1.7 Oligonucleotides

The oligonucleotides used in this study were synthesized by Eurogins MWG Operon. The sequences of these oligonucleotides are listed in the Appendix Table 7.

### 2.1.8 PAMPs

*Bacillus subtilis* PGN was purchased from Invivogen (San Diego, CA). *Micrococcus luteus* cells were purchased from Sigma-Aldrich (Hamburg, Germany). Chitin hexamer was purchased from Carbosynth (UK), chitin octamer was purchased from Seikagaku (Japan), and crab chitin was purchased from Sigma-Aldrich. All PAMPs, unless stated otherwise, are dissolved or resuspended in ddH$_2$O.

Flg22 peptide QRLSTGSTINSAKDDAAGLQIA was purchased from Selleckchem company and was dissolved in ddH$_2$O at high concentration (1 M) and in 1mg/mg BSA.
and 0.1M NaCl at low concentration (1mM). Elf18 peptide SKEKFERTKPHVNVGTIG was a gift from George Felix.

2.2 Methods

2.2.1 Plant growth

2.2.1.1 Growth conditions

*Arabidopsis* seeds were sown on steam-sterilized GS90-soil (Gebr. Patzer GmbH) mixed with vermiculite or after surface-sterilization with chlorine gas on sterile ½ MS plates. After stratification of the seeds for two days at 4°C in the dark, the plants were grown in either long-day (16 h light, 8 h darkness) or short-day (8 h light, 16 h darkness) environmental chambers under standard conditions (150μmol/cm²s light, 40-60 % humidity, 22°C). *N. benthamiana* plants were cultivated in a mixture of soil and sand containing 0.1 % (v/v) Confidor in the greenhouse (13 h light, 11 h darkness).

2.2.1.2 Seed surface sterilization

*Arabidopsis* seeds were sterilized by chlorine gas treatment. Seeds were transferred to Eppendorf tubes and placed in a desiccator. In a glass beaker placed in the desiccator, 2 ml of 37 % HCl were added to 50 ml of 12 % sodium hypochloride solution forming the chlorine gas. The lid of the desiccator was immediately closed and a vacuum was generated to get an airtight seal. The seeds were incubated for 4 h.

2.2.1.3 Cell suspension cultures

*Arabidopsis* and *Oryza sativa* suspension cell cultures were grown in MS medium (4.41g/l MS salt, 6% sucrose, 50 mg/l MES, 2mg/l 2, 4-D) at 150 rpm and sub-cultured every week.

2.2.2 Microbe cultivation

2.2.2.1 Growth of *Escherichia coli*
E. coli strains were cultivated overnight at 37°C either on LB-plates in oven or in liquid LB medium at 230 rpm in shaker. Antibiotics were added into the media according to the resistance cassettes of each strains.

2.2.2.2 Growth of Pseudomonas syringae

P. syringae pv. tomato DC3000 were grown for 24-48 hours at 28°C either on King’s B-plates in an oven or in liquid King’s B medium at 180 rpm in a shaker. For the determination of bacterial growth in infection assays the Pseudomonas strains were re-isolated from plant material and plated on LB plates containing cycloheximide in addition to other antibiotics.

2.2.2.3 Growth of Agrobacterium tumefaciens

A. tumefaciens strains were cultivated for 48 hours at 28°C on LB-plates in an oven or in liquid LB medium at 230 rpm in a shaker. Additional antibiotics were added into the media according to the plasmid-DNA the strains were carrying.

2.2.3 Molecular biology

2.2.3.1 Isolation of plasmid DNA from E.coli

For alkaline lysis, a bacterial pellet from a 2 ml overnight culture was resuspended in 100 µl lysis buffer (50 mM Tris/HCl pH 8.0, 50 mM EDTA pH 8.0, 15 % Saccharose), and lysed with 200 µl alkaline SDS-buffer (200 mM NaOH, 1 % (w/v) SDS) for a maximum of 5 min and finally 150 µl Potassium acetate buffer (3 M Potassium acetate, 11.5 % (v/v) acetic acid) was added. The mix was centrifuged and the aqueous phase containing plasmid DNA was precipitated with 0.6 Vol Isopropanol. The pellet was washed with 70 % ethanol and dissolved in TE-buffer (10 mM Tris/HCl, 1 mM EDTA, pH 8) or deionized water.

Alternatively to the classical alkaline lysis, for mini scale isolation, plasmid DNA was extracted from 3 ml overnight culture using the MiniPrep Kit (Thermo Scientific) according to the manufacturer’s instructions.
For midi scale isolation, plasmid DNA was extracted from 25 ml (high copy plasmid) or 100 ml (low copy plasmid) overnight culture using the PureLink HiPure kit (Invitrogen) according to the manufacturer’s instructions.

For large scale plasmid isolation, a manual protocol was used as described below. A single colony was pre-cultured in 2.5 ml LB for 3 hours and was enlarged with pre-warmed 500 ml LB culture for overnight growing at 37°C. Cells were collected by centrifuging for 30 minutes at 5000 rpm and manipulated from this step onward on ice or at 4°C. Bacterial pellets were suspended in 2.5 ml freshly-prepared lysozyme solution (10 mg/ml), mixing them by up-and-down pipetting followed by 5 minutes incubation on ice. 2 ml of 0.5 M EDTA pH 8 was added, mixed and incubated for 5 minutes on ice. The solution was mixed with 100 µl RNAase A (20 mg/ml) and 150 µl of 10% Triton X-100 and topped up to 1 ml with 0.02xTE. The mixture was then incubated for 60 minutes on ice. The supernatant was separated by centrifugation at 18000 rpm for 60 minutes and then transferred to a new tube and mixed 1:1 with phenol (equilibrated with 0.1% 8-hydroxyquinoline pH 8) and shaken vigorously for 1 minute. The solution was centrifuged for 20 min. The upper aqueous phase was carefully recovered and transferred to a new tube containing 1 volume chloroform, followed by shaking and spinning as before. The aqueous phase was recovered to a 30 ml corex tube, adding 10% 5M NaClO₄ of water volume and 80% isopropanol of water volume. The tube was sealed with parafilm and mixed by turning upside down. Then the tube was centrifuged at 10000 rpm for 15 minutes. The pellet was air-dried, and re-suspended in 500 µl TE by incubation overnight. The DNA was checked by agarose gel electrophoresis (2.2.3.6).

### 2.2.3.2 Genomic DNA isolation from plant material

Leaf pieces were ground in a 1.5 ml Eppendorf tube with 200 µl Edwards buffer (200 mM Tris/HCl pH 7.5, 250 mM NaCl, 25 mM EDTA pH 8, 0.5 % (w/v) SDS) using a homogenizer machine. The samples were centrifuged for 5 minutes at 13000 rpm. The supernatant was transferred to a new tube with the same volume isopropanol and thoroughly mixed. Genomic DNA was precipitated at room temperature for 5-10 minutes, and centrifuged at 4°C for 10 minutes at 14000 rpm. The pellet was washed with 200 µl
ethanol (70%, v/v) and was air-dried at room temperature. Finally, the DNA pellet was dissolved in ddH$_2$O.

### 2.2.3.3 Total RNA isolation from *Arabidopsis* seedlings

3-5 seedlings (7-10 day old) were harvested in a 1.5 ml Eppendorf tube and frozen in liquid N$_2$. The tube was placed on a precooled plate and the seedlings were ground into fine powder with a precooled pestle. 300 µl Trizol (Chomczynski and Sacchi, 1987) was added immediately to the tube, followed by vortex mixing for 10 seconds and incubation for 10 minutes at room temperature. 60 µl chloroform was then added, and the mixture was vortexed again. The samples were incubated for 10 minutes at room temperature and then centrifuged at 14000 rpm for 10 minutes at room temperature. The upper aqueous phase was transferred into a new tube filled with 150 µl isopropanol and mixed by vortex. The samples were incubated for 1 hour at -80°C or overnight at -20°C. The precipitated RNA was collected by centrifugation at 14000 rpm for 10 minutes at 4°C. The RNA pellet was washed with 500 µl 70% (v/v) ethanol. Finally, the air-dried pellet was dissolved in 10 µl ddH$_2$O.

### 2.2.3.4 Complementary DNA synthesis by Reverse Transcription (RT)

2-5 µg total RNA in 1.5 µl was mixed with 0.5 µl 30 µM oligo-dT and denatured at 70°C for 10 minutes. Then 8 µl RT-mixture (2 µl 5x buffer, 1µl 2.5 mM each dNTPs, 0.5 µl RNAase inhibitor (Fermentas), 1 µl reverse transcriptase (Fermentas), 4.25 µl H$_2$O) was added and incubated at 42°C for 90 min, followed by 70°C for 10 min.

### 2.2.3.5 Standard PCR

In a 30µl reaction volume, 1 µl template was added into the mixture (3 µl 10x buffer, 0.5 µl 2.5 mM dNTPs, 0.5 µl 10 µM forward and reverse primer, 0.5 µl lab-made Taq polymerase). The reaction was performed in a PCR machine (PTC-200) with the following program: 94°C for 2 min, 30 cycles of 94°C 30 seconds, 55°C for 30 seconds, 72°C for 1 min/kb, 72°C 10 min.

### 2.2.3.6 DNA agarose gel electrophoresis

...
DNA electrophoresis was performed on a 0.8-1.5 % agarose gel in 1x TAE buffer (40 mM Tris/acetate pH 8.0, 2 mM EDTA pH 8.0) at 60-120 V. A 1 kb ladder (Fermentas) was used as size marker. Ethidium bromide (0.5 µg /ml) or 6% (v/v) peqGREEN (Peqlab) present in the gel helped the visualization of DNA by a UV-Transilluminator (Infinity-3026 WL/26 MX, Peqlab).

2.2.3.7 Sequencing

The constructs and PCR products were sequenced by GATC Biotech AG (Konstanz). 5µl DNA template with either 80-100 ng/µl plasmid or 20-80 ng/µl PCR product was added to 5 µl 5 µM sequencing primer. The results were analyzed using DNAsstar or CLC main workbench software.

2.2.3.8 Quantitative fluorescent real time PCR

2.5 µL seedling RNA or 1 µg leaf RNA was used for the cDNA synthesis (in 5µl total reaction volume). Leaf cDNA was diluted 3 to 5 fold for RT-qPCR experiments, whereas seedling cDNA was used undiluted. RT-qPCR amplifications and measurements were performed with the iQ5 Real Time PCR detection system from Bio-Rad. RT-qPCR amplifications were monitored using the SYBR Green Fluorescein Mix (Thermo Scientific). The gene expression data was quantified using the $2^{-\Delta\Delta CT}$ method (Livak and Schmittgen, 2001). The normalization of the expression levels was done using the CT values obtained for the EF-1α gene. The presence of a single PCR product was further verified by dissociation analysis in all amplifications. All quantifications were made in duplicate on RNA samples obtained from three independent experiments, each performed with a pool of two leaves or 3-5 seedlings.

2.2.3.9 Restriction endonuclease digestion

DNA was digested in 20µl reaction volume with 1 U/µl DNA at 37°C (or the appropriate temperature for the given restriction endonuclease) for 1 hour according to the manufacturer's recommendations.

2.2.3.10 Isolation of DNA fragments from agarose gels
Materials and Methods

PCR or DNA fragments were separated via agarose gel electrophoresis. DNA bands were excised from the gel with clean razor blades and extracted using the GeneJet Gel Extraction Kit (Fermentas) according to manufacturer’s manual.

### 2.2.3.11 DNA ligation

In 10µl reaction volume, 50 ng vector and insert DNA were incubated with ligation buffer and T4 DNA ligase (Fermentas) at 16°C overnight.

### 2.2.3.12 Gateway cloning

In a BP reaction, 100 fmol of the insert and 100 fmol of donor vector were mixed with 2 µl BP recombinase mix (Invitrogen). After brief vortexing, the mix was incubated overnight at 25°C. The reaction was stopped by adding 1 µl of Proteinase K (Invitrogen) and the samples were incubated for 15 min at 37°C. 2µl of the reaction volume was used to transform *E. coli* DH5α (2.2.3.13).

To add an A-overhang to PCR products, 40 µM dATP was added to the reaction mixture after PCR and the PCR product was insert into the pCR8 vector using the pCR8/GW/TOPO Cloning Kit (Invitrogen) to generate an entry clone. Then, a LR reaction was performed to create a destination construct. 200 to 250 ng of the destination vector and TE buffer pH 8.0 up to a final volume of 4 µl was added to a reaction mix containing 50 to 150 ng of the entry vector 1 µl of the LR Clonase II mix (Invitrogen) was added and the mixture was incubated overnight at 25°C. The reaction was stopped by adding 1 µl of Proteinase K (Invitrogen) and the samples were incubated for 15 min at 37°C. 2 µl of the reaction volume was used to transform *E. coli* DH5α (2.2.3.13).

### 2.2.3.13 Preparation and transformation of chemically competent *E.coli* cells

A single colony of *E. coli* DH5α was grown in 5ml LB overnight at 37°C and 250 rpm shaking. 400 ml of LB medium was inoculated with 4 ml overnight culture in sterile 2-liter flasks and grown at 37°C and 250 rpm to an OD590 = 0.375. The culture was divided into 8 pre-chilled falcon tubes and incubated for 10 minutes on ice. After centrifugation at 1600g and 4°C for 7 minutes, pellets were suspended in 10 ml ice cold CaCl₂ solution
(60 mM CaCl$_2$, 15% Glycerol, 10 mM PIPES pH 7.0) and incubated on ice for 30 minutes, followed by washing 2 times by centrifuging at 1100 x g and 4°C for 5 minutes and then incubation for 30 minutes on ice. The pellet was resuspended in 2ml of CaCl$_2$ and again put on ice, then aliquoted in 50µl into chilled Eppendorf tubes on ice and frozen in liquid nitrogen and stored at -80°C.

50µl chemically competent cells were taken from -80 °C and thawed on ice. Plasmid or ligation or recombination products were added to the aliquot and kept on ice for 20 minutes. The mixture was heat-shocked at 42°C for 45 seconds and immediately put on ice for 2-3 minutes. Then 800µl LB medium was added. The mixture was incubated at 37°C for 1 hour on a shaker. 200µl transformed cell culture was spread onto LB medium plates with appropriate antibiotics.

2.2.3.14 Preparation and transformation of electrical competent cells of A. tumefaciens

The agrobacterium strain was streaked out onto a LB plate with selective antibiotics and grown at 28°C for 2 days. A single colony was picked to inoculate 5 ml LB liquid medium with appropriate antibiotics. The culture was grown overnight at 28 °C and then enlarged with 500ml LB and grown to an OD600 of 0.5-1.0. Subsequently, the culture was chilled on ice for 15-30 minutes. From this time point onwards the cells were manipulated at 4°C. The cells were centrifuged at 4500 rpm for 15 minutes at 4°C and the pellet was suspended in 200 ml ice-cooled sterile water. The cells were again centrifuged at 4500 rpm for 15 minutes at 4°C and suspended in 100 ml of ice-cooled sterile water and centrifuged as described above. The cells was suspended in 4ml of ice-cold 10% glycerol and centrifuged as described above. The cells were resuspended in 1 ml of ice-cold 10% glycerol. 40 µl aliquots were frozen in liquid nitrogen and stored at -80°C.

40 µl electrically competent cells were thawed on ice for 20 minutes and 100 ng plasmid DNA was added, the mixture then was transferred to a precooled electroporation cuvette. The cells were pulsed once with 1500 voltage for 5 ms (Eppendorf, Hamburg), the cuvette was put back on ice and immediately 500 µl LB medium was added to the
Cuvette. Cells were quickly resuspended by gentle pipetting up and down and then transferred to a 1.5 ml Eppendorf tube. The tube was incubated for 1 hour in a rotary shaker at 28°C. The mixture was plated onto selective LB plates and incubated for 2 days in a 28°C incubator.

2.2.4 Plant methods

2.2.4.1 Isolation of mesophyll protoplasts from *Arabidopsis*

Isolation of mesophyll protoplasts from leaves of 4-5 week-old *Arabidopsis* plants was performed according to the protocol of (Yoo et al., 2007) with minor changes. Briefly, thin leaf strips were dipped into 1.5% cellulose ‘Onozuka’ R10 – 0.4% macerozyme R10 solution (Yakult Pharmaceutical Industry), vacuum-infiltrated for 30 min and digested for 3 h at 20°C in the dark. After two subsequent washing steps with W5 buffer *Arabidopsis* protoplasts were suspended in MMG buffer to a concentration of 2*10^5 cells/ml prior to polyethylene glycol-mediated transfection. 100 µg plasmid DNA/ml protoplast suspension was used during transfection. Protoplasts samples were then incubated in W1 buffer at 20°C in the dark for 12 to 16 h allowing plasmid gene expression.

