Characterisation of the Role of LysM Receptor-Like Kinases and the CHIA Chitinase in the Perception of Peptidoglycan and in the Innate Immunity of Arabidopsis thaliana

Dissertation

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Heini Marjatta Grabherr
geb. Lajunen
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1. Berichterstatter: Prof. Dr. Thorsten Nürnberger
2. Berichterstatter: Prof. Dr. Georg Felix
Table of contents

1 Introduction ........................................................................................................... 1
  1.1 Host-pathogen interaction ................................................................................ 1
  1.2 Plant innate immunity ..................................................................................... 1
    1.2.1 PAMP-triggered immunity ........................................................................ 1
    1.2.2 Effector-triggered immunity ..................................................................... 4
    1.2.3 Chitinases in plant pathogen defense ..................................................... 5
  1.3 Peptidoglycan - A bacterial cell wall constituent ......................................... 6
  1.4 PGN processing and perception in animals ............................................... 9
  1.5 Lysin motif (LysM) as mediator in carbohydrate signaling in plants .......... 14
  1.6 Aims of the thesis ......................................................................................... 15

2 Materials and methods ...................................................................................... 16
  2.1 Materials ....................................................................................................... 16
    2.1.1 Chemicals ............................................................................................... 16
    2.1.2 Media ..................................................................................................... 16
    2.1.3 Vectors .................................................................................................. 18
    2.1.4 Primers ................................................................................................. 18
  2.2 Organisms ...................................................................................................... 19
    2.2.1 Bacteria and fungi .................................................................................. 19
    2.2.2 Arabidopsis thaliana lines ...................................................................... 19
  2.3 Cultivation conditions of the organisms ...................................................... 20
    2.3.1 Growth of Escherichia coli .................................................................... 20
    2.3.2 Growth of Pseudomonas syringae ........................................................ 21
    2.3.3 Growth of Agrobacterium tumefaciens ................................................. 21
    2.3.4 Growth of Pichia pastoris ...................................................................... 21
    2.3.5 Growth of Alternaria brassicicola and Botrytis cinerea ....................... 21
    2.3.6 Growth of Arabidopsis thaliana and Nicotiana benthamiana ............ 21
  2.4 Methods ......................................................................................................... 22
    2.4.1 Isolation of peptidoglycan from Gram-negative bacteria .................... 22
    2.4.2 Isolation of peptidoglycan from Gram-positive bacteria ..................... 23
    2.4.3 General molecular biology methods ..................................................... 23
    2.4.4 Cloning .................................................................................................. 23
    2.4.5 DNA isolation ........................................................................................ 24
    2.4.6 RNA isolation ......................................................................................... 24
    2.4.7 Semi-quantitative RT-PCR .................................................................... 24
    2.4.8 Quantitative Real-time PCR .................................................................. 24
    2.4.9 Isolation of mesophyll protoplasts from Arabidopsis ......................... 25
    2.4.10 Stable transformation of Arabidopsis thaliana ..................................... 25
    2.4.11 Transient transformation of Nicotiana benthamiana ......................... 25
    2.4.12 Crossing Arabidopsis thaliana plants .................................................. 26
    2.4.13 Generation of knock-down lines ......................................................... 26
Table of contents

2.4.14 Generation of overexpression lines ....................................................... 27
2.4.15 Generation of pCHIA::GUS lines ......................................................... 27
2.4.16 Generation of constructs for expression in E.coli .................................. 28
2.4.17 Generation of constructs for expression in P.pastoris ......................... 28
2.4.18 Generation of constructs for the yeast two-hybrid system ................. 28
2.4.19 Genotyping analysis of T-DNA insertion lines ..................................... 28

2.5 Biochemical methods ............................................................................. 29
2.5.1 Protein expression in E.coli ................................................................... 29
2.5.2 Ni\textsuperscript{2+}-NTA affinity purification ........................................ 30
2.5.3 Protein expression in P.pastoris ........................................................... 30
2.5.4 Protein extraction from plant tissue .................................................... 30
2.5.5 Immunoprecipitation ........................................................................... 31
2.5.6 Determination of protein concentration ............................................. 31
2.5.7 SDS-PAGE .......................................................................................... 31
2.5.8 Western blot analysis ........................................................................... 31
2.5.9 Coomassie blue stain .......................................................................... 32
2.5.10 Silver stain .......................................................................................... 32
2.5.11 Turbidity assay (PGN-hydrolysis assay) ............................................ 33
2.5.12 4-MUCT assay (Chitin-hydrolysis assay) ......................................... 33
2.5.13 Colloidal chitin hydrolysis assay ....................................................... 33
2.5.14 Yeast two-hybrid ................................................................................ 34