2.2.4.2 Stable transformation of *Arabidopsis thaliana*

*A. thaliana* plants were stably transformed by the floral dip-method (Clough and Bent, 1998). 500 ml liquid LB medium containing appropriate antibiotics was inoculated with a pre-culture of selected agrobacteria and further cultivated for 18 – 24 hours. The cells were pelleted for 20 minutes at 4500 x g and resuspended in fresh 5 % (w/v) saccharose solution at a density of 0.8 (OD600 nm). After addition of 0.02 % (v/v) Silwet young Arabidopsis inflorescences were dipped for one minute into the bacterial suspension. Afterward the plants were incubated at 100 % humidity for 24 hours. Seeds from floral-dipped plants were then screened for resistance against Basta (glufosinate-ammonium) or kanamycin.

2.2.4.3 Transient transformation of *Nicotiana benthamiana*

*Agrobacterium tumefaciens*-mediated transient transformation was used for the transient expression of proteins in *N. benthamiana*. The bacterial strain carrying the appropriate
expression vector was cultured as described in 2.2.2.3. After cells were harvested at 4°C for 10 minutes at 2000 x g, they were washed two times with 10mM MgCl₂. The density of the culture was diluted to 0.1 of OD600 and 150 μM acetosyringone was added. The bacterial suspension was then incubated by shaking at RT for 3-6 hours. Afterwards the suspension was mixed 1:1 with a suspension of bacteria carrying an expression construct of p19 (Voinnet et al., 2003) and the mixture was then infiltrated into 3 week-old tobacco leaves. The leaf tissue was analyzed 2-4 days post infection for the presence of the protein.

2.2.4.4 Generation of knock-down lines

Artificial microRNA-mediated gene silencing was used to specifically knock-down cpk15 in the Arabidopsis Col-0 background. The Web microRNA Designer (WMD; http://wmd.weigelworld.org) was used to select the primers (see Appendix Table 8) for the generation of an artificial 21mer microRNA (Schwab et al., 2005). The insert was generated in four PCR-steps with pRS300 as PCR template and then cloned into the pCR8 vector. The amiRNA in pCR8 was then cloned into the pB2GW7 destination vector. The transformation of the resulting vector into agrobacteria was mediated using electrical transformation. The stable transformation of the construct into the Arabidopsis genome was performed using the floral dip method. Offspring were screened for phosphinothricin (Basta) resistance. Analysis of the cpk15 transcript level in the cpk15 knock-down line (cpk15-kd) was performed by quantitative RT-PCR using primers listed in Table 8.

2.2.4.5 Genotyping analysis of T-DNA insertion lines

The T-DNA lines used in the frame of this work were analyzed for their genotype. Since diploid plants contain two copies of each gene and are thus able to segregate in offspring, it was necessary to confirm that the T-DNA insertion lines used for the experiments were homozygous. The discrimination between WT, heterozygous insertion and homozygous insertion lines was achieved by two sets of PCR reactions. In the WT-PCR, two primers bind two regions flanking the T-DNA insertion and thus amplify the
product only in the WT plants, because the large size of the T-DNA insertion inhibits the amplification in mutants.

In the second PCR a T-DNA specific left border amplifying primer (Lba primer) is used in a combination with a gene-specific primer allowing an amplification product only in plants carrying a T-DNA insertion. Thus, homozygous plants should show a product only in the Lba-PCR whereas heterozygous plants produce an amplicon in both WT-PCR and Lba-PCR.

**2.2.5 Protein biochemistry**

**2.2.5.1 Protein extraction from plant tissue**

Total protein was extracted from plant tissue using either a protein extraction buffer specific for acidic chitinases (20mM sodium acetate, pH 5.2/15mM β-mercaptoethanol supplemented with 1 proteinase inhibitor cocktail tablet/10ml from Roche) or an extraction buffer containing detergents for solubilization of membrane-bound proteins (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 % (v/v) Nonidet P40 and 1 protease inhibitor cocktail tablet/10 ml from Roche). The plant tissue was first homogenized in liquid N₂ and after addition of the extraction buffer the sample was incubated for 30 minutes on ice. Afterwards the soluble proteins were separated from cell debris using centrifugation at 4°C and the supernatant was used for further analysis.

**2.2.5.2 Immunoprecipitation**

Leaf protein was extracted from the *LYS1* overexpression plants and approximately 200 μg total proteins was used for the immunoprecipitation of LYS1-GFP. Next, protein extracts were incubated for 90 minutes at 4°C with gentle rotation either with 15 μl α-YFP (rabbit) or α-GFP (goat) antibody (Acris). In control protein samples no antibody was added.

Meanwhile, 400 μl agarose A bead solution (Roche) was washed three times with 800 μl water (1 min 2000 rpm 4°C), and once with 800 μl buffer A (50 mM Tris-HCl, pH 7.5, 5 mM EDTA, pH 8, 2 mM DTT, proteinase inhibitor cocktail). Finally, the agarose A beads were resuspended in buffer A (600 μl) and 50 μl bead solution was incubated with the
protein/antibody mixture for an additional 30 minutes in a rotator at 4°C. Afterwards, the beads were washed two times with 500 μl buffer A (1 min 1500 x g 4°C) and once with 500 μl buffer A containing 1 M NaCl. The immunoprecipitated proteins were then further analysed by immunoblot or activity assay.

2.2.5.3 LYS1 purification

500g 5-week-old leaves from LYS1OE leaves were frozen in liquid nitrogen and ground to a fine powder. Buffer A (20mM NaAc, pH5.2, 0.01% β-Mercaptoethanol) was added to the sample and incubated on ice for 4 hours. The sample was filleted through four layers of cheesecloth. The homogenate was centrifuged at 10000g for 30 minutes. The supernatant was loaded on a cation exchange column (SP Sepharose, GE Healthcare, Germany) equilibrated with buffer A. The proteins were eluted with a 0 to 1M NaCl gradient in buffer A. The elution fractions were monitored for LYS1 activity by the 4-MUCT assay and LYS1 presence was further confirmed by SDS-PAGE, coomassie blue staining and mass spectrometry. 4-MUCT active fractions were pooled and concentrated with vivaspin columns (GE Healthcare). Protein concentration was determined using the Bradford assay.

2.2.5.4 Transient expression and co-immunoprecipitation

Single colony Agrobacterium tumefaciens containing the indicated construct were inoculated in 3 ml liquid LB overnight at 37°C shaker. 100 μl culture was enlarged in 5ml fresh LB media in a 15 ml falcon tube overnight at 37°C shaker. The cells were collected by centrifugation for 10 min at 5000 rpm. The pellet was washed 2 times with 10 mM MgCl2 and resuspended in 10mM MgCl2 and 150µm acetosyringone to OD6001.0. The culture was keep at room temperature for 1-4 hours and mixed 1:1 with a p19 culture. The mixture was syringe-infiltrated into 4-week-old N. benthamiana leaves. After 2 days, the leaves were harvested by immediate freezing in liquid nitrogen. For total protein extraction, leaves were ground in liquid nitrogen and extraction buffer (50 mM Tris-Cl pH7.5, 150 mM NaCl, protease inhibitor cocktail (Roche) and phosphatase inhibitor, and 0.5% NP40) were added. The mixture was rotated at 4°C for 1 hour at 5-7 rpm and centrifuged at 13000rpm for 15 minutes at 4°C and repeated 2-3 times. Meanwhile, the
GFP-Trap beads (chromotek) were washed 3 times with extraction buffer. The supernatant was added to 40 µl beads and rotated at 5 rpm for 2 hours at 4°C. The bead mixture was washed 1 time with extraction buffer, 1 time with washing buffer 1 (50 mM Tris-Cl pH7.5, 150 mM NaCl) and 1 time with washing buffer 2 (50 mM Tris-Cl pH7.5) by spinning down at 2000 rpm for 1 minute at 4°C. As much of the supernatant as possible was removed from the beads, which were then resuspended in 40 µl SDS loading buffer. Then the samples were subjected to Laemmli SDS-PAGE for western blot detection.

2.2.5.5 Determination of protein concentration

The protein concentration was measured using the Bradford method (Bradford, 1976) and the Roti-Quant solution (Carl Roth). A standard curve was calculated using bovine serum albumin (BSA).

2.2.5.6 SDS-PAGE

SDS polyacrylamide gel electrophoresis was performed using the gel chamber system PROTEAN II from BioRad. 12 % SDS-PA gels were used as separating gels (with 5 % stacking gels) for the discontinuous SDS-PAGE using the Laemmli method (Laemmli, 1970) unless mentioned otherwise. The Pre-stained Protein Ladder Mix (Fermentas) was used as a protein marker.

2.2.5.7 Coomassie Brillant Blue staining

Proteins present in a gel or on a membrane after Western blotting were colored with staining solution (0.125 % (w/v) Coomassie blue R-250, 50 % (v/v) MeOH; 10 % (v/v) acetic acid). After incubation for 10 min at RT the unspecific stain was removed by destaining solution (50 % (v/v) MeOH, 10 % (v/v) acetic acid).

2.2.5.8 Western blot analysis

For the western blot analysis the proteins were transferred after SDS-PAGE onto a Hybond nitrocellulose membrane (GE Healthcare) using a Mini Trans-Blot® Electrophoretic Transfer Cell (BioRad) for one hour at 100 V. The protein transfer was controlled by Ponceau S red stain (0.1 % (w/v) Ponceau S red and 5 % (v/v) acetic acid).
Unspecific binding sites were blocked by incubation of the membrane for 1 hour at RT with 5 % (w/v) milk in either 1 x TBST (150 mM NaCl, 20 mM Tris-HCl; pH 7.6 and 0.1 % (v/v) Tween 20) or 1 x PBST (140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, 0.1 % (v/v) Tween 20).

Afterwards the membrane was incubated with a primary antibody overnight at 4°C. Then the membrane was washed for 3 x 5 minutes with 1 x TBST or 1 x PBST and incubated for 1.5 hours with a secondary antibody. The signal of a horseradish peroxidase-coupled secondary antibody was detected using the Enhanced Chemiluminescence Kit (GE Healthcare) according to the manufacturer’s instructions. For the detection of an alkaline phosphatase-coupled secondary antibody the membrane was washed with 1 x TBST for 3 x 5 minutes and then equilibrated for 2 minutes with a Tris 9.5-buffer (150 mM Tris-HCl; pH 9.5, 5 mM MgCl2 and 100 mM NaCl). The staining reaction was performed with 1 x BCIP/NBT in Tris 9.5-buffer (5-bromo-4-chloro-3-indolylphosphate; 200 x stock solution 50mg/ml in 70 % (v/v) dimethylformamide; Nitro-blue tetrazolium chloride; 200 x stock solution 50mg/ml in 100 % (v/v) dimethylformamide). After staining the membrane was washed with water.

2.2.5.9 CTAB western blotting

After CTAB-PAGE separation (zymography, 2.2.5.14), proteins were transferred to a nitrocellulose membrane with 1/20 lower buffer with 0.05% CTAB, 200 ml methanol 200 V 350 mA for 2 hours. The subsequent steps were the same as with western blotting above (2.2.5.8).

2.2.5.10 MAPK kinase assay

Total plant crude protein extract was isolated and subjected to a 12 % SDS-PAGE with 20 µl per lane, transferred to a nitrocellulose membrane, and detected with a primary antibody against phospho-p44/42-MAPK and secondary antibody.

2.2.5.11 Turbidity assay (PGN-hydrolysis assay)

The turbidity assay was performed as described in Park et al. (Park et al., 2002). In brief, lytic activity towards Micrococcus luteus cell wall or Bacillus subtilis peptidoglycan
(Sigma, Invivogen) was measured and compared to that of 1 µg hen egg white lysozyme (HEWL, Sigma). 1 ml 0.02 % (w/v) M. luteus cells or PGN in 20 mM sodium acetate, pH 5.2 were incubated at 37°C together with the enzyme, and the decrease in absorbance at 570 nm of the suspension was measured with a spectrophotometer over time. Approximately 60 µg total protein of the leaf extract and 15 µg total proteins of the protoplast samples were added to the reaction solutions.

**2.2.5.12 4-MUC cellulose assay**

The cellulose hydrolysis assay was performed using 4-methylumbelliferyl-β-D-cellobioside (4-MUC, Sigma-Aldrich) as substrate. 1 mM 4-MUC was incubated in 20 mM sodium acetate (pH 5.2) at 37°C for 1 hour in a 96 well plate with either 40 µg purified LYS1 or cellulose (Duchefa, Haarlem, The Netherlands) in a total volume of 100 µl. The reaction was stopped with 0.2 M Na₂CO₃ and the intensity of the fluorescence was monitored with an MWG Sirius HT fluorescence microplate reader, using excitation and emission wavelengths of 365 nm and 455 nm, respectively.

**2.2.5.13 4-MUCT assay (Chitin-hydrolysis assay)**

The 4-MUCT assay was performed as described in Brunner et al. (Brunner et al., 1998). In brief, the hydrolytic activity towards the substrate 4-methylumbelliferyl-β-D- N, N', N'' triacetylchitotriose (4-MUCT, Sigma) was measured and compared to that of 2 µg Streptomyces griseus chitinase (Sigma). After enzyme incubation in 250 µl final volume of 0.05 % (w/v) 4-MUCT in 20 mM sodium acetate, pH 5.2 at 37°C, 20 µl of the reaction mixture were removed and added to 980 µl 0.2 M sodium carbonate solution. Free 4-MU (Sigma) was used for the generation of a standard curve. The intensity of the 4-MU fluorescence in the samples was monitored with an MWGt Sirius HT fluorescence microplate reader (absorbance at 360 nm and emission at 450 nm). The same protein amounts were used as for the turbidity assay (see above).

**2.2.5.14 Zymography**

A discontinuous CTAB polyacrylamid gel electrophoresis was performed using a 12% separating gel (43 mM KOH, 280 mM acetic acid, pH4.0, 12% acrylamide bisacrylamide
Materials and Methods

37.5:1, 8% glycerol, 1.3% ammonium persulphate and 0.16% TEMED), overlaid by a 4% stacking gel (64 mM KOH; 94 mM acetic acid, pH5.1, 4% acrylamide, 1.25% ammonium persulphate and 0.125 mM TEMED). Prior to loading, the gel was pre-run using anode buffer (40 mM beta-alanine, 70 mM acetic acid, 0.1% CTAB, pH4.0) and cathode buffer (50 mM KOH, 56 mM acetic acid, pH5.7, 0.1% CTAB) for 1 hour at 250 volt. Crude protein extracts were mixed with an equal volume of loading buffer (5 M urea, 25 mM KAc pH6.8, Methylene blue) and separated for 2 hours at 150 volt and 4°C. After electrophoresis, the CTAB gel was washed with 20 mM NaAc and then sprayed with 0.00625% 4-MUCT in 20 mM NaAc, pH5.2 and incubated at 37°C for 30 minutes. Fluorescent bands were documented under UV light using the infinit-3026 WL/26MX gel imaging system (PeqLab, Erlangen, Germany).

2.2.5.15 HPLC analysis of PGN fragments

500 µg/ml B.subtilis PGN was incubated with 140 µg LYS1 purified from LYS1OE plants or controls in 20 mM NaAc pH5.2 at 37°C with shaking for the indicated time. After stopping the reaction by heating at 100°C for 10 minutes, the reaction was centrifuged and the supernatant analysed by HPLC. The analyses were done by CeCo labs on an Agilent 1200 system with a Prontosil C18-RP column (Bischoff Chromatography, Leonberg, Germany).

2.2.6 Bioassays

2.2.6.1 Infection with Pseudomonas syringae

For the bacterial infection assay, an overnight culture of Pseudomonas syringae pv. tomato DC3000 was harvested by centrifugation, washed once with 10 mM MgCl2 and finally diluted with 10 mM MgCl2 to a density of 1 x 10^4 cfu/ml (OD600 ~2 x 10^-5) and was then infiltrated with a 1ml-needleless syringe into the leaf apoplast. Two leaves per plant and 8 plants were infected per plant genotype. The growth of bacteria was determined after 0 and 2 days post infection. For the quantification, infected leaves were harvested (2 leaves at 0 dpi and 3 leaves at 2 dpi) and washed for one minute in both 70 % (v/v) EtOH and water. Afterwards 2 leaf discs per leaf with a diameter of 5 mm were cut out and homogenized in 200 µl 10 mM MgCl2. 10 µl of each homogenate were
then plated undiluted and in different dilutions onto LB agar plates and incubated at 28°C for 24-48 hours. The growth of bacteria was determined by colony counting, and subsequently mean values and standard deviations were determined.

2.2.6.2 Elicitation assays in leaves or seedlings

Leaves of 4-6 week old plants were infiltrated using a needle-less syringe with solutions of PAMPs and harvested after indicated time points. For the seedling elicitations, seedlings were first cultivated on sterile ½ MS plates for 5-6 days in long-day. Then they were transferred into liquid MS medium supplemented with 1 % (w/v) saccharose (4-6 seedlings in 200 µl medium/well, 48-er well plate) and equilibrated overnight. After addition of the PAMPs, the seedlings were incubated with gentle shaking and harvested at indicated time points.

The PAMPs were used in elicitation assays in the following concentrations: 1 μM flg22 and 100 μg/ml chitin or PGN.

2.2.6.3 Medium alkalization assay

Medium alkalization in cell culture upon PAMP treatment was performed as described previously (Gust et al., 2007). In brief, 300µl cultured cells were transferred to 48-well plates and equilibrated at 150 rpm for 30 minutes. After addition of PAMPs, the pH in the cell culture was monitored with an in Lab Microelectrode (Mettler Toledo, Gießen, Germany). The changes in pH were monitored and recorded by the Observer II program (Brainchild Electronics Co., Ltd, Taipei, Taiwan).

2.2.6.4 Oxidative burst assay

Leaf discs were excised from 6 week-old Arabidopsis plants and incubated in water overnight. The following day, the discs were transferred to a solution of 20 μM luminol L-012 (Wako Pure Chemical Industries) in a 96-well plate supplemented with PAMPs. The plates were analyzed for a period of at least 30 min using a multiplate reader Centro LB 900 (Berthold Technologies). For each data point, at least 6 replicates were measured.
2.2.7 Microscopy

The visualization of fluorescence in samples was done using confocal laser scanning microscopy (TCS SP2, Leica). The images were taken using the 63x/1,2 Plan Apo H₂O objectives. The Software LCS Lite Version 2.61 was used for the processing of the images.

Statistical analysis

Statistical analysis was performed using Microsoft Office Excel. The data represent the average of replicates with standard deviation (SD) or standard error (SE). Statistical significance between two groups has been checked by using a two-tailed unpaired Student’s t test. For multiple comparisons, the one-way ANOVA method was performed combined with the Tukey’s honest significant difference (HSD) test. Significant differences are indicated with different letters (p < 0.01). Asterisks represent significant differences (*p < 0.05; **p < 0.01; ***p < 0.001).
3. Results

3.1 Identification of an *Arabidopsis* PGN hydrolase

3.1.1 An induced PGN-degrading activity in *Arabidopsis* is not due to PGN receptors LYM1 and LYM3

Both, complex insoluble PGN fractions prepared from bacterial cell walls as well as mixtures of soluble oligomeric PGN fragments have previously been shown to stimulate plant immune responses in *Arabidopsis* (Gust et al., 2007; Erbs et al., 2008; Willmann et al., 2011), suggesting that partially hydrolyzed ligands could potentially serve PRR-mediated immune activation. Therefore we aimed to identify such PGN-hydrolase activity in *Arabidopsis*. Plant genomes do not encode lysozyme-like proteins, but many plant species engage in lysozyme-like activities under induced conditions (Brunner et al., 1998; van Loon et al., 2006). In *Arabidopsis*, we tested if lysozyme-like activities can also be induced by bacteria-derived PAMPs. To do this, a standard lysozyme assay (Park et al., 2002) was employed to determine PGN-degrading activity based on turbidity reduction in suspensions of Gram-positive *Micrococcus luteus* cell wall preparations. As shown in Figure 1, significant cell wall-degrading activities were detected in the extracts from leaves infiltrated with both Flg22 and PGN, compared with the water infiltration, indicating that these PAMPs induce *Arabidopsis* leaves to generate lysozyme-like lytic activities.
Figure 1. Bacterial PAMPs induce lysozyme-like activities in *Arabidopsis*.

*Micrococcus luteus* cell wall preparations were incubated with 40 µg protein crude extracts from *Arabidopsis* leaves 6 hours post infiltration with water, 1 µM Flg22 or 100 µg/ml PGN. PGN hydrolytic activity was assayed in a turbidity assay at the indicated time points. *M.luteus* cells were incubated together with the crude extracts and the turbidity in absorbance at 570 nm of the suspension was measured with a spectrophotometer over time. As positive control, *Micrococcus luteus* cell wall preparations were incubated with 0.5 µg/ml hen egg white lysozyme. Means ± SD of three replicates per sample are given. Statistical significance compared with the water infiltration (***(p<0.0001, Student’s t test) is indicated by asterisks.

In metazoan, some PGN recognition proteins (PGRPs) harbor PGN lytic activities (Gelius et al., 2003; Wang et al., 2003). Here, we investigated whether LYM1 and LYM3, the two *Arabidopsis* PGN receptors (Willmann et al., 2011), were also able to catalyze PGN degradation in a standard lysozyme assay. As shown in Figure 2, neither recombinantly expressed and purified LYM1 nor LYM3 displayed cell wall-degrading
activity. Thus we propose that the induced lysozyme activities in *Arabidopsis* result from some other, yet unknown PAMP-induced enzyme(s).

![Figure 2](image)

**Figure 2. LYM1 and LYM3 do not possess PGN hydrolytic activity.**

*Micrococcus luteus* cell wall preparations were incubated with 20 μg affinity-purified His6-tagged LYM1 or LYM3 or 0.5 μg hen egg-white lysozyme and PGN hydrolytic activity was assayed in a turbidity assay at the indicated time points. As negative control (nc), non-induced His6-tagged LYM3 bacterial lysates were used for affinity purification and elutes were subjected to turbidity assays. Means ± SD of three replicates per sample are given. Statistical significance compared with the negative control (**p<0.001, ***p<0.0001, Student’s t test) is indicated by asterisks. The data for this figure were kindly provided by Roland Willmann.

### 3.1.2 Identification of LYS1 as a potential PGN hydrolase

Lysozymes (EC 3.2.1.17) hydrolyze β (1, 4) linkages between GlcNAc and MurNAc residues in PGNs and between GlcNAc residues in chitodextrins (enzyme.expasy.org). Although lysozyme-like sequences were not found in plant genomes, some plant chitinases were reported to display lysozyme-like activities (Audy et al., 1988; Sakthivel et al., 2010). For example, hevamine, one class III chitinase from the rubber tree has been shown to harbor PGN hydrolysis activity (Bokma et al., 1997). Similarly, two
tobacco class III chitinases, lysb1 and lysb2 were also found to harbor lysozyme activity (Brunner et al., 1998).

Figure 3. Protein sequence alignment of the 24 *Arabidopsis* chitinases and cartoon diagram of the 3D structure of LYS1. (A) Full length amino acid sequences were aligned with the ClustalW2 algorithm and subgroups were classified according to their sequences and structures (Passarinho and de Vries, 2002). *Arabidopsis* lysozyme 1 (LYS1, At5g24090, formerly also named CHIA) represents the only member of class III. (B) Full length amino acid sequences of LYS1 (At5g24090) was predicted using Phyre sever (Kelley and Sternberg, 2009) and a ribbon diagram was created with the Pymol program. A typical (βα)8 barrel fold is shown in the image. N and C indicate N-terminus and C-terminus of LYS1, respectively.

Chitinases (EC 3.2.1.14) hydrolyze β (1, 4) linkages between GlcNAc in chitin and chitodextrins (enzyme.expasy.org). *Arabidopsis* chitinases fall into five groups (Figure 3A) (Passarinho and de Vries, 2002), and are grouped into structurally unrelated families 18 and 19 of glycosyl hydrolases (Henrissat, 1991), respectively. As mentioned above, the class III chitinases (glycosyl hydrolase family 18) from rubber tree and tobacco displayed bifunctional activities: chitinase and lysozyme activity. Of the 24 annotated members of *Arabidopsis* chitinases, LYS1 (At5g24090, formerly also named CHIA) is the only member of class III (Figure 3A). Alignment of LYS1 with rubber tree hevamine resulted in approximately 70% identity (Grabherr, 2011). LYS1 was predicted to have a
typical (βα)₈ barrel fold (Figure 3B). All these information make LYS a perfect PGN hydrolase candidate. Thus, we chose LYS1 as a putative lysozyme candidate for further studies.

3.1.3 Expression of epitope-tagged LYS1 and identification of active LYS1

In order to analyze the enzymatic properties of LYS1 in vitro, heterologous expression of LYS1 was attempted. Overexpression in E.coli failed to produce active enzyme and LYS1 production in eukaryotic Pichia pastoris entirely failed to produce recombinant protein (Grabherr, 2011). Thus, LYS1 overexpression (LYS₁⁰ᴱ) lines carrying a p35S::LYS₁-GFP cassette were created in the Col-0 background using the floral dipping method (Grabherr, 2011). We first examined LYS1 protein levels in LYS₁⁰ᴱ lines using two different antibodies for Western blotting analysis. The first antibody was anti-GFP which could be used to detect the fused LYS1-GFP in the LYS₁⁰ᴱ plants. The second antibody, which was raised in rabbit against the tobacco class III chitinases (kindly provided by Dr. Frédéric Brunner), could be tested for recognition of the native LYS1 and the LYS1-GFP fusion. The α-GFP antibody detected specific bands in leaf extracts of LYS₁⁰ᴱ lines with sizes of approximately 60 kDa and 30 kDa (Figure 4, right panel), which correlated with the expected sizes of the LYS1-GFP fusion (60.1 kDa) and free GFP (27 kDa).

The detection of LYS1 using the α-class III chitinase antibody revealed a more complex pattern of protein bands. In the LYS₁⁰ᴱ lane a clear band appeared below the 35 kDa marker band (Figure 4, left panel). This protein band possibly represents the free LYS1 protein, which has the calculated size of 33.1 kDa without the GFP tag. A weak band in the size of the LYS1-GFP fusion protein at about 60 kDa was also present in the LYS₁⁰ᴱ lane. It is possible that the large GFP tag interferes with the recognition of the LYS1 protein by the α-class III chitinase antibody, leading to weaker detection of the GFP-tagged LYS1. Alternatively, a major amount of LYS1 in the LYS₁⁰ᴱ line is not present as GFP-fusion but as free LYS1 (Figure 4, left panel). In the same immunoblot analysis we also included an amiRNA knockdown line, which will be introduced in chapter 2.4.1. Here, the two antibodies detected no band either in the WT samples or in the LYS₁¹ᴷ⁰
lanes, indicating undetectable levels of native LYS1 in the WT and absence of any GFP in both plant types (Figure 4).

![Image](image.png)

**Figure 4. Analysis of LYS1 protein levels in LYS1<sup>OE</sup> lines.** Immunoblot analysis of protein extracts from leaves of two independent LYS1<sup>OE</sup> lines, a LYS1 knock-down line (LYS1<sup>KD-1</sup>, see chapter 3.4.1) and wild-type plants (WT). Total leaf protein was separated by SDS-PAGE and blotted onto a nitrocellulose membrane. Immunodetection was carried out using antibodies raised against α-tobacco class III chitinase (α-Chit) or green fluorescent protein (α-GFP). Ponceau S red staining of the large subunit of RuBisCO served as loading control.

To analyze LYS1 enzymatic properties, we resorted to isolating active LYS1 from plant hosts including the stable LYS1 expressor-<i>Arabidopsis</i> lines LYS1<sup>OE</sup> and transient LYS1 expression in <i>N. benthamiana</i>. At first an immunoprecipitation (IP) approach was exploited to enrich and purify LYS1 from LYS1<sup>OE</sup> plants. Protein-G agarose beads were coupled with the anti-GFP antibody and then used to pull down LYS1-GFP from protein extracts of LYS1<sup>OE</sup> plants (Figure 5A). Protein G-isolated LYS1 was subjected to a PGN hydrolysis assay (Figure 5B) and immunoblotting (Figure 5C). As shown in Figure 5C, western blot analysis showed a clear band of LYS1-GFP in the lane with the Protein-G-bound sample from LYS1<sup>OE</sup> plants, indicating successful isolation and enrichment of LYS1-GFP by this IP. However, the PGN activity assay revealed that the immunopurified LYS1-GFP protein did not display any PGN hydrolytic activity in comparison to respective samples derived from the wild type control (Figure 5B). If we compare PGN
hydrolytic activities of fractions shown in Figure 5B with their corresponding Western blot band intensities shown in Figure 5C, obviously LYS1-GFP amounts did not correlate with PGN hydrolytic activity.

Figure 5. Isolation and analysis of LYS1-GFP via immunoprecipitation (IP). (A) Schematic drawing of purification steps. Protein-G beads were incubated with crude protein extracts of LYS1OE (oe) or wild type (wt) plants, and washed afterwards four times with the indicated buffers (P.E., protein extraction buffer), collecting the washing supernatant (ws) after each step (0-4). Bound proteins were collected and used for a PGN hydrolysis assay (B) and Western blotting analysis (C). (B) Bacillus subtilis PGN suspension was incubated with the fractions indicated in (A). Relative PGN hydrolytic activities were calculated after 2 hours incubation using hen egg-white lysozyme as standard (set to 1). Shown are the means ± SD of 3 replicates. (C) Fractions indicated in (A) were separated by SDS-PAGE and blotted onto a nitrocellulose membrane. Immunodetection was carried out using an anti-GFP antibody. Arrowheads indicate positions of LYS1-GFP and GFP.
We thus assume that the IP-isolated LYS1-GFP does not have any lysozyme-like activity, irrespective of higher PGN hydrolytic activities measured in the \( \text{LYS}^{\text{OE}} \) input samples expressing \( p35S::\text{LYS1-GFP} \) than in wild type samples (Figure 5B).

To further confirm our assumption that LYS1-GFP does not harbor lysozyme-like activity, a CTAB-PAGE zymography assay was performed to analyze the protein extracts derived from \( \text{LYS}^{\text{OE}} \) plants. In a conventional Laemmli SDS-PAGE (Laemmli, 1970) in which proteins are separated based on their sizes, SDS often destroys the protein activities. In contrast, in a CTAB-PAGE protein activities can be retained while proteins are also separated based on their sizes (Akins et al., 1992). The CTAB-PAGE system was therefore employed here to identify the responsible protein for lysozyme-like activity in \( \text{LYS}^{\text{OE}} \) lines. Following some initial experiments with various buffer systems based on \textit{in silico} design (Jovin, 1973), an acidic buffer system was finally chosen as the only one that yielded LYS1 activity in a following zymography (see details in the method section).

As shown in Figure 6A, the zymography results revealed that in the LYS1-GFP sample fluorescent bands appeared at the expected position of LYS1 (33.1 kDa) but not at the expected size of LYS1-GFP (60.1 kDa) (Figure 6A, left panel), although a parallel immunoblot detection showed visible bands for LYS1-GFP in the LYS1-GFP sample (Figure 6A, right panel), suggesting that free LYS1 but not the LYS1-GFP fusion protein harbors enzyme activity. As a negative control, protein extracts from plants stably expressing secreted GFP (secGFP) (Teh and Moore, 2007) were included. The result is in agreement with the one shown in Figure 5, where IP-isolated LYS1-GFP was also not responsible for PGN hydrolytic activity (Figure 5B). Likewise, CTAB-PAGE-zymography analysis of transiently expressed LYS1-GFP from \textit{N. benthamiana} revealed that LYS1-GFP did not display hydrolytic activity (Figure 6B).
Figure 6. CTAB-PAGE zymography and immunoblot analysis of extracts of \textit{LYS}^\text{OE} \textit{plants} and \textit{N. benthamiana} transiently expressing \textit{LYS1-GFP}. (A) Protein extracts from \textit{LYS}^\text{OE}-1 or \textit{secGFP} plants were separated on a cetyltrimethylammonium bromide (CTAB)–polyacrylamide (PA) gel and hydrolytic activity was assayed in a zymogram by overlaying the gel with the substrate 4-MUCT. BSA served as negative control. Fluorescent bands are indicative of substrate cleavage (left panel). The proteins were blotted from the CTAB gel onto a nitrocellulose membrane and detected using an anti-GFP antibody (right panel). Arrowheads indicate the positions of \textit{LYS1-GFP}, \textit{LYS1} and GFP. (B) Protein extracts from \textit{N. benthamiana} leaves transiently expressing \textit{p35S::LYS1-GFP}, \textit{p35S::del-LYS1-GFP} (deletion of signal peptide) or p19 were separated on a CTAB-PA gel, and hydrolytic activity (left panel) or protein expression using an anti-GFP Western blot (right panel) were analysed as described in (A). Indicated by arrowheads are the positions of the \textit{LYS1-GFP} fusion and free \textit{LYS1} or GFP and of an unspecific background band appearing due to \textit{Agrobacterium} infiltration, thus also present in the p19 negative control.