2.6 Bioassays ................................................................................................ 34
2.6.1 Infection with Pseudomonas syringae ............................................... 34
2.6.2 Infection with Alternaria brassicicola .................................................. 34
2.6.3 Infection with Botrytis cinerea ............................................................. 35
2.6.4 Elicitation assays in leaves or seedlings .............................................. 35
2.6.5 Microarray analysis ............................................................................. 35
2.6.6 pH assay .............................................................................................. 36

2.7 Microscopy and Histochemistry ............................................................... 36
2.7.1 Confocal microscopy ........................................................................... 36
2.7.2 Aniline blue stain ................................................................................ 36
2.7.3 Trypan blue stain ................................................................................. 36
2.7.4 GUS stain ............................................................................................ 37

2.8 Statistical analysis ................................................................................... 37

3 Results ..................................................................................................... 38
3.1 Isolation and analysis of PGN from P.syringae and other bacteria .......... 38
3.2 Identification of putative PGN receptor(s) among the LysM-RLKs ......... 41
3.2.1 Analysis of the LYK T-DNA insertion lines ......................................... 43
3.2.1.1 Phenotypic analysis of lyk3 and lyk5 single and double mutants ... 45
3.2.2 Role of LYKs in fungal resistance ....................................................... 47
3.2.3 Influence of LYKs on bacterial resistance ......................................... 49
3.2.4 Analysis of peptidoglycan responsiveness in lyk mutants ............... 51
3.2.4.1 The LysM-receptor kinase CERK1 mediates sensitivity to PGN .... 52
3.2.5 Analysis of potential redundancy among LYK genes .................. 55
3.3 The role of PGN hydrolysis in the PGN sensing process ........... 56
3.4 Identification of a putative PGN hydrolase in Arabidopsis thaliana .... 57
  3.4.1 Analysis of the class III chitinase CHIA .................................. 58
  3.4.2 Expression pattern of the CHIA gene upon biotic stress ........... 60
  3.4.3 Analysis of the transgenic CHIA lines ..................................... 64
    3.4.3.1 Phenotypic analysis of the CHIA overexpression and knock-down
    lines .......................................................................................... 66
  3.4.4 Subcellular localisation of CHIA .............................................. 67
  3.4.5 Detection of CHIA protein in the transgenic CHIA lines .......... 71
  3.4.6 Posttranslational modification of the CHIA protein ................. 72
  3.4.7 Expression of CHIA protein using heterologous expression systems ... 73
  3.4.8 Analysis of homo-oligomerisation properties of CHIA .......... 76
  3.4.9 Assessment of the chitin- and PGN-hydrolase activity of CHIA ...... 77
  3.4.10 Role of CHIA in fungal resistance .......................................... 82
    3.4.10.1 Infection of the transgenic CHIA lines with Botrytis cinerea ....... 82
    3.4.10.2 Infection of the transgenic CHIA lines with Alternaria brassicicola . 83
  3.4.11 Impact of CHIA on bacterial resistance ................................... 86
4 Discussion ....................................................................................... 88
  4.1 LYKs contribute to plant innate immunity .................................. 88