Here a construct expressing \textit{LYS1-GFP} without the \textit{LYS1} secretion peptide (del-LYS1-GFP) (Grabherr, 2011) was included. Notably, no activity was detected at the expected position of \textit{LYS1} (33.1 kDa) in the del-LYS1-GFP sample (Figure 6B, left panel), although del-LYS1-GFP was detected in immunoblotting analysis (Figure 6B, right panel), indicating that proper secretion is required for \textit{LYS1} activity. We thus assume
that LYS1 undergoes some posttranslational processing to generate an active form in a secretion pathway.

Figure 7. Transiently expressed LYS1 is a glucan hydrolase. (A) Protein extracts from *N. benthamiana* leaves expressing *p35S::LYS1* constructs containing different epitope tags (*LYS1_myc*, *LYS1_HA* or *LYS1_GFP*) were separated on a SDS-polyacrylamid gel and analysed by western blot using antibodies raised against the myc-, HA- or GFP-epitope tags. As control, plants were infiltrated with agrobacteria harboring the p19 suppressor of silencing construct (p19). Protein sizes (kDa) are indicated on the left. (B) *N. benthamiana* protein extracts expressing LYS1 (*LYS1_myc*, *LYS1_HA* or *LYS1_GFP*) or as control extracts from non-infiltrated (ni) or p19-infiltrated leaves were separated on a CTAB-polyacrylamid gel and hydrolytic activity was assayed by overlaying the gel with the substrate 4-MUCT. Fluorescent bands are indicative of substrate
cleavage. Arrowheads indicate the positions of epitope-tagged LYS1, an unspecific band in all samples infiltrated with agrobacteria is labelled with an asterisk.

We assumed that the large GFP tag in the LYS1-GFP fusion protein disturbed LYS1 activity while small tags might not. To address this question, p35S::LYS1 with small and large epitope tags (myc, HA, GFP) were transiently expressed in N. benthamiana. Western blotting analysis showed that the three proteins were successfully expressed (Figure 7A), and a following CTAB-PAGE zymography revealed that all three extracts contain the active LYS1 at the expected position (Figure 7B). However, as myc and HA are very small epitope-tags and as the CTAB-PAGE has a poor resolution, it cannot be distinguished whether 4-MUCT-hydrolytic activity is derived from epitope-tagged LYS1 or the cleaved, native LYS1. Nevertheless, it was also observed that all three extracts displayed lysozyme-like activity towards 4-MUCT, M. luteus cells and B. subtilis PGN as shown in Figure 8A-C. Notably, the PGN hydrolytic activity of extracts expressing LYS1myc was the highest at about pH 5-6 (Figure 8D), which is approximately the same pH optimum observed for extracts from LYS1OE plants (Grabherr, 2011; Liu et al., 2014).
Figure 8. Analysis of the activity of transiently expressed LYS1. Protein extracts from *N. benthamiana* leaves expressing p35S::LYS1 constructs containing different epitope tags (LYS1<sub>GFP</sub>, LYS1<sub>HA</sub> or LYS1<sub>myc</sub>) were assayed for chitinolytic activity with 4-MUCT substrate (A) or for PGN hydrolytic activity in a turbidity assay using *M. luteus* cells (B) or *Bacillus subtilis* PGN (C). Relative activities (2 hours incubation) were calculated using *Streptomyces griseus* chitinase (A) or hen egg-white lysozyme (B-D) as standards (set to 1). As control, plants were left untreated (c) or infiltrated with agrobacteria harboring the p19 suppressor of silencing construct (p19). (D) Lysis of *M. luteus* cells was determined in a turbidity assay with LYS1<sub>myc</sub> leaf protein extracts as described in (B) at the indicated pH. Means ± SD of two replicates per sample are given. Significant differences in enzyme activities relative to those in the p19 control are indicated (*, p ≤ 0.05; Student’s t-test).
Next, the active LYS1 in the extracts expressing \( LYS1_{myc} \) was separated by CTAB-PAGE, followed by pull-down and subsequent SDS-PAGE, and then analyzed by nano-LC MS/MS, which confirmed the identity of LYS1 (Figure 9). However, in the following PGN hydrolytic activity assay, IP-isolated LYS1-HA and LYS1-myc fusion proteins did not display enriched PGN-degrading activities either (Figure 10).

**Figure 9.** Isolation and identification of active LYS1 transiently expressed in *N. benthamiana*. (A) Total *N. benthamiana* protein extracts expressing LYS1\(_{myc}\) were prepared (Total), separated on a CTAB-gel and hydrolytic activity was assayed by overlaying the gel with the substrate 4-MUCT (middle panel). Fluorescent bands were excised, and the eluted proteins were precipitated with insoluble chitin and separated by SDS-PA gel electrophoresis followed by Coomassie Brilliant Blue staining (right panel). A
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flow chart illustrates the purification steps (left panel). (B) The band visible at approximately 35 kDa in (A) (arrow) was excised and the eluted proteins were subjected to nano-LC MS/MS analysis. Shown is a summary of protein hits obtained by MS/MS analysis. Only hits with a peptide number ≥ 2 and a Mascot score sum ≥ 37 were retained. Identification number (ID) from the tobacco proteome database and molecular weight (MW) of the identified proteins are indicated.

From these results we reasoned that all tested tags (GFP, HA or myc) were interfering with LYS1 enzymatic activity and the only active form is the untagged LYS1 derived from overexpression of epitope-tagged LYS1 in both Arabidopsis and N. benthamiana. As the LC-MS/MS analysis of the enzymatically active band shown in Figure 9 did not yield any chitinase-like enzymes in the cut-out band (Figure 9B), we could, however, rule out that a N. benthamiana protein is responsible for the observed 4-MUCT-cleavage.

Figure 10. Isolation and analysis of LYS1-myc and LYS1-HA via immunoprecipitation (IP). (A) Schematic drawing of purification steps. Protein-G beads were incubated with crude protein extracts from N. benthamiana leaves expressing p35S::LYS1-HA, p35S::LYS1-myc or control leaves which were infiltrated with agrobacteria harboring the p19 suppressor of silencing construct (p19), and afterwards washed four times with the indicated buffers (P.E., protein extraction buffer), collecting the washing
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supernatant (ws) after each step (0-4). (B) Proteins bound to Protein-G beads were collected and used for a PGN hydrolysis assay. *B. subtilis* PGN suspension was incubated with the fractions indicated in (A). Relative PGN hydrolytic activities were calculated after 2 hours incubation using hen egg-white lysozyme as standard (set to 1). Means ± SD of 3 replicates per sample are given.

### 3.1.4 Purification of active LYS1 using FPLC

Since only the active LYS1 form seems to be the free LYS1 in *Arabidopsis* and *N. benthamiana*, and additional unspecific hydrolytically active proteins were induced upon *Agrobacterium*-infiltration in *N. benthamiana* (Figure 7B and 8), we next aimed at isolating the free LYS1 directly from LYS1OE leaf extracts. Based on some properties of the tested LYS1, we developed a method based on fast protein liquid chromatography (FPLC) to isolate and purify the active, untagged LYS1. Firstly, the isoelectric point of LYS1 was predicted to be 9.3 (arabidopsis.org), hence a cation-exchange chromatography was selected to separate LYS1 as a basic protein. Secondly, the optimum pH for LYS1 enzyme activity is between pH 5 and 6 (Figure 8D), and thus an acidic NaAc pH 5.2 buffer was used as FPLC buffer (Figure 11A). As shown in Figure 11, the collected fractions from the FPLC were validated for their activities by a 4-MUCT assay. The results revealed that fractions 41-45 displayed strong fluorescent signals, indicative of cleavage of the 4-MUCT substrate (Figure 11B, left panel). Further SDS-PAGE analysis with subsequent Coomassie Brilliant Blue staining of these fractions confirmed a band at around 35 kDa, the expected size of native LYS1 (Figure 11B, right panel). This 35 kDa-band was excised from the gel for subsequent nano-LC MS/MS analysis. The peptide mass fingerprint not only confirmed the identity of LYS1 in this band, but also yielded peptides spanning the whole protein sequence, except for the first 53 amino acids which are predicted to present a signal peptide (Figure 11C).
Figure 11. FPLC purification of active LYS1 from LYS1\textsuperscript{OE}. Cleaved untagged LYS1 was isolated via FPLC-purification (see materials and methods) from leaf tissue of transgenic LYS1-overexpressing (LYS1\textsuperscript{OE}) Arabidopsis lines. (A) Flowchart of purification of LYS1. (B) FPLC elution fractions 36 - 48 were assayed for enzymatic activity using the 4-MUCT assay (left panel) and the presence of LYS1 protein was confirmed by SDS-PAGE and Coomassie Brilliant Blue staining (CBB, right panel). (C) Fraction 42 and 43 of the FPLC elution were separated on a SDS-PAGE and stained with Coomassie Brilliant Blue (left panel). The protein band at the expected size of LYS1 was excised and subjected to nano-LC MS/MS analysis. The MS identified peptides (highlighted in light grey letters in the sequence and indicated by grey bars below the sequence) were matched to the LYS1 amino acid sequence (right panel).

3.2 Characterization of LYS1

3.2.1 LYS1 is a bifunctional enzyme with lysozyme- and chitinase-activity

FPLC-purified LYS1 was tested for its chitinolytic activity in a 4-MUCT assay. As shown in Figure 12A, compared with the control (respective purification with WT protein extract), purified samples from LYS1\textsuperscript{OE} extracts exhibited significant cleavage activities towards
4-MUCT. Thus, LYS1 indeed has the expected chitinase activity. Next, LYS1 was tested for its ability to solubilize complex PGN presented by intact Gram-positive *M. luteus* cells and to cleave preparations of complex, insoluble *B. subtilis* PGN. Again, purified samples from LYS1\(^{OE}\) extracts exhibited significant cell-decomposing (Figure 12B) and PGN-degrading activities (Figure 12C) in comparison to the WT control.

Figure 12. *Purified LYS1 from LYS1\(^{OE}\) has glucan-hydrolase activity.* FPLC purified LYS1 protein (LYS1) from LYS1\(^{OE}\) protein extracts and, as a control, a comparable FPLC-purified fraction from wild type leaf material (WT) were assayed for hydrolytic activity towards glycan substrates. (A) Assay for chitinolytic activity using the 4-MUCT substrate. Relative activities at 1 and 3 hours post treatment were calculated using *Streptomyces griseus* chitinase as standard. *S. griseus* chitinase activity was set to 1. (B, C) *Micrococcus luteus* cells (B) or *Bacillus subtilis* PGN (C) were subjected to hydrolysis by FPLC-purified LYS1 and hydrolytic activity was calculated at 1 and 3 hours after treatment using hen egg-white lysozyme as standard (set to 1). Significant differences compared with the buffer control (buffer) are indicated by asterisks (*p<0.05; Student's t test).

In addition, we also tested if LYS1 might have a potential cellulase activity. As shown in Figure 13, LYS1 did not display any cellulase activity compared to a commercial
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cellulase. Hence, we conclude that LYS1, formerly identified as a chitinase, harbors a bifunctional lysozyme/chitinase activity.

Figure 13. LYS1 is devoid of cellulose hydrolytic activity. LYS1 was purified from 5-week-old LYS1OE plants and used for cellulase activity assays. The substrate 4-methylumbelliferyl-β-D-cellobioside was incubated for 1 hr with purified LYS1, commercial reference cellulose, or buffer as control. Fluorescence was determined (ex/em = 365 nm/455 nm) after stopping the reaction with 0.2 M sodium carbonate. Means ± SD of three replicates per sample are given. Statistical significance compared with the buffer control (***p<0.001, Student’s t test) is indicated by asterisks.

3.2.2 Enzyme Kinetics of LYS1

To determine specific enzyme activities, untagged LYS1 was purified from LYS1OE Arabidopsis lines by FPLC and used for enzyme assays. In a 4-MUCT assay LYS1 yielded a K_m of 70 ± 14 μM and a V_max of 378 ± 42 μM min^{-1} mg^{-1} for LYS1, and a K_m of 53 ± 27 μM and a V_max of 397 ± 145 μM min^{-1} mg^{-1} for commercial S. griseus chitinase. In a turbidity assay with M. luteus cell wall preparations, a K_m of 18.2 ± 2.5 mg/ml and V_max of 4.4 ± 0.6 mg mg^{-1} min^{-1} were obtained for LYS1, and a K_m of 8.4 ± 0.8 mg/ml and V_max of 192 ± 120 mg mg^{-1} min^{-1} for commercial hen egg white lysozyme. Thus, the K_m values for LYS1 are comparable to the commercial enzymes.
3.2.3 LYS1 localizes to the apoplast

**Figure 14. LYS1 is localized to the apoplast.** Apoplastic washes were prepared from leaves of wild-type *Arabidopsis* plants or the *LYS1*<sup>OE</sup>-1 and *LYS1*<sup>KD</sup>-1 lines. Apoplastic fluids (concentrated tenfold) or total leaf protein extracts were subjected to western blot analysis using antibodies raised against green fluorescent protein (α-GFP), tobacco class III chitinase (α-chit), or the cytoplasmic mitogen-activated protein kinase 3 (MPK3).

LYS1 is predicted to contain a signal peptide and to be secreted into the plant apoplast. To confirm this localization of LYS1, we prepared apoplastic washes from *LYS1*<sup>OE</sup> *Arabidopsis* lines. Both the LYS1-GFP fusion protein as well as free LYS1 were detectable in concentrated apoplastic fluids whereas the cytoplasmic mitogen-activated protein kinase MPK3 was only present in the total leaf protein samples (Figure 14). Moreover, previous identification within the *Arabidopsis* cell wall proteome (Kwon et al., 2005) suggests that LYS1 acts in the plant apoplast. Furthermore, Heini Grabherr could demonstrate in her thesis that, in preparations of protoplasts from the *LYS1*<sup>OE</sup> line, increased hydrolytic activity as compared to the wild-type was mostly found in the
protoplast medium, again indicating that LYS1 has been secreted by the protoplasts (Grabherr, 2011; Liu et al., 2014).

3.3 Role of LYS1 in the generation of immunogenic PGN fragments

3.3.1 LYS1 generates plant immunogenic PGN fragments

As LYS1 was shown to harbor PGN hydrolytic activity, we subsequently wanted to analyze immunogenic activities of PGN cleavage products generated by LYS1. Untagged LYS1 was purified from LYS1OE Arabidopsis lines by FPLC and used for degradation of B. subtilis PGN. Solubilized PGN fragments found in the supernatant of LYS1-digested PGN were subsequently analyzed by high performance liquid chromatography (Figure 15A). Few peaks could be detected in the supernatant of PGN incubated with a buffer control or with heat-inactivated LYS1. In contrast, PGN-digests produced by native FPLC-purified LYS1 (see 3.1.4) yielded several characteristic peaks that were also detectable in the supernatants of PGN preparations treated with mutanolysin, which has been shown to cleave O-glycosidic bonds between GlcNAc and MurNAc residues in complex PGN (Yokogawa et al., 1975)
Results

Figure 15. Purified LYS1 generates immunogenic PGN fragments. LYS1 was purified from 5 weeks old *LYS1*OE plants and used for PGN digestion. (A) 500 µg of *Bacillus subtilis* PGN were digested for 7 hours with either mutanolysin (50 µg/ml), native purified LYS1 (140 µg/ml), heat-denatured purified LYS1 (140 µg/ml) or the reaction buffer alone and subjected to HPLC fractionation. Shown are the peak profiles of representative runs. The signal intensity is given in milli absorbance units (mAU). (B) *B. subtilis* PGN
was digested for 4 h as described in (A) and Arabidopsis wild type seedlings or the indicated mutant lines were treated for 6 h with 25 µl/ml digest supernatant containing solubilized PGN fragments. Total seedling RNA was subjected to RT-qPCR using Flagellin responsive kinase (FRK1) specific primers. EF1α transcript was used for normalization; water treatment served as control and was set to 1. (C) Supernatants of digested PGN (25 µl/ml) were added to cultured rice cells and medium alkanilization was determined at 20 min post addition. Treatment with water or MES buffer served as control. All data represent triplicate samples ± SD, and bars with different letters are significantly different based on one-way ANOVA (p < 0.05; B, C). (D) B. subtilis PGN was digested with native purified LYS1 for the indicated times or overnight (o/n), and digest supernatant was used to trigger medium alkanilization in rice cells as described in (C). All data represent triplicate samples ± SD; asterisks indicate significant differences compared to the buffer control (* p < 0.05; ** p < 0.01; *** p < 0.001; Student’s t test).

LYS1-generated PGN-fragments were subsequently tested for their abilities to trigger plant immunity-associated responses (Figures 15B-C). Firstly, supernatants of PGN preparations treated with either native or heat-denatured LYS1 were used to trigger immune marker gene FRK1 expression in Arabidopsis seedlings. Importantly, only supernatants from PGN-digests produced by native LYS1 or mutanolysin induced FRK1 expression whereas buffer controls or digests produced by heat-inactivated LYS1 did not release immunogenic soluble fragments from complex PGN (Figure 15B). Notably, activation of immune responses by LYS1-generated PGN-fragments was dependent on Arabidopsis PGN receptor complex components LYM1, LYM3 and CERK1 (Willmann et al., 2011) as the respective mutant genotypes failed to respond to immunogenic PGN fragments (Figure 15B). Secondly, we tested whether LYS1-generated PGN fragments were able to trigger an immunity-associated response, medium alkanilization, in rice cell suspensions. This plant was chosen for testing as a PGN receptor system, because it has recently been reported to be very similar to that in Arabidopsis (Liu et al., 2012a) and because rice cells have been observed to be much more sensitive to PGN than Arabidopsis cells (Roland Willmann, personal communication). As shown in Figure 15C, LYS1-released PGN-fragments triggered medium alkanilization in cultured rice cells, suggesting that immune defence stimulation by soluble PGN fragments is not restricted to Arabidopsis only.

3.3.2 LYS1-overdigested PGN induces weaker immunity responses
We further investigated the kinetics of PGN fragment release from complex PGN. As shown in Figure 15D, release of immunogenic PGN-fragments into solution occurred rapidly within 10 min of incubation with native LYS1. Incubation of complex PGN with LYS1 yielded the highest immunogenic activity of the digested supernatant after 30 min, suggesting that at that time point the maximum amount of immunogenic PGN fragments was generated. However, prolonged incubation with LYS1 again resulted in a loss of activity with overnight digestion completely abolishing stimulatory activity of the PGN digest. We assume that LYS1 is capable of releasing immunogenic fragments from complex PGN, but extensive or complete digest into PGN-monomers or small PGN fragments appears to abolish the immunogenic activity of PGN fragments. This result is in accordance with previous observations that prolonged digestion of PGN with mutanolysin diminishes its defence-inducing activity (Gust et al., 2007).

3.4 LYS1 is required for immune responses to PGN

3.4.1 Characterization of \( LYS1^{KD} \) lines

To examine a role of LYS1 in plant immunity, attempts were undertaken to genetically inactivate \( LYS1 \) expression. Three independent \( LYS1 \) T-DNA insert lines were obtained from the Nottingham Arabidopsis Stock Center (NASC) and the Cold Spring Harbor Laboratory (CHSL), but transcription analysis of the three lines revealed similar \( LYS1 \) transcript levels in these mutants to those in the corresponding wild types (Grabherr, 2011). Alternatively, transgenic \( LYS1 \) knockdown (\( LYS1^{KD} \)) lines were successfully generated using artificial microRNA technology (Schwab et al., 2006; Grabherr, 2011). To ensure silencing of \( LYS1 \) in every generation of plants used for experiments, we analyzed the transcript levels of \( LYS1 \) in \( LYS1^{KD} \) lines (generated by Heini Grabherr) (Grabherr, 2011). As shown in Figure 16, \( LYS1^{KD} \) lines contained only approximately 5-20% of WT \( LYS1 \) transcript amounts.
**Figure 16. Analysis of LYS1 knockdown lines.** RT-qPCR analyses of transcript levels in mature leaves of three independent amiRNA knockdown lines (LYS1\textsuperscript{KD}) relative to expression levels in wild-type leaves, which was set to 1. EF1\textalpha transcript was used for normalization. Error bars, SD (n = 3). Statistical significance compared with wild-type (***p<0.001, Student’s t test) is indicated by asterisks.

To exclude the possibility that transcripts of potential amiRNA off-target genes were degraded in LYS1\textsuperscript{KD} lines, we identified potential off-target genes using the Web microRNA Designer and investigated transcripts of the top four hits of the identified off-target genes by RT-qPCR. As shown in Figure 17, the results revealed that transcript levels of these potential off-target genes are not affected.
Results

Figure 17. Determination of putative LYS1 amiRNA off-targets. (A) Predicted LYS1 gene structure (exons, black bars; introns, black lines; untranslated regions, grey). The region targeted by the amiRNA construct is indicated by an arrowhead. (B) Off-target genes for the LYS1-amiRNA construct were identified using the Web microRNA Designer (WMD; http://wmd.weigelworld.org). The region targeted by the amiRNA is given for each gene, mismatches are indicated with grey boxes. Potential off targets either possess more than one mismatch at positions 2–12 or have mismatches at position 10 and/or 11 which will limit amiRNA function. (C) Transcript levels of the four top hits shown in (B) were determined by RT-qPCR in untreated seedlings of two independent transgenic LYS1-amiRNA knock-down lines (LYS1 KD-1, LYS1 KD-2) using gene-specific primers for At4g02540, At1g05615, At5g58780, and At3g51010. EF1α transcript was used for normalization. Error bars, SD (n = 3). No statistically significant differences to the wild-type control (which was set to 1 for each primer set) could be observed (Student’s t test).

3.4.2 Lack of LYS1 PGN-degrading activity dampens plant immunity.
To examine a role of LYS1 in immunity to bacterial infection, we infected wild type plants or LYS1$^{KD}$ and LYS1$^{OE}$ lines with virulent PtoDC3000. Two independent LYS1$^{KD}$ lines exhibited hypersusceptibility to bacterial infection (Grabherr, 2011; Liu et al., 2014), suggesting that lack of PGN-degrading activity results in reduced plant immunity. Likewise, immunity to hypovirulent PtoDC3000 ΔAvrPto/PtoB was compromised in these lines (Grabherr, 2011; Liu et al., 2014). Transcriptional up-regulation of defence-related genes is one of the PTI responses (Felix et al., 1999). Thus, we further examined whether LYS1 protein levels in the transgenics might affect the up-regulation of resistance-related genes. The results showed that expression of the immune marker gene FRK1 upon administration of complex PGN was greatly impaired in the LYS1$^{KD}$ mutants (Figure 18). These findings suggest that the enzymatic activity of LYS1 on PGN contributes substantially to plant immunity against bacterial infection.

![Figure 18. Manipulation of LYS1 levels causes a loss of PGN-triggered immune responses.](image)

Leaves of wild type plants or transgenic LYS1 plants were treated for 6 hours with 100 µg B. subtilis PGN and total RNA was subjected to RT-qPCR using FRK1 specific primers. EF1α transcript was used for normalization. Data represent means ± SD of triplicate samples, and shown is the result of one out of three independent experiments. Statistical significance compared to wild-type (* p < 0.05, Student’s t-test) is indicated by asterisks.

Unexpectedly, bacterial growth on LYS1$^{OE}$ lines was also significantly enhanced as compared to those observed on wild type plants (Grabherr, 2011; Liu et al., 2014). Likewise, FRK1 transcript accumulation upon administration of complex PGN was also strongly reduced in LYS1-overexpressors (Figure 18), indicating that a manipulation of
LYS1 protein levels, irrespective of increasing or decreasing them, results in an impaired immune response towards PGN treatment.

To exclude a direct effect of LYS1-overexpression on PGN receptor abundance, we examined transcript levels of LYM1, LYM3 and CERK1 but found no effect on the transcription of these receptor genes in the LYS1OE lines (Figure 19A). Also, CERK1 protein levels were unaltered in the LYS1OE lines, whereas there was no CERK1 protein detectable in the cerk1-2 mutant (Figure 19B).

Figure 19. LYS1 overexpression does not affect PGN receptor expression. Transcript levels of LYS1 and the PGN receptors LYM1, LYM3 and CERK1 in the strong LYS1 overexpressor line, LYS1OE-1, compared to the weak overexpressor line LYS1OE-3. Total RNA from untreated seedlings (A) or mature leaves (B) was subjected to RT-qPCR using specific primers for LYS1, LYM1, LYM3 or CERK1. EF1α transcript was used for normalization. Data represent means ± SD of triplicate samples. For mature leaves, also CERK1 protein levels were determined using an anti-CERK1 antibody (B, inset). Ponceau S red staining of the large subunit of RuBisCO served as loading control.

Moreover, we included the LYS1OE-3 line with only moderately increased LYS1 transcript and protein levels in mature leaves (Figure 19B and 20A). Susceptibility to Pseudomonas infection in the LYS1OE-3 line was only slightly but not significantly increased (p = 0.064, Student’s t-test) (Figure 20B). These results indicate that lowering LYS1 expression levels, accompanied by lower LYS1 hydrolytic activity on PGN brings down these lines close to wild-type. Thus, massive LYS1 overexpression and loss-of-
function mutations are phenocopies of each other, irrespective of the fact that $LYS1^{KD}$ and $LYS1^{OE}$ lines show dramatic differences in LYS1 enzymatic activities (Grabherr, 2011; Liu et al., 2014).

Figure 20. Impact of weak LYS1 overexpression. (A) Immunoblot analysis of protein extracts from leaves of two independent LYS1$^{OE}$ lines ($LYS1^{OE}$-1, $LYS1^{OE}$-3) and wild type plants. Total leaf protein was subjected to Western blot analysis using α-tobacco class III chitinase (α-Chit) or α-GFP (both from rabbit) and an anti-rabbit HRP-coupled secondary antibody. Ponceau S red staining of the large subunit of RuBisCO served as loading control. (B) Growth of PtoDC3000 was determined 2 days post infiltration of $10^4$ colony forming units ml$^{-1}$ (cfu/ml). Data represent means ± SD of six replicate measurements/genotype/data point. Statistical significance compared to wild-type (* p < 0.05; ** p < 0.01, Student’s t-test) is indicated by asterisks.

3.4.3 LYS1 is able to decompose PtoDC3000 cell but does not inhibit their growth

To further elucidate the role of LYS1 in the interaction between host and pathogen, we compared the effect of LYS1 on PtoDC3000 and E.coli cells in a digestion assay and a growth inhibition assay. Firstly, we tested the ability of LYS1 to degrade bacterial cell walls in a turbidity assay. As shown in Figure 21, LYS1 displayed direct hydrolytic activities towards E.coli (Figure 21A) but not against PtoDC3000 cells (Figure 21B). In a following growth inhibition assay, LYS1 displayed its inhibitory activity on bacterial growth only towards E.coli cells (Figure 21C) but not PtoDC3000 (Figure 21D), suggesting that PtoDC3000 has probably evolved some mechanism to counteract LYS1
activity. However, these are preliminary data are still awaiting confirmation. If this holds true, it will be interesting in the future to investigate how PtoDC3000 interferes with the ability of LYS1 to inhibit bacterial growth.

Figure 21. LYS1 can digest bacteria cells and inhibit bacterial growth. (A, B) Overnight cultured PtoDC3000 and E.coli were incubated with LYS1 purified from 5-week-old LYS1 overexpressing (LYS1\textsuperscript{OE}) Arabidopsis plants or a protein preparation from wild type Arabidopsis plants as a control. After 3 hours the turbidity reduction of the respective culture was determined (A, E.coli; B, PtoDC3000) and set in relation to lysozyme activity, which was set to 1. (C, D) An aliquot of each remaining culture (C, E. coli; D,Pto) was re-cultured overnight and plated on LB medium. Bacterial growth given in colony forming units (cfu per ml) was determined after 24 hours.

Altogether, we propose that LYS1 contributes to plant immunity against bacterial infection by decomposition of bacterial PGN and generation of soluble PGN-derived
patterns that trigger immune activation in a LYM1-LYM3-CERK1 receptor-complex-dependent manner.

### 3.5 Identification of a CERK1-interacting calcium-dependent protein kinase

#### 3.5.1 Identification of putative CERK1 interactors from a Y2H database

To better understand the signaling pathways induced by PGN and chitin, respectively, we next aimed at identifying novel interacting proteins for CERK1. The yeast two-hybrid system is a useful and powerful genetic technique for the in vivo analysis of protein-protein interactions (Bartel and Fields, 1995). However, its application is limited in that hybrid proteins generated in the two-hybrid assay are targeted to the nucleus. Therefore, integral membrane proteins, which exist in the lipid biolayer, are excluded because they are unlikely to be able to enter the nucleus, or will be misfolded if they do. To overcome limits of the conventional yeast two-hybrid system, an alternative split-ubiquitin system was developed in 1994 (Johnsson and Varshavsky, 1994) and then modified for the in vivo analysis of membrane proteins on a small-scale basis (Stagljar et al., 1998). Later on, a mating-based split ubiquitin system (mbSUS) was developed for systematic identification of interactions between membrane proteins as well as between membrane and soluble proteins on a large-scale basis (Obrdlik et al., 2004). In the mbSUS, two integral membrane proteins are fused to the two halves of ubiquitin and expressed in yeast cells of opposite mating types. Upon mating, the diploid yeast cell co-expresses the proteins. An interaction of membrane proteins brings the two halves of ubiquitin into close proximity, forming a reconstituted molecule that is cleaved by ubiquitin-specific proteases, releasing the transcription factor to enter the nucleus and activate reporter gene transcription.

With this mbSUS system, Frommer and his colleagues used a library of more than 3000 Arabidopsis membrane proteins and soluble signaling proteins to screen over 3 million binary interactions. The results lead to a membrane-based interactome network database (M.I.N.D.) containing 12102 high confidence protein-protein interactions that were identified in repeated rounds of interaction screening (associomics.org).
Table 7. Putative CERK1 interacting proteins identified from the Membrane-based Interactome Network Database (M.I.N.D. 0.5).

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<tr>
<td>AT1G58520</td>
<td>RXW8__lipases; hydrolases, acting on ester bonds</td>
</tr>
<tr>
<td>AT1G79820</td>
<td>SGB1__Major facilitator superfamily protein</td>
</tr>
</tbody>
</table>

In order to identify novel components interacting directly with CERK1, we searched CERK1-interacting proteins in the database of the M.I.N.D and obtained 20 putative interacting proteins (Table 1).

Of these 20 putative CERK1-interacting proteins, three protein kinases were selected for further study: a protein kinase without known function (PK, At1g69910), CBL-interacting protein kinase 7 (CIPK7, At3g23000) and calcium-dependent protein kinase 15 (CPK15, At4g21940).
3.5.2 Analysis of M.I.N.D.-interactors for CERK1

Figure 22. Gene models and genotyping of T-DNA insertion mutants of three putative CERK1 interactors. Gene models of CIPK7 (A), PK (B) and CPK15 (C, D) including the positions of the T-DNA insertions. Exons and introns are indicated by grey bars and grey lines, respectively. 5'- and 3'-UTR regions are represented by light grey bars and the T-DNA insertions by grey triangles. Promoter regions are indicated by thin lines. For genotyping the mutants, leaf genomic DNA was isolated and genotyping PCRs were performed. (A) Genotyping of pk-1 and pk-2 was done with primer pairs Salk-Lba/pk1-rp and GK-Lba/pk2-rp for the Lba-PCR, and pk1-lp/pk1-rp and pk2-lp/pk2-rp for the WT-PCR. (B) Genotyping of cipk7-1 and cipk7-2 was done with primer pairs Salk-Lba/cipk1-rp and GK-Lba/cipk2-rp for the Lba-PCR, and cipk1-lp/cipk1-rp and cipk2-lp/cipk2-rp for the WT-PCR. (C) Genotyping of cpk15-1 and cpk15-2 was
done with primer pairs Sail-Lba/cpk15-rp and F3/JL-270 for the Lba-PCR, and cpk15-lp/cpk15-rp and F3/R3 for the WT-PCR. For transcript analysis using semi-quantitative RT-PCR total RNA was isolated from leaves and transcribed into cDNA. EF1a-s and EF1a-as primers were used to amplify the transcript of the house-keeping gene EF1α. F3/R2 and F3/R1 primer pairs were used to amplify the transcripts of CPK15. (D) Two gene models of CPK15 are indicated by GM1 and GM2.