    4.1.1 Effects of LYK gene deletions in plant fungal and bacterial resistance . 88
    4.1.2 CERK1 serves together with LYM3 peptidoglycan recognition .......... 90
  4.2 CHIA chitinase is involved in plant innate immunity ...................... 93
    4.2.1 Chitinolytic activity of CHIA and its impact on fungal immunity .......... 94
    4.2.2 PGN-lytic activity of CHIA and its importance to bacterial immunity .... 95
  4.3 Peptidoglycan perception systems have arisen through convergent evolution in
     metazoans and plants ...................................................................... 97
5 Summary .......................................................................................... 99
6 Zusammenfassung ........................................................................... 101
7 References ....................................................................................... 103
8 Appendix .......................................................................................... i
### Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>aa</td>
<td>Amino acid</td>
</tr>
<tr>
<td>amiRNA</td>
<td>Artificial microRNA</td>
</tr>
<tr>
<td>At</td>
<td>Arabidopsis thaliana</td>
</tr>
<tr>
<td>Avr</td>
<td>Avirulence factor</td>
</tr>
<tr>
<td>bp</td>
<td>Base pair</td>
</tr>
<tr>
<td>cDNA</td>
<td>Complementary DNA</td>
</tr>
<tr>
<td>cfu</td>
<td>Colony forming unit</td>
</tr>
<tr>
<td>Col-0/Col-2</td>
<td>Columbia-0/Columbia-2, ecotypes</td>
</tr>
<tr>
<td>DAMP</td>
<td>Danger-associated molecular pattern</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleid acid</td>
</tr>
<tr>
<td>EF-Tu</td>
<td>Elongation factor Tu</td>
</tr>
<tr>
<td>Elf18</td>
<td>Peptide from EF-Tu with the sequence SKEFERTKPHVNVGTIG</td>
</tr>
<tr>
<td>ETI</td>
<td>Effector-triggered immunity</td>
</tr>
<tr>
<td>ETS</td>
<td>Effector-triggered susceptibility</td>
</tr>
<tr>
<td>Flg22</td>
<td>Peptide from flagellin with the sequence QRLSTGSRINSAKDDAAGLQIA</td>
</tr>
<tr>
<td>FLS2</td>
<td>Flagellin-sensitive 2</td>
</tr>
<tr>
<td>GFP</td>
<td>Green fluorescent protein</td>
</tr>
<tr>
<td>HEWL</td>
<td>Hen egg-white lysozyme</td>
</tr>
<tr>
<td>HR</td>
<td>Hypersensitive reaction</td>
</tr>
<tr>
<td>kb</td>
<td>Kilobase</td>
</tr>
<tr>
<td>kDa</td>
<td>KiloDalton</td>
</tr>
<tr>
<td>KO</td>
<td>Knock-out</td>
</tr>
<tr>
<td>Ler</td>
<td>Landsberg erecta, ecotype</td>
</tr>
<tr>
<td>LRR</td>
<td>Leucine-rich repeat</td>
</tr>
<tr>
<td>LPS</td>
<td>Lipopolysaccharide</td>
</tr>
<tr>
<td>LysM</td>
<td>Lysin motif</td>
</tr>
<tr>
<td>MAMP</td>
<td>Microbe-associated molecular pattern</td>
</tr>
<tr>
<td>MAPK/MPK</td>
<td>Mitogen-activated protein kinase</td>
</tr>
<tr>
<td>Os</td>
<td>Oryza sativa</td>
</tr>
<tr>
<td>PAMP</td>
<td>Pathogen-associated molecular pattern</td>
</tr>
<tr>
<td>PCD</td>
<td>Programmed cell-death</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>PGN</td>
<td>Peptidoglycan</td>
</tr>
<tr>
<td>PGRP</td>
<td>Peptidoglycan recognition protein</td>
</tr>
<tr>
<td>PR</td>
<td>Pathogenesis-related</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>Pseudomonas syringae pv. tomato</td>
</tr>
<tr>
<td>pv.</td>
<td>Pathovar</td>
</tr>
<tr>
<td>R-Gene</td>
<td>Resistance gene</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
</tr>
<tr>
<td>R-Protein</td>
<td>Resistance protein</td>
</tr>
<tr>
<td>RLK</td>
<td>Receptor-like kinase</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive oxygen species</td>
</tr>
<tr>
<td>SA</td>
<td>Salicylic acid</td>
</tr>
<tr>
<td>SAR</td>
<td>Systemic acquired resistance</td>
</tr>
<tr>
<td>T-DNA</td>
<td>Transfer-DNA</td>
</tr>
<tr>
<td>TLR</td>
<td>Toll-like receptor</td>
</tr>
<tr>
<td>UTR</td>
<td>Untranslated region</td>
</tr>
<tr>
<td>WT</td>
<td>Wild type</td>
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</tbody>
</table>
List of Figures

Figure 1-1: Plant pattern-recognition receptors and their signaling adapters...........................3
Figure 1-2: The schematic structure of peptidoglycan .............................................................7
Figure 1-3: Gram-positive and Gram-negative cell walls .........................................................8
Figure 1-4: Peptidoglycan sensing in animals ........................................................................11
Figure 1-5: Metazoan PGN hydrolytic activities .....................................................................13
Figure 2-1: Working mode of amiRNA-mediated gene silencing ...........................................27
Figure 2-2: Genotyping analysis of T-DNA lines by gene- and insertion-specific primer ......29
Figure 3-1: Peptidoglycan induces defense responses in *Arabidopsis thaliana* ......................40
Figure 3-2: *Arabidopsis* LysM receptor-like kinase family ....................................................42
Figure 3-3: Characterisation of CERK1, LYK3 and LYK5 and their mutants ............................44
Figure 3-4: Characterisation of LYK2 and LYK4 and their mutants .......................................45
Figure 3-5: Phenotypes of *lyk3* and *lyk5* single and double mutants .................................46
Figure 3-6: Infection of *lyk* mutants with *Botrytis cinerea* ................................................48
Figure 3-7: Infection of *lyk* mutants with *Pseudomonas syringae* strains ............................50
Figure 3-8: Analysis of PGN responsiveness in *lyk* mutants ..............................................52
Figure 3-9: Functional characterisation of PGN-induced genes by GO analysis .....................53
Figure 3-10: CERK1 mediates PGN sensitivity .....................................................................54
Figure 3-11: Generation of triple *lyk* knock-out mutants ....................................................55
Figure 3-12: Muropeptide-induced gene expression requires LYM3 and CERK1 .................57
Figure 3-13: Sequence alignment of the *Arabidopsis* chitinases .........................................58
Figure 3-14: CHIA protein sequence and annotated features ...............................................59
Figure 3-15: CHIA expression profile upon biotic stress ......................................................61
Figure 3-16: CHIA expression analysis using a *pCHIA::GUS* reporter line .........................63
Figure 3-17: Characterisation of CHIA T-DNA insertion lines ..............................................64
Figure 3-18: Characterisation of CHIA knock-down and overexpression lines ....................66
Figure 3-19: Phenotypes of CHIA overexpression and knock-down lines .............................67
Figure 3-20: Localisation of CHIA-GFP in *Arabidopsis* ......................................................68
Figure 3-21: Localisation of CHIA-GFP and CHIAΔSP-GFP in tobacco .................................70
Figure 3-22: Detection of CHIA protein in *Arabidopsis* ........................................................72
Figure 3-23: Deglycosylation of CHIA-GFP and CHIAΔSP-GFP ............................................73
Figure 3-24: Expression of His6-CHIA in *Escherichia coli* ................................................75
Figure 3-25: Expression of CHIA in *Pichia pastoris* ............................................................76
Figure 3-26: Yeast two-hybrid analysis ..................................................................................77
Figure 3-27: Chitin- and PGN-hydrolysis activity of CHIA leaf protein ...............................79
Figure 3-28: Chitin- and PGN-hydrolysis activity of CHIA protoplast samples ....................81
Figure 3-29: Infection of the transgenic CHIA lines with \textit{Botrytis cinerea} .........................83
Figure 3-30: Infection of the transgenic CHIA lines with \textit{Alternaria brassicicola} ..................85
Figure 3-31: Infection of the transgenic CHIA lines with \textit{Pseudomonas syringae} ..................87
Figure 4-1: Plant perception and signaling of carbohydrate PAMPs/MAMPs .........................92
Figure 8-1: ClustalW2 protein sequence alignment of CHIA (At5g24090) and selected plant class chitinases ......................................................... iv

\section*{List of Tables}

Table 2-1: Used media ...........................................................................................................17
Table 2-2: Used antibiotics ...................................................................................................17
Table 2-3: Used vectors .......................................................................................................18
Table 2-4: Used bacterial strains ..........................................................................................19
Table 2-5: Knock-out and complementation lines used in this work ....................................20
Table 8-1: Used oligonucleotides ....................................................................................... iii
1 Introduction