Two independent T-DNA mutant lines of each selected gene (*PK*, *CIPK7*, and *CPK15*, Table 1) were obtained from the Nottingham Arabidopsis Stock Center (NASC). Genotyping analyses of these mutants were performed by Lba-PCR and WT-PCR, revealing that these mutants were homozygous T-DNA insertion mutants (Figure 22, A-C).

The accumulation of ROS in an oxidative burst is an early response triggered by PAMPs and is also observed in *Arabidopsis* upon treatment with chitin (Miya et al., 2007). We thus investigated if the oxidative burst was impaired in these potential CERK1 interacting protein mutants in response to chitoctamers in a leaf ROS assay. As shown in Figure 23, no significant changes were detected in the two *PK* mutants and the two *CIPK7* mutants in comparison to the Col-0 wild type. However, *cpk15-1* plants displayed a significant reduction in ROS production whereas *cpk15-2* mutants did not (Figure 23, right panel). We thus chose CPK15 for a more detailed investigation as a putative CERK1 interactor.
Figure 23. Screening analysis of T-DNA mutants of \( PK \), \( CIPK7 \) and \( CPK15 \) in a ROS oxidative burst assay. Leaf pieces from 5-week old Col-0, \( pk \), \( cipk7 \), \( cpk15 \) and \( cerk1-2 \) mutants were elicited with 1 µM chitooctamer (C8) and the oxidative burst was measured in a 96-well plate using a plate reader (for more details see method section). The \( cerk1-2 \) mutant served as a negative control. Mean and standard error of 6 replicates are presented. Asterisks indicate significant differences of C8 treatment in comparison to Col-0 \((^*p<0.05, ^{*}*p<0.001\), Student’s t test).}

3.6 \( CPK15 \) is involved in the CERK1-mediated PTI pathway

3.6.1 Characterization of \( cpk15 \) T-DNA insertion lines

It was predicted that there are two possible gene models of CPK15 as shown in Figure 22D, with gene model 1 (GM1) comprised of 8 exons while gene model 2 (GM2) contains 9. In order to examine which of two gene models is correct, the constructs \( p35S::GM1-YFP \) and \( p35S::GM2-YFP \) were expressed in \( N. benthamiana \). YFP fluorescence signals could only be observed in leaves expressing \( GM1-YFP \) (Tina Romeis, personal communication), suggesting GM1 is the correct model.

Next we analyzed \( cpk15 \) T-DNA mutant lines more thoroughly. Sequence comparison of two T-DNA flanking regions with CPK15 genomic DNA revealed that the T-DNA was inserted in the promoter region in \( cpk15-1 \) and in the third exon in \( cpk15-2 \) as shown in Figure 22C. To examine the transcript levels of \( CPK15 \) in the two T-DNA insertion
mutants, total RNA was isolated from leaves of the two mutants and used for semiquantitative RT-qPCR analysis with gene-specific primer pairs (Figure 22C). The agarose gel electrophoresis analysis of the amplified products revealed that no 3’-end but the 5’-end transcript of the CPK15 could be detected in the two cpk15 mutants (Figure 22C). Since cpk15-2 did not display significant differences in the ROS-based screening, we next focused on the cpk15-1 line. Morphology observation of cpk15-1 mutant plants revealed no differences between Col-0 and cpk15-1 mutants except that cpk15-1 leaves displayed curved edges (Figure 24). However, a southern blot to exclude multiple T-DNA insertion events in the cpk15-1 line is still outstanding. Thus, so far we cannot rule out that this leaf phenotype is caused by a secondary T-DNA insertion and not by genetic inactivation of the CPK15 gene itself.

![Figure 24. Morphological phenotypes of cpk15-1 T-DNA insertion lines. 5-week old plants of the cpk15-1 line and the Col-0 wild type were photographed to show whole plants from the top (left) and leaves from the adaxial and abaxial sides (right).](image)

3.6.2 cpk15-1 produces less ROS in response to complex or soluble chitin

In order to investigate if cpk15-1 mutants were compromised in ROS accumulation in response to complex chitin as well, five-week old leaves of cpk15-1 and Col-0 plants were treated with Flg22, complex crab chitin and soluble chitooctamer and subsequent ROS production was determined in a 96-well plate reader. The results showed that upon treatment with both crab chitin and C8, cpk15-1 mutants exhibited a significant reduction of ROS accumulation compared to Col-0 plants (Figure 25). Notably, cpk15-1 displayed a tendency of higher ROS levels in response to Flg22 treatment (Figure 25). The cerk1-
2 mutation almost abolished ROS accumulation in response to chitin, although in other experiments this mutant showed a normal Flg22-induced response level (data not shown).

**Figure 25. Determination of ROS accumulation in cpk15-1 mutant plants.** Leaf pieces from 5-week old Col-0, cpk15-1, and cerk1-2 plants were treated with 100 µg/ml crab chitin or 1 µM chitooctamer (C8). The oxidative burst was measured at 20 min after treatment in a 96-well plate reader (for more details see method section). The oxidative burst triggered by 100 nM Flg22 served as a positive control. The cerk1-2 mutant served as a negative control only for chitin treatment. Mean values with standard errors of 6 replicates are represented. Asterisks indicate significant differences of each treatment in comparison to the Col-0 control (**p<0.01, ***p<0.001, Student’s t test).

### 3.6.3 Chitin-induced activation of MAPK is not affected in cpk15-1

In addition to the production of reactive oxygen species, early cellular events upon perception of PAMPs also involve the post-translational activation of mitogen-activated protein kinase (MAPK) cascades (Boller and Felix, 2009). To investigate if the activation
of MAPKs is affected in the *cpk15*-1 mutant line, comparative studies with Col-0 upon treatment with chitin were performed. MAPK activities were analyzed by immunoblot assays using the p44/12 antibody raised against phosphorylated MAPKs (Boller and Felix, 2009). As shown in Figure 26, chitohexamers (C6) strongly activated the defence-associated MPK3, MPK4/11 and MPK6 in *cpk15*-1, which was indistinguishable from the induction pattern obtained in Col-0 plants. In *cerk1*-2 plants used as negative control, C6-induced activation of MAPKs was completely abolished, and water treatment did not induce the activation of MAPKs in all tested plants. In contrast, as positive controls, Flg22 induced a high level of activation of MAPKs in all plant types (Figure 26). Hence, we conclude that CPK15 is involved in a signaling pathway independent from chitin-induced MAPK activation.

![Figure 26. Chitin induced MAPK activation is not impaired in the *cpk15*-1 mutant.](Image)

**Figure 26. Chitin induced MAPK activation is not impaired in the *cpk15*-1 mutant.** 7-day old seedlings of the *cpk15*-1 or *cerk1*-2 mutant, or Col-0 as a control, were collected at indicated time points after treatment with 10 µM chitohexamer (C6). Treatment with water and 100 nM Flg22 served as negative and positive controls, respectively. The crude protein extracts from these seedlings were separated on a SDS-PA gel and blotted onto a nitrocellulose membrane. Immunodetection was carried out using the anti-phospho p44/42 antibody. Arrowheads indicate the positions of MAP kinases 6, 3 and 4/11.

### 3.6.4 CPK15 localizes to the plasma membrane
Most CDPKs have a predicted N-myristoylation site involved in membrane targeting (TermiNator, http://www.isv.cnrs-gif.fr/terminator2/index.html). This irreversible co-translational acylation requires a second post-translational signal to maintain the membrane association, such as reversible palmitoylation (Martin and Busconi, 2000), leading to most CDPKs anchored in the membrane.

**Figure 27. Subcellular localization of the CPK15-YFP fusion protein.** Constructs \textit{p35S::CPK15-fl-YFP} (upper panel) and \textit{p35S::CPK15-vk-YFP} (middle panel) were expressed in Arabidopsis mesophyll protoplasts. VK represent a truncated CPK15 version (variable domain and kinase domain). The localization was visualized by laser scanning confocal microscopy. \textit{p35S::GFP} was used as a control (lower panel).

CPK15 has a predicted N-myristoylation site and could be localized at the plasma membrane. To examine the localization of CPK15, we expressed a \textit{p35S::CPK15-fl-YFP} construct and a construct for a truncated version \textit{p35S::CPK15-vk-YFP} (containing only the variable and kinase domain of CPK15, see also Figure 29, constructs kindly
provided by Prof. Tina Romeis) in *Arabidopsis* mesophyll protoplast. As shown in Figure 27, confocal images revealed that YFP-fluorescence signals appear on the plasma membranes of protoplasts expressing the full length and the truncated version of CPK15, confirming a localization of CPK15 at the plasma membrane as predicted, whereas the control-GFP signal could only be found in the cytoplasm. Moreover, these results indicate that the variable and kinase domain of CPK15 are sufficient to target the protein to the plasma membrane.

3.6.5 CERK1 physically interacts with CPK15

![Figure 28. CERK1 physically interacts with CPK15. Constructs p35S::CERK1-HA and p35S::CPK15-GFP were transiently (co-)expressed in *N. benthamiana* leaves by agrobacteria-mediated transformation. After 36 hours, CPK15-GFP was immunoprecipitated from total protein extracts using GFP-Trap and detected with an anti-GFP antibody. Co-immunoprecipitated CERK1-HA was detected with anti-HA antibody. Western blot analysis with anti-HA and anti-GFP of corresponding total proteins served as input controls.]

Identification via the M.I.N.D. already suggested that CERK1 has the ability to interact with CPK15 in a yeast-based system. Thus, we next investigated if CERK1 directly physically interacts with CPK15 *in planta*. To do this, CERK1-HA and CPK15-GFP were transiently co-expressed in *N. benthamiana* and protein extracts were used for a co-immunoprecipitation analysis. An immunoprecipitation of CPK15-GFP with GFP-Trap (a GFP-binding protein coupled to agarose beads) was performed and co-immunopurified
proteins were subjected to immunoblotting analysis with HA antibodies. The results revealed that CERK1-HA could be co-immunoprecipitated by CPK15-GFP (Figure 28), indicating a direct physical interaction between CPK15 and CERK1. Thus, CERK1 and CPK15 can interact both in yeast cells and in plant tissue.

3.7 Characterization of knockdown lines of CPK15 and CPK15/CPK21/CPK23

3.7.1 Structure and classification of Arabidopsis CDPKs

![Diagram of Arabidopsis CDPKs]

Figure 29. Structure and subfamilies of Arabidopsis CDPKs. (A) Schematic general structure of CDPKs. N, N-terminal variable domain; CaM, calmodulin-like domain. The four bars within the CaM-like domain represent the EF hand Ca$^{2+}$-binding sites. (B) Subfamilies of Arabidopsis CDPKs. The complete protein sequences of the Arabidopsis CDPKs were aligned and analyzed by the Treeview 1.6.5 program (http://taxonomy.zoology.gla.ac.uk/rod/rod.html). The uprooted distance tree reveals the presence of four distinct, branched subgroups (I–IV). The branch lengths are proportional to divergence, with the scale of “0.1” representing 10% change. CPK15 and its closest homologs CPK21/23 belong to group II. This figure was modified from Cheng et al. (Cheng et al., 2002).
There are 34 Arabidopsis CDPKs in *Arabidopsis* which are highly homologous to each other (Boudsocq and Sheen, 2013) (Figure 29B). Pair-wise analyses with the full protein sequences indicated that the overall identities and similarities are 39% to 95% and 56% to 96%, respectively. High homologies may indicate redundant functions. Based upon sequence homology, the CDPKs of *Arabidopsis* cluster into four subgroups (I–IV). Subgroup IV is the least complex, with three members, and subgroup II is the most complex, with 13 members. Subgroups I through III are closer in sequence identity to each other than to subgroup IV. CPK15 belongs to subgroup II and its closest homologs are CPK21 and CPK23.

### 3.7.2 Generation of artificial microRNA lines of *CPK15*, *CPK21* and *CPK23*.

![Diagram of gene editing](image)

**Figure 30. Generation of *CPK15* single and *CPK15/21/23* triple knockdown lines.** (A) *CPK15* gene with regions targeted by the artificial microRNA. Two sets of amiRNA constructs were generated with one targeting the region from 334-354 nucleotides (nt) and another targeting the region from 1792-1802 nt of the *CPK15* gene. (B) For the multiple gene knockdowns one construct was generated with an amiRNA targeting the three genes *CPK15*, *CPK21* and *CPK23*.

To generate a second *CPK15* knockdown (*CPK15*\(^{KD}\) allele and to genetically inactivate *CPK15* together with its homologs, we made use of an artificial microRNA system
(Schwab et al., 2006) to create single CPK15 knockdown plant (Figure 30A), in which only CPK15 is targeted, and triple knockdown plants, in which CPK15, CPK21 and CPK23 are targeted (Figure 30B). T1 seeds of CPK15\textsuperscript{KD} have been harvested and need to be further confirmed by qPCR and the constructs for cpk15/21/23 amiRNA triple knockout mutants are under construction.
4. Discussion

Although plant-microbe interactions have occurred for several millions years, a close look at the underlying molecular mechanism was initiated only some fifty years ago. It is understood that plants activate their immune responses by perceiving microbe-derived molecular patterns via cell surface-localized immune receptors. However pre- and post-perception events remain largely a mystery. It is unclear if and how microbial pattern supermolecules are processed prior to perception by cell surface receptors. The mechanism linking perception with diverse downstream immune responses is also largely unknown. Therefore, this thesis focused on these topics and aimed at answering these important questions.

It is generally little understood whether, and if so, how microbial patterns derived from complex extracellular assemblies, such as bacterial cell walls, are accessible to host PRRs for host immune activation in eukaryotes. This is true not only for bacterial PGNs, but also for other microbial patterns such as bacterial LPS, flagellin, fungal chitin or glucan structures, all of which have been identified as inducers of innate immunity in metazoans and plants (Boller and Felix, 2009; Kumar et al., 2013; Newman et al., 2013; Pel and Pieterse, 2013). Recent research into the 3D structure of ligand-PRR complexes, as well as studies on ligand structural requirements for plant immune activation, suggest that small ligand epitopes are crucial for binding to host PRRs (Liu et al., 2012b; Sun et al., 2013). It is thus generally assumed that soluble fragments derived from complex microbial matrices serve as ligands for host PRRs and subsequent immune activation in both metazoans and plants.

4.1 LYS1 is involved in the apoplastic battle between plants and pathogens

In general, upon invasion foliar plant pathogens encounter plant defence in three battlegrounds: the plant surface, plant cell apoplast, and the plant cell symplast (Senthil-Kumar and Mysore, 2013). Plant defences in the three battlegrounds are generally classified into two types: preformed defence and inducible defence. Pathogens that attempt to invade plants are initially exposed to a wide range of preformed plant defence including both physical barriers (Gottwald and Graham, 1992; Xiao et al., 2004) and
chemical inhibitors (Dixon et al., 2002; Aires et al., 2009; Che et al., 2011). Subsequently, pathogens that overcome these preformed defence encounter the plant inducible defence (Li et al., 2005; Nicaise et al., 2009; Sohn et al., 2012). Both preformed and inducible defence are integrated by plants across the three battlegrounds and deployed during infection to combat pathogens.

Preformed defence, although important, are passive and constitutive responses. Inducible defence, in contrast, are active and stimuli-oriented responses (Senthil-Kumar and Mysore, 2013). Thus, the inducible defence allows plants to induce immune responses at appropriate levels in response to a given pathogen infection and thus is rather cost-saving (Heil and Baldwin, 2002), compared to the constitutive defence which use the same plant energy resources. As a result, plants can divert the saved cost from defence to growth and reproduction, which is a benefit for plant fitness (Heil and Baldwin, 2002).

Of the three battlegrounds, the apoplast is the first place where the pathogens establish their physical contact with plant cells. On the one hand, pathogens attempt to adapt to and colonize the apoplast. On the other hand, plants impede these colonization attempts via preformed and induced barriers. This opposition makes the apoplast a key battleground where the fate of both the pathogens and the plants will be determined during infection.