1.1 Host-pathogen interaction

Plants, animals and other organisms are continuously confronted with potential disease-causing pathogens. Host organisms provide the successful invaders nutrients, protection, convenient transmission routes or a platform for propagation. Luckily, the host barriers are seldomly overcome by the pathogenic microbes due to effective defense mechanisms. Animals rely on two types of surveillance and defense systems. The innate immunity renders animals a first line of protection against many pathogens due to the recognition of conserved microbial signatures by pattern recognition receptors (PRRs) mediating host inflammatory responses. The second line of defense is brought about by the adaptive immune system, which is comprised of specialised immune cells and immunoglobulins with immense diversity and specificity for macromolecules derived from invading pathogens. Invertebrates like *Drosophila* and *Caenorhabditis* are solely dependent on the innate immunity, suggesting this immune system to be more ancient than that of jawed vertebrates, which rely on both adaptive and innate immunity (Medzhitov and Janeway, 2000).

1.2 Plant innate immunity

Although plants lack the recombinatorial adaptive immunity known from the animal kingdom, also they possess an effective immune system. The basis of the plant innate immunity is the ability to differentiate between self and non-self or modified self. Furthermore, the plant innate immune system is branched into two forms of immunity, termed pathogen-associated molecular pattern (PAMP)-triggered immunity (PTI) and effector-triggered immunity (ETI), formerly also known as basal disease resistance and resistance (R) gene-based disease resistance, both aiming at a successful restriction of pathogen growth to defeat pathogenic attack (Abramovitch et al., 2006; Chisholm et al., 2006; Jones and Dangl, 2006).

1.2.1 PAMP-triggered immunity

PAMPs or MAMPs (microbe-associated molecular patterns) are highly conserved molecules and often essential for the fitness and survival of the microbial organisms (Nürnberger and Brunner, 2002). They include for instance proteinaceous signatures such as bacterial flagellin, elongation factor Tu (EF-Tu), a 13 aa-fragment of a *Phytophthora* transglutaminase (Pep-13) and cell wall components like fungal chitin, oomycete heptaglucan and bacterial lipopolysaccharides and peptidoglycan (Cosio et al., 1990; Dow et al., 2000; Erbs et al.,
The perception of such non-self structures is mediated by cell surface PRRs (Figure 1-1), which transduce the signal into the plant cell ultimately triggering a generic, broad-range defense response leading to PTI. Many of the PRRs identified so far belong to the large receptor-like kinase (RLK)/Pelle gene family (Shiu and Bleecker, 2001). They typically harbour an extracellular domain dedicated for ligand perception and possibly protein-protein interaction, a transmembrane domain and a cytoplasmic protein kinase domain. Motifs found within the extracellular domain vary, a prominent example is the leucine-rich repeat (LRR) domain present in two well-studied receptor proteins, FLS2 (FLAGELLIN-SENSING 2) and EFR (EF-Tu receptor) (Gómez-Gómez and Boller, 2000; Zipfel et al., 2006). The LRR domain is also a feature of some of the animal PRRs, the Drosophila Toll and the mammalian Toll-like receptors (TLRs) (Imler and Hoffmann, 2001; Medzhitov et al., 1997). Intriguingly, the perception of bacterial flagellin is mediated through LRR domain-containing receptors both in plants and in animals by FLS2 and TLR5, respectively (Gómez-Gómez and Boller, 2000; Hayashi et al., 2001). However, the extracellular LRR domains of the two proteins and also the flagellin epitopes recognised by these receptors differ greatly suggesting that the perception systems are the product of convergent evolution. The PRR-dependent activation of the basal immune reactions truly contributes to resistance towards pathogens. For instance, the depletion of FLS2 was shown to lead to enhanced bacterial susceptibility and the mutant line lacking the gene encoding the chitin elicitor receptor kinase 1 (CERK1) was less resistant towards fungal pathogens than wild type plants (Miya et al., 2007; Wan et al., 2008; Zipfel et al., 2004).
Figure 1-1: Plant pattern-recognition receptors and their signaling adapters

The bacterial PAMPs flagellin (flg22) and elongation factor Tu (EF-Tu) are perceived by the Arabidopsis LRR receptor kinases FLS2 and EFR, respectively. FLS2 and EFR form hetero-oligomers with the coreceptor BAK1 (the BRI1-associated kinase 1) in a ligand-dependent manner. AtCERK1 mediates the recognition of an so far unknown bacterial PAMP and is also essential for chitin perception and fungal resistance. In rice, OsCERK1 acts together with the CEBiP protein in the chitin detection. In tomato, xylanase is detected by the RLPs LeEIX1 and LeEIX2. In leguminous plants, oomycete heptaglucan (HG) is bound by the glucan-binding protein (GBP), which subsequently activates the plant immune responses in a receptor-dependent manner. The AtPep peptides act as danger-associated molecular patterns (DAMPs) and are recognised by the LRR-RLK PEPR1. From Zipfel (2009).

The PRR-based recognition of microbial signatures generates a battery of basal defense reactions in plants. One of the very early responses is the change of ion fluxes across the plasma membrane. Thereby, the influx of Ca\(^{2+}\) and H\(^{+}\) and the efflux of K\(^{+}\) and anions like chloride and nitrate are increased (Boller, 1995; Nürnberger et al., 2004; Wendehenne et al., 2002). Experimental proof for the role of Ca\(^{2+}\) as an intracellular second messenger activating calcium-dependent protein kinases (CDPKs) and membrane channels has been provided (Blume et al., 2000; Brunner et al., 2002; Lecourieux et al., 2002; Ranf et al., 2008). Additional early responses are the generation of reactive oxygen species (ROS) and the activation of mitogen-activated protein kinases (MAPKs) (Apel and Hirt, 2004; Asai et al., 2002; Rodriguez Suarez et al., 2010). The MAPK signaling cascade culminates in the activation of transcription factors, such as members of the WRKY family, leading ultimately to induction of defense-responsive genes. Among these genes are for instance camalexin biosynthesis genes and genes encoding antimicrobial enzymes or receptor proteins (Boudsocq et al., 2010; Gust et al., 2007; Miya et al., 2007; Zipfel et al., 2006; Zipfel et al., 2004). The changes in the transcriptome induced by different PAMPs strongly overlap.
indicating that the PRR pathways are at some point merging thereby giving rise to a generic plant immune response. An additional defense response is the accumulation of β-1,3-glucan (callose) deposits at the plant cell periphery (Flors et al., 2005; Luna et al., 2011). Callose deposition is detectable only after several hours upon infection and thus belongs to the so-called late immune responses.

In addition to non-self signals also signals resulting from altered host molecules can induce the primary plant defense response. Such danger-associated molecular patterns (DAMPs) are released and sensed upon pathogen attack (Hückelhoven, 2007). Among active DAMPs are polysaccharides like oligogalacturonides (OGAs) (D’Ovidio et al., 2004) and cutin monomers (Schweizer et al., 1996), which are released from the plant cell wall, but also endogenous peptides. The *Arabidopsis* peptide *At*Pep1 acts as a DAMP molecule and induces upon perception by the PEPR1 and PEPR2 receptors defense gene expression and also the induction of the *At*Pep1 precursor protein (Huffaker and Ryan, 2007; Krol et al., 2010; Ryan et al., 2007; Yamaguchi et al., 2006).

Up to date, many more PTI-triggering structures, such as peptidoglycans (see chapter 1.3), have been characterised than corresponding plant receptors, leaving intriguing gaps in the knowledge of PAMP perception and PTI.