Plant apoplastic defence involves physical barriers, antimicrobial chemical compounds and antimicrobial enzymatic activities of pathogenesis-related (PR) proteins. For instance, cell wall reinforcement by callose, lignin, and suberin deposition is induced by pathogens as part of the plant defence against pathogens (Senthil-Kumar and Mysore, 2013). In addition, some chemical compounds, such as phytoanticipins and phytoalexins, are also present constitutively or inducibly in the apoplast and act as antimicrobial agents (Senthil-Kumar and Mysore, 2013). Moreover, important components of the plant defence response against microbial pathogens in the apoplast involve proteins such as PR proteins that are produced upon pathogen perception in order to restrict pathogen growth. The apoplastic PRs, which actually comprise all PR families except the ribonuclease-like PR-10, can be found in cell wall appositions in response to pathogen attack. PRs have various antimicrobial activities, some of which harbor carbohydrate-
degrading hydrolytic enzyme activities. In the case of the β-1,3-glucanase PR-2 or the chitinases PR-3/4/8/11, these hydrolytic enzymes have been implicated in plant immunity via their enzymatic attacks on major components of fungal cell walls (van Loon et al., 2006).

In this study, we showed that the immunity-associated lysozyme-like protein LYS1 functions in the apoplast. LYS1, a PR-8 protein, is predicted to have a 22-amino acid secretion signal at its N-terminus (Grabherr, 2011). This signal peptide directs LYS1 to be secreted into the apoplast and is then removed by a peptidase to produce a mature LYS1. This is in accordance with the results of our LYS1 MS analysis, which showed that none of the identified peptides of the purified mature LYS1 were mapped to this N-terminal signal peptide region (Figure 11C). Moreover, immunoblotting analysis of the apoplastic fluid of LYS1OE plants further confirmed the presence of LYS1 in the Arabidopsis apoplast (Figure 14). Finally, a previous apoplast proteome analysis supported our finding that LYS1 was an apoplastic protein (Kwon et al., 2005).

Many secretory proteins undergo some posttranslational modifications, (e.g. glycosylation), which are often required for enzyme activities (Tekoah, 2004). The glycosylation of LYS1 was investigated and confirmed by deglycosylation experiments (Grabherr, 2011). In contrast, LYS1 lacking the signal peptide cannot properly be secreted into the apoplast (Grabherr, 2011) and displayed no enzymatic activity (Figure 6B).

In addition to posttranslational modification, the correct folding of a mature protein is also critical for its activity. The prediction of the LYS1 three dimensional structure using the Phyre server (Kelley and Sternberg, 2009) revealed that LYS1 has a (ββα)₈ barrel fold and that the substrate-binding cleft is formed by the C-terminal residues of barrel β-strand and subsequent loops (Figure 3B). Therefore, C-terminally fused tags might interfere with the correct folding of LYS1 and the tight binding of substrates, resulting in a loss of the enzymatic activity. Indeed, in this study all three LYS1 fusions with C-terminal tags (HA, myc and GFP) displayed no detectable enzymatic activities (Figure 7 and 10).

Taken together, LYS1 is an apoplastic protein whose activity depends on the required posttranslational modifications and correct folding. Since LYS1 lysozyme-like activity is
considered to be related to plant immunity, we conclude that LYS1 functions as an immunity-associated protein in the apoplast battleground.

4.2 LYS1 is involved in plant inducible defence

The inducible defence responses can occur through PAMP-triggered immunity (PTI) or through effector-triggered immunity (ETI) (Jones and Dangl, 2006). In PTI, some highly-conserved, microbe-derived molecular patterns can be recognized by plant surface-localized pathogen recognition receptors (PRRs) to induce immune responses. Structural analyses have demonstrated that ligand-induced oligomerization of PRR is a common theme for the activation of plant PRRs (Chinchilla et al., 2007; Liu et al., 2012b; Sun et al., 2013), suggesting that small ligand epitopes are crucial for binding to the host PRRs. It is generally assumed that PAMPs are present as soluble fragments in the plant apoplast to gain better access to the plasma membrane-localized PRRs and thus activate the immune responses. In fact, it was reported that soluble chitin oligomers are bound by the LysM domains of two receptor monomers, resulting in the receptor dimerization and the activation of subsequent signaling (Liu et al., 2012b; Hayafune et al., 2014). It has also been shown that soluble oligomeric PGN fragments stimulate plant immune responses in Arabidopsis (Gust et al., 2007; Erbs et al., 2008; Willmann et al., 2011).

Soluble PAMPs are proposed to be generated from macromolecular assemblies in two possible ways. First, imperfect recycling of bacterial wall components might serve passively as sources of soluble ligands for host PRRs sensing (Nigro et al., 2008). Second, host hydrolytic enzymes might release actively soluble ligands from bacterial envelope structures (Wang et al., 2006). The second option prompted us to investigate whether this way also is important in Arabidopsis immunity, and if so, which protein is responsible for the generation of soluble PGN from the insoluble PGN complex. The results in this study showed that LYS1 harbored PGN hydrolytic activity (Figure 12) and was able to release soluble fragments from insoluble PGN (Figure 15). More importantly, these LYS1-generated soluble PGN fragments harbored immunogenic activity and triggered immunity-associated responses in a PGN receptor-dependent manner (Figure 15B).
Therefore, we conclude that lysozyme-like LYS1 contributes to the inducible defence in *Arabidopsis* by generating the soluble PGN fragments to activate PTI.

It was demonstrated that soluble chitin oligomers induced the formation of a receptor complex at the plasma membrane to activate subsequent signaling in *Arabidopsis* and rice (Cao et al., 2014; Hayafune et al., 2014). Likewise, it was postulated that soluble PGN oligomer binding to LYM1 and LYM3 results in the formation of a tripartite complex of LYM1, LYM3 and CERK1 to activate subsequent immune responses (Willmann et al., 2011), but the direct evidence of physical LYM1/LYM3/CERK1 interaction *in planta* is still lacking. It remains to be elucidated if LYM1, LYM3 and CERK1 form a protein complex, and if so, how the receptor complex formation is associated with the PGN ligand.

Recent advances in the study of chitin receptor structures revealed that a sandwich-type dimerization of receptor monomers is induced by chitin oligomers and is required for activation of immune signaling in *Arabidopsis* and rice (Liu et al., 2012b; Cao et al., 2014; Hayafune et al., 2014). In both *Arabidopsis* and rice, the fragment sizes of eight GlcNAc were defined as the minimum length for inducing dimerization of the receptor complex. The similarity of receptors and substrates between the chitin-perception system and the PGN-perception system raises the question if the minimum immunogenic unit model in the chitin-perception system can be generalized to the PGN-perception system. Previous data showed that a PGN glycan chain that is longer than the disaccharide can display immunogenic activity, but the exact minimum immunogenic unit of PGN is still unknown (Gust et al., 2007). Attempts to identify the minimum immunogenic motif of PGN have failed so far due to technical limits. The synthesized PGN oligomers up to heptamers (kindly provided by Koichi Fukase and Yukari Fujimoto, Osaka University) showed no immunogenic activity, and higher oligomers could not be obtained (data not shown). Although LYS1-digested PGN can be efficiently separated by HPLC (Figure 15A), the amount of PGN in each collected peak is so far not enough to perform subsequent immune assays. These limitations could be overcome in the future by advancements in HPLC purification and PGN synthesis.

In summary, based on the activation mechanism of the chitin receptors, it is most likely that plants can only perceive PGN ligands within a defined range, possibly generated by
LYS1, whereas insoluble complex PGN and smaller units do not have immunogenic activity.

4.3 LYS1 is a bifunctional lysozyme/chitinase

Plants do not contain chitin in their cell walls, whereas major agricultural pests such as most fungi and insects do, which leads to the hypothesis that plant chitinases act as a defence mechanism against pathogens. Indeed, plant chitinases are often considered as PR-proteins, since their activities can be induced by abiotic and biotic stress (Graham and Sticklen, 1994).

Plant chitinases play diverse physiological roles in plants (Grover, 2012) and are classified into two classes, GH18 and GH19, which do not share amino acid sequence similarity (Henrissat, 1991). These two chitinase families have completely different 3-dimensional structures and molecular mechanisms despite shared chitinolytic activity (Hamid et al., 2013), possibly indicating convergent evolution of the two families from different ancestors. Chitinases of the GH18 class are ubiquitously found in all organisms, whereas those of the GH19 class are found almost exclusively in plants.

A comparative study of class I (GH19) and class III (GH18) chitinases revealed that an excess of amino acid replacements can be found at the active site and substrate binding cleft of class I chitinases (Bishop et al., 2000). This highly unusual pattern of replacements in the plant class I chitinases was proposed as a rapid adaption to the fungal pathogens that had evolved to overcome plant chitinolytic activity through enzymatic inhibition or modification of their cell wall. Such an anomaly suggests that class I chitinases face a highly positive selective pressure from the fungal pathogens. Therefore, similar to R genes, plant defence proteins such as chitinases might also undergo a rapid adaptive evolution driven by an arms race between plants and fungal pathogens. This raises the question why the class III chitinases did not display such frequent amino acid substitutions within the active site cleft. One possibility is that the similarity of the protein fold and function between plant class III and microbial class III chitinases could protect the plant class III chitinases from evolutionary pressure. In other words, if a given pathogen evolves to produce an inhibitor for plant class III chitinases, this inhibitor likely also inhibits its own class III chitinase, thereby interfering with
physiological functions of this chitinase in the pathogen and likely resulting in a disadvantage for the pathogen.

Of plant GH18 chitinases, the basic class III chitinases were found to display high lysozyme activities such as hevamine in rubber trees, lysb1 and lysb2 in tobacco or OsChib1b in rice (Beintema et al., 1991; Brunner et al., 1998; Park et al., 2002). These chitinases are basic proteins with high isoelectric points. Structurally, these proteins possess a (βα)_8 barrel domain, and a substrate-binding cleft at the C-terminus containing two aspartate and one glutamate residue separated by phenylalanine and isoleucine (DxDxE) (Tsuij et al., 2010) and belong to the PR-8 protein family (van Loon et al., 2006).

LYS1, as the only members of class III chitinases in Arabidopsis, contains all above-mentioned features. LYS1 is predicted to have an isoelectric point of 9.3. As a basic protein, a net positive charge (pI 9.3) on the surface would facilitate LYS1 binding to the negatively charged bacterial cells to perform its lysozyme activity. Moreover, LYS1 has a predicted (βα)_8 barrel structure and a C-terminal substrate binding cleft (Figure 3B). This architecture has been confirmed by a crystal structure study of hevamine, a bifunctional class III lysozyme/chitinase in rubber tree (Terwisscha van Scheltinga et al., 1994). Based on the shared architecture in class III members, we assumed that LYS1 may harbor hydrolytic activities towards both chitin and PGN as other known plant class III chitinases do. The results of this study confirmed that LYS1 is able to hydrolyze β (1, 4) linkages between MurNAc and GlcNAc in PGN and between GlcNAc in chitin oligomers (Figure 12), indicating its potential role in PGN-induced immunity.

Although LYS1 was initially named as chitinase A (CHIA), we demonstrated in this study that LYS1 also harbors lysozyme activity (Figure 12B-C). LYS1 was first isolated in 1990 and its gene expression was investigated upon fungal infection (Samac et al., 1990; Samac and Shah, 1991). The investigation revealed that LYS1 transcript accumulation was dependent on the fungal strain used for infection and only occurred at the infection site. Despite its classification as a chitinase and the fact that it indeed has chitinase activity (Figure 12A), LYS1 did not display significant anti-fungal function upon infection of B. cinera and A. brassicicola (Grabherr, 2011).
Discussion

2011) (Grabherr, 2011). For example, plants with LYS1/CHIA levels less than 10% of that of the wild-type showed no sign of increased susceptibility to *Botrytis cinerea* (Samac and Shah, 1994). These results are in agreement with our lab’s finding that LYS1<sup>KD</sup> plants did not display higher susceptibility to *B. cinerea* and *A. brassicicola* (Grabherr, 2011). This may suggest that LYS1 has lost its role in defence against these fungi possibly due to an unknown counteractive mechanism co-evolved by the fungal pathogens. However, LYS1 still plays an important role in plant defence against bacterial pathogens like *PtoDC3000* (Figure 20), and this anti-bacterial function is associated with its lysozyme activity (Figure 15).

Taken together, the bifunctional LYS1 has likely lost its anti-fungal activities towards *B. cinerea* and *A. brassicicola* but retained anti-bacterial activities towards *PtoDC3000* possibly due to co-evolutionary interaction of plants and pathogens, and thus it plays a unique role in plant immunity in *Arabidopsis*.

4.4 How does LYS1 contribute to defence: direct killing or generating PAMPs?

When microbes invade the apoplast of plant cells, the host cells secret hydrolytic enzymes, such as chitinases and glucanases, to target the constituents of the microbial cell wall and thus disrupt cell wall integrity (Schlumbaum et al., 1886). Thus, hydrolytic enzyme activities can have a dual function for the host plant: Firstly, releasing PAMP molecules that further stimulate immune responses and secondly, causing cell collapse in the invader to inhibit its growth (Kombrink et al., 2011).

In mammals, PGN hydrolytic enzyme activities such as lysozymes have been ascribed functions in direct bacterial killing (Cho et al., 2005) and in generating soluble PGN fragments as ligands for PRRs (Cho et al., 2005; Dziarski and Gupta, 2010; Davis et al., 2011). For example, human PGRP-S, one PGN recognition protein, is co-located with lysozyme in the granules of human neutrophils. The two proteins act together to kill bacteria trapped in the neutrophil extracellular traps (Cho et al., 2005). In addition, lysozymes in mice can release PGNs as PAMPs for recognition by NOD2 to activate immune responses (Davis et al., 2011).

A role for plant glycosyl hydrolases in immunogenic PAMP generation and immune activation has also been proposed previously (Mithöfer et al., 2000; Fliegmann et al.,
An extracellular soluble bipartite soybean glucan binding protein (GBP) was shown to harbor 1,3-β-glucanase activity and binding activity for glucan fragments of DP > 6 derived of intact glucans. Complex glucans constitute major constituents of various *Phytophthora* species, many of which are plant pathogens (Kroon et al., 2011). It was hence suggested that during infection GBP endoglucanase activity produces soluble *Phytophthora*-derived oligoglucoside fragments as ligands for the high-affinity binding site within this protein (Fliegmann et al., 2004). While this study supported the concept of plant hydrolases tailor-making ligands for plant PRRs, causal evidence for the involvement of the endoglucanase activity in plant immunity was not provided.

In this study, LYS1 was identified as a plant lysozyme-like enzyme which harbors PGN hydrolytic activity (Figure 12). We assume two possible scenarios as to how LYS1 is involved in plant defence. First, LYS1 might directly kill bacterial cells via hydrolyzing bacterial cell walls. In an *in vitro* digestion and inhibition assay, LYS1 could indeed digest *E. coli* cells directly, subsequently leading to a reduced growth of *E. coli* on agar plate (Figure 21C). The bactericidal activity of LYS1 could contribute to plant basal immunity against non-adapted bacteria and even some adapted bacterial pathogens. In response, adapted bacterial plant pathogens may have evolved a counteractive mechanism to avoid digestion of their cell wall. As we observed in Figure 21, LYS1 could not digest *Pto*DC3000 cells and could not inhibit its growth, indicating that phytopathogenic *Pto*DC3000 might have evolved inhibitory mechanisms against *Arabidopsis* LYS1. This could also explain our observation that *LYS1*OE transgenics are more susceptible and not more resistant to *Pto*DC3000 in spite of high LYS1 levels in the *LYS1*OE lines (Figure 20B), which argues against an important role for LYS1 in killing *Pto*DC3000. Second, LYS1 most likely provides the PGN receptors with soluble immunogenic ligands that stimulate the plant immune system to fend off invading bacterial pathogens. Our analysis showed that LYS1 can generate soluble fragments from PGN complex, and these fragments displayed immunogenic activities in rice and *Arabidopsis* (Figure 15). These immune responses were PGN receptor-dependent, thereby confirming that LYS1-generated PGN fragments are subsequently recognized by the receptors to trigger downstream responses.
The two LYS1 functions involved in defence may co-exist in plants to promote synergetic plant immunity. However, the second LYS1 function of generating soluble PAMPs to induce the immune system seems to be more efficient. In order to directly and completely kill bacteria via bactericidal activity, LYS1 protein is most likely required in relatively high amounts in the plant apoplast. In comparison, relatively little amounts of LYS1 are able to release PGN ligands, which initiate signaling and later even amplify the immune responses. Moreover, in the co-evolution of pathogens and plants, plant-secreted LYS1 could be targeted by inhibitors of pathogens and lead to reduced enzymatic activity. Such compromised enzymatic activity of LYS1 could not kill bacteria directly, but could still release PGN ligands from bacterial cells to induce immune responses.

Indeed, although we did not find LYS1 hydrolytic activity towards \textit{PtoDC3000} cells (Figure 21), \textit{LYS1}\textsuperscript{KD} lines are more susceptible to \textit{PtoDC3000} than WT (Figure 20B). We assume that LYS1 in WT could still retain weak hydrolytic activity at levels sufficient to produce enough PAMPs for induction of immune responses in plants (Figure 20B). Taken together, LYS1 contributes to plant resistance to \textit{PtoDC3000} by generating immunogenic ligands rather than directly killing bacteria.

4.5 Dynamics of LYS1 protein levels are related to immunity performance

In plants, there is generally a trade-off between growth and immunity. In order to reduce the fitness cost, plants fine-tune appropriate immune responses to invading pathogens. LYS1-mediated immunity is no exception.

In wild type plants, LYS1 was constitutively expressed at low level (Figure 4). When pathogenic bacteria enter the apoplast, positively-charged LYS1 at low levels could efficiently bind to the invading negatively-charged bacterial cell walls and hydrolyze them, at least partially. Then LYS1-released PGN ligands get better access to plasma membrane-localized receptors and bind to them to activate subsequent immune responses, including increased LYS1 levels at the infected site. The induced, high-level LYS1 could contribute to immunity via direct bacterial killing (in case of non-adapted pathogens) and by generating more soluble ligands (in case of adapted and non-
adapted pathogens), serving to enhance immune activation, not only locally, but also systemically. Not surprisingly, in \( LYS1^{KD} \) plants, a lack of LYS1 led to compromised plant immunity and resulted in a higher susceptibility to pathogens (Figure 20B).

In wild type plants, once the invading pathogens are successfully warded off, pathogen numbers will decrease accordingly, thus reducing the amount of newly released immunogenic PGN fragments. The high LYS1 levels in the apoplast might continually degrade the soluble PGN fragments into smaller units that do not induce immunity, but only bind to receptors. As a result, receptor stimulation will gradually be attenuated, followed also by a decrease in LYS1 levels back to the un-induced basal levels. Therefore, in \( LYS1^{OE} \) plants where LYS1 is constitutively expressed at high levels, this LYS1 will most likely always over-digest PGN into small fragments, which have no immunogenic activities (Figure 15D). These over-digested short fragments might compete with long immunogenic fragments for binding to their receptors and consequently block the receptor-mediated immune signaling, resulting in a reduced activation of the immune system. Direct killing of bacteria in \( LYS1^{OE} \) lines, however, was not observed and is likely inhibited by adapted pathogens, resulting in a total reduced resistance to adapted pathogens. This model could explain our observation that \( LYS1^{OE} \) plants with high constitutive LYS1 levels displayed an elevated susceptibility to \( PtoDC3000 \) in comparison to wild type plants (Figure 20).

In conclusion, LYS1 levels are fine-tuned in response to external flexible stimuli to defend against pathogens. Inappropriate LYS1 levels will compromise plant immunity as shown in \( LYS1^{KD} \) and \( LYS1^{OE} \) mutant lines. There is another prominent example where overexpression of a protein mimics the effect of genetic inactivation. Strong overexpression of BAK1, a co-receptor of leucine-rich-repeat receptor kinases, triggers inappropriate plant cell death (Belkhadir et al., 2012) as is also observed in \( bak1 \) mutants (Kemmerling et al., 2007). Hence, it can be assumed that wild-type levels of proteins are optimized during evolution and it is therefore not surprising that either too little or too much activity might have a deleterious effect on plant.

These results should also remind us to be aware of possible complications arising from manipulation of protein levels in transgenic plants used in agriculture.
4.6 Known signaling components in CERK1-mediated immunity

Plant LysM-type PRRs recognize GlcNAc-containing ligands, such as fungal chitin, bacterial PGN (PGN), and thus have functions in innate immunity (Böhm et al., 2014). However, downstream signaling events are so far poorly understood.

In rice, upon perceiving chitin, the chitin elicitor binding protein (CEBiP) dimer likely recruits *Oryza sativa* chitin elicitor receptor kinase 1 (*OsCERK1*), which is subsequently (auto)-phosphorylated and thus activates the downstream immune response (Hayafune et al., 2014). In the process, *OsCERK1* phosphorylates *OsRacGEF1* (*Oryza sativa* RAC guanine nucleotide exchange factor 1), which activates the small GTPase *OsRAC1* (Akamatsu et al., 2013). In turn *OsRAC1* was proposed to interact with downstream signaling components and regulate the final steps of the signaling pathways, including the generation of reactive oxygen species, phytoalexins, lignins, the activation of MAPK cascades, and the expression of PR proteins (Akamatsu et al., 2013). In addition, two RLKs, *Oryza sativa* receptor-like cytoplasmic kinase 185 (*OsRLCK185*) and *Oryza sativa* receptor-like cytoplasmic kinase 176 (*OsRLCK176*), were shown to mediate chitin-induced signaling, potentially constituting additional links between PRRs and the MAPK cascades (Yamaguchi et al., 2013; Ao et al., 2014). Finally, two LysM-receptor-like proteins, *OsLYP4* and *OsLYP6*, were reported to heterodimerize with *OsCERK1* upon chitin binding and thereby induce immune responses (Liu et al., 2012a; Ao et al., 2014). Interestingly, *OsRLCK176*, *OsCERK1*, *OsLYP4* and *OsLYP6* do not only respond to chitin, but also have been implicated in PGN responses. Thus, *OsCERK1* most likely acts as an adaptor for signal transduction of multiple LysM-type PRRs and is not solely involved in chitin-induced but also PGN-induce immunity (Ao et al., 2014; Kouzai et al., 2014).

In *Arabidopsis*, it was suggested that chitin oligomers (DP>6) are bound by the LysM domains of two CERK1 monomers, resulting in receptor dimerization and trans-phosphorylation of their cytoplasmic kinase domains (Liu et al., 2012b). This dimerization model is based on relatively long (DP=8), not short (DP=5 or 6), chitin oligomers, despite the findings that short chitin oligomers can also induce chitin-triggered responses via a currently unknown mechanism (Felix et al., 1993; Liu et al.,
2012b). Upon perception of chitin, CERK1 immediately phosphorylates two receptor-like cytoplasmic kinases (RLCKs), BOTRYTIS-INDUCED KINASE1 (BIK1) and PBS1-like protein 27 (PBL27), and then regulates partially overlapping but also different downstream signaling pathways. Both RLCKs were involved in the regulation of MAMP-induced callose deposition, but only BIK1 was involved in MAMP-induced accumulation of ROS. Likewise, only PBL27 was involved in the activation of MAPK cascades (Shinya et al., 2014).

Interestingly, LYM2, which has high affinity binding with chitin, mediates immunity via a CERK1-independent pathway (Faulkner et al., 2013). Recently, LYK5, a kinase-inactive LYK, was proposed to form a chitin-inducible complex with CERK1, which leads to CERK1 phosphorylation and the induction of chitin-activated immunity (Cao et al., 2014). Such a sandwich-like complex model, as similarly proposed for OsCERK1 in rice, would argue that chitin perception systems of rice and Arabidopsis are somehow similar to each other.

4.7 CDPKs in plant immunity

In order to identify further novel components of the chitin- and PGN-induced CERK1-mediated signaling pathways, we searched the M.I.N.D. yeast-two-hybrid database for putative CERK1 interacting proteins. One candidate protein chosen for further analysis was the calcium-dependent protein kinase CPK15.

Upon perception of PAMPs, plant PRRs immediately activate opening of Ca\(^{2+}\) channel in the plasma membrane and induce rapid Ca\(^{2+}\) influx, thus resulting in a rapid increase of cytosolic Ca\(^{2+}\) concentration (Lecourieux et al., 2006). Ca\(^{2+}\) has long been recognized as a conserved second messenger and principal mediator in plant immune and stress signaling through Ca\(^{2+}\) sensors (Boudsocq and Sheen, 2013). Plants possess three main families of calcium sensors: calmodulins (CaM), calcineurin B-like proteins (CBL) and CDPKs. Unlike CaM and CBL that relay the Ca\(^{2+}\)-induced conformational change to protein partners, CDPKs have the unique feature that both Ca\(^{2+}\) sensing and receiver domains are combined within a single protein to directly translate Ca\(^{2+}\) signals into phosphorylation (Boudsocq and Sheen, 2013).
Discussion

CDPKs are encoded by 34 members in *Arabidopsis* and divided into four subgroups (Cheng et al., 2002). Typically, a CDPK harbors an N-terminal variable domain, a Ser/Thr kinase domain, an auto-inhibitory junction region and a regulatory calmodulin-like domain (CaM-LD) (Harper et al., 2004). The CaM-LD is composed of the four EF-hand Ca\(^{2+}\)-binding motifs which have high Ca\(^{2+}\) affinities resulting in Ca\(^{2+}\)-mediated CDPK regulation. In the basal state, the interaction between the auto-inhibitory region and the kinase domain keeps the kinase in an inactive state by a pseudosubstrate mechanism (Harper et al., 2004). The C-terminal lobe of the CaM-LD exhibiting high Ca\(^{2+}\) affinity interacts with the auto-inhibitory region at low Ca\(^{2+}\) levels to stabilize the structure (Weljie and Vogel, 2004). Ca\(^{2+}\) binding to the low-affinity N-terminal lobe of the CaM-LD induces a conformational change that releases the auto-inhibition (Harper et al., 2004), resulting in activation of the protein kinase domain. N-variable domains of CDPKs often have a myristoylation and a palmitoylation posttranslational modification, which determines protein subcellular localization and substrate recognition (Asai et al., 2013). Mutations of N-terminal myristoylation and palmitoylation sites in the N-terminal domain eliminated the predominantly plasma membrane localization and the capacity of StCDPK5 to activate StRBOHB (Respiratory burst oxidase homolog B) *in vivo* (Asai et al., 2013). CPK15 in this study is predicted to have both myristoylation and palmitoylation modifications (Cheng et al., 2002) and could be shown to be localized at the plasma membrane (Figure 27). This localization of CPK15 would facilitate CERK1-interaction to mediate a CERK1-dependent reactive oxygen species (ROS) burst (Figure 25), probably also through association with the NADPH oxidase RBOHD (Respiratory burst oxidase homolog D).

The 34 CDPK members of *Arabidopsis* are cluster into four subgroups (Figure 29). The members in one subgroup often display redundancy along with functional specificity, exemplified by CDPK4/5/6/11 in orchestrating immune gene expression, CDPK1/2/4/11 in ROS production, and CDPK1/2/5/6/ in programmed cell death (PCD) (Gao et al., 2013). The functional specificity of CDPKs is maintained apparently via phosphorylation and activation of different substrates in distinct subcellular compartments. The apparent functional redundancy of closely related CDPKs often challenges genetic investigation of special function. In this study, CDPK15 displayed only a 30-50 % reduction in chitin-
induced ROS accumulation (Figure 23 and 25), which may indicate the presence of functional redundant members such as CDPK21 and CDPK23, the two closest homologs of CDPK15 (Figure 29). It will be interesting to investigate roles of CDPK21 and CDPK23 in chitin-induced signaling.

PAMP perception immediately induces the activation of MAPK and CDPK, leading to expression of defence genes. Unexpectedly, CDPKs and MAPK cascades differentially regulate flg22-induced early genes in at least four regulatory programs (Boudsocq et al., 2010). In this study, CPK15 did not affect MAPK cascades and may thus function independently of MAPK cascades (Figure 26).

Unlike most CDPKs, which are involved in plant immune signaling as positive regulators (Boudsocq et al., 2010), it was reported recently that CPK28 acts as a negative regulator in PTI signaling (Monaghan et al., 2014). CPK28 associates with RBOHD and BIK1, and phosphates the latter, an important convergent substrate of multiple PRR complexes such as CERK1 and FLS2 (Zhang et al., 2010; Kadota et al., 2014). In this study, CPK15 is involved in chitin- but not flg22-induced ROS (Figure 25). Thus we postulate that CPK15 should be placed upstream of BIK1 to specifically act in CERK1-mediated PTI signaling. Indeed, our results showed that CPK15 directly interacts with CERK1 (Figure 28). Further investigation of the possible association between CPK15 and BIK1 or between CPK15 and RBOHD will be very interesting.

Taken together, CPK15 directly targets CERK1 and positively regulates CERK1-mediated PTI signaling in Arabidopsis.
5. Summary

As one part of a two-tiered pathogen-detection system in plants, PTI is not only sufficient to ward off most microbes, but also contributes to basal immunity during infection.

Although PGN and the LYM1-LYM3-CERK1 receptor complex have been identified as one PAMP-PRR pair in *Arabidopsis*, upstream and downstream events of PGN perception remain elusive. Specifically, it is unclear whether and, if so, how PGN supermolecules are processed prior to perception by the LYM1-LYM3-CERK1 receptor complex. Moreover, the mechanisms linking PGN perception with diverse downstream immune responses are little understood.

In this study, a lysozyme-like hydrolase (lysozyme 1, LYS1) was identified as an enhancer in PGN-induced immunity. Upon bacterial infection or exposure to bacterial patterns, *Arabidopsis* produces LYS1 to release soluble PGN fragments from insoluble bacterial cell walls. LYS1-released soluble PGNs trigger typical immunity-associated responses, such as medium alkalinization and up-regulation of resistance-related genes. Importantly, these immune responses are dependent on the PGN receptor complex. LYS1 mutants exhibit super-susceptibility to bacterial infection similar to that observed in PGN receptor mutants. We propose that plants employ hydrolytic activities for the decomposition of complex bacterial structures and that the subsequent generation of soluble patterns might aid PRR-mediated immune activation in cell layers adjacent to infection sites.

In addition, we identified the calcium-dependent protein kinase CPK15 as an interactor of CERK1, which is not only involved in PGN perception but also in chitin recognition. CPK15 was shown to interact with CERK1 in the yeast-two-hybrid system and in plant tissue, and *cpk15* mutants displayed reduced ROS accumulation upon chitin treatment. These findings indicate that CPK15 is involved in CERK1-mediated PTI signaling.

Summing up, this study aimed at improving our understanding of PGN- and chitin-triggered immunity by identifying and characterizing critical components of the plant immune system, such as LYS1, involved in upstream events of PGN perception, and CPK15, involved in downstream events of CERK1-mediated glucan perception.
6. Zusammenfassung

Als ein Teil eines zweistufigen pflanzlichen Pathogen-Erkennungssystems ist die PAMP-getriggerte Immunität (PTI) nicht nur ausreichend zur Abwehr der meisten Mikroben, sondern trägt auch zur basalen Immunität während der Infektion bei.

Obwohl PGN und der LYM1-LYM3-CERK1 Rezeptorkomplex als ein PAMP/PRR-Paar in *Arabidopsis* identifiziert worden sind, sind sowohl die vor- als auch die nachgeschalteten Ereignisse bei der PGN-Erkennung noch wenig untersucht. Insbesondere ist unklar, ob, und wenn ja, wie PGN-Makromoleküle vor ihrer Perzeption durch den LYM1-LYM3-CERK1 Rezeptorkomplex prozessiert werden. Darüber hinaus ist wenig über die Mechanismen bekannt, die die PGN-Erkennungsproteine mit den verschiedenen nachgeschalteten Immunreaktionen verbindet.


Zusammenfassend sollte diese Studie dazu beitragen, unser Verständnis der PGN-ausgelösten Immunität zu verbessern. So wurden hier kritische Komponenten des pflanzlichen Immunsystems identifiziert und charakterisiert, wie zum einen LYS1, welches an der PGN-Wahrnehmung vor Rezeptorbindung beteiligt ist, und zum anderen CPK15, welche wichtig für die durch CERK1 vermittelte Signalweiterleitung nach Glucan-Wahrnehmung ist.
7. Reference


References


Fliegmann, J., Mithofer, A., Wanner, G., and Ebel, J. (2004). An ancient enzyme domain hidden in the putative beta-glucan elicitor receptor of soybean may play an active part in the perception of pathogen-associated molecular patterns during broad host resistance. The Journal of biological chemistry 279, 1132-1140.


References


SUPPRESSOR OF BIR1-1/EVERSHELD Mediate Innate Immunity to Necrotrophic Fungi. The Plant cell 25, 4227-4241.


8. Appendix.

Table 9. Oligonucleotides used in this study

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<td>59,3</td>
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9 Acknowledgements

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10 Curriculum vitae

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Publications