### 1.2.2 Effector-triggered immunity

Virulent pathogens have found ways to overcome the first inducible layer of defense, the PTI, in the plant. Phytopathogenic Gram-negative bacteria, for instance, use a sophisticated secretion system to smuggle a complete battery of proteins, called effectors, into the host cell. One of the best studied bacterial protein-secretion systems is the type III secretion system (T3SS). The T3SS forms a macromolecular infection apparatus with more than 20 subunits, that is able to rupture the plant cell surface (Büttner and Bonas, 2006). Once in the host cytoplasm the effectors start to disable the defense mechanisms and manipulate the host metabolism resulting in profit for the pathogen (Abramovitch et al., 2006; Chisholm et al., 2006). The *P. syringae* effector proteins AvrPto, AvrPtoB and AvrRpt2 suppress responses induced by PAMPs or MAMPs by inhibiting proteins involved in the PTI pathway, such as PRR receptors (Hauck et al., 2003; He et al., 2006; Kim et al., 2005). The toxin coronatine mimicks the plant hormone jasmonate and can reverse the MAMP-induced stomatal closure thus allowing pathogenic bacteria to gain entry into the host (Melotto et al., 2006). As a result of the effector activities the plant resistance mechanisms are impaired, and the pathogen can proliferate. This phenomenon is termed effector-triggered susceptibility (ETS) (Jones and Dangl, 2006). The effectors and more importantly their action in the host
cell served, however, as a new and more pathogen-specific surveillance platform mediated by specific host disease resistance (R) proteins. Most of these R proteins are cytoplasmic and contain a nucleotide binding (NB) and an LRR domain (NB-LRR proteins) (Caplan et al., 2008). The recognition of an effector or its activity by a NB-LRR protein triggers an efficient, prolonged defense response accompanied by a programmed cell death leading renewed to immunity (effector-triggered immunity, ETI) (Jones and Dangl, 2006). In tomato, the host Pto kinase mediates the association between the effector AvrPto and the host NB-LBB receptor protein Prf leading to successful defense (Mucyn et al., 2006). Also AvrPtoB can be recognised by the tomato Pto, which upon phosphorylation inactivates its E3 ligase activity needed for virulence, and subsequently signals ETI through Prf (Ntoukakis et al., 2009). Due to natural selection pathogens able to evade the ETI are favored and so the arms race between plants and pathogenic microbes continues.

1.2.3 Chitinases in plant pathogen defense

Bacterial and fungal chitinases often have housekeeping functions in nutrition processes or within morphogenesis of the cell wall (Cohen-Kupiec and Chet, 1998). In contrary, the plant and animal chitinases mainly play a role in the host self-defense against pathogen attack (Cohen-Kupiec and Chet, 1998; Kasprzewska, 2003; Patil et al., 2000). Therefore, plant chitinases are grouped among other defense-related enzymes and proteins into the large family of pathogenesis-related (PR) proteins (van Loon et al., 2006). Pathogens often try to enter the plants via natural openings, like stomata and hydathodes, hence the apoplastic space is an ancient battleground for plant-pathogen interactions. In addition to other host PR proteins also the acidic chitinase forms are secreted into the apoplast (Sahai and Manocha, 1993). The basic chitinases accumulate mainly in the vacuole. The role of chitinases as antimicrobial proteins is supported by experimental data showing that the expression of many chitinases is induced upon infection with fungal pathogens (Majeau et al., 1990; Samac and Shah, 1991). Moreover, chitinases have been reported to degrade fungal cell walls and inhibit fungal growth in vitro, especially when combined with β-1,3 glucanases (Arlorio et al., 1992; Mauch et al., 1988; Schlumbaum et al., 1986). The first chitinase gene isolated in Arabidopsis encodes the basic chitinase ATHCHIB (At3g12500), which could inhibit growth of Trichoderma reesei in vitro (Samac et al., 1990). The pathogen-induced expression of ATHCHIB was shown to be ethylene-dependent (Thomma et al., 1999a). Interestingly, pathogens have evolved counter attack mechanisms to evade the antimicrobial activity of plant chitinases and so promote virulence. One example for such a mechanism is the production of chitinase-specific inhibitors (Misas-Villamil and van der Hoorn, 2008). Moreover, the Cladosporium fulvum effector Avr4 was reported to protect the fungal cell wall
against chitinolytic degradation by masking the cell wall chitin, suggesting that the chitin-binding properties of Avr4 are also part of a counter-defensive arsenal of the fungi (van den Burg et al., 2006). Another fungal effector, Ecp6, inhibits as scavenger the release of free chitin fragments needed for PTI-activation (de Jonge et al., 2010). Chitinases (EC.3.2.1.14) cleave the glycosidic \( \beta(1\rightarrow4) \) bond in biopolymers of N-acetylglucosamine, present mainly in chitin. Chitin is the major building block of fungal cell walls and also of the exoskeleton of insects and crustaceans. According to sequence similarity of the catalytic glycosyl hydrolase domains, chitinases are divided in families 18 and 19 (Henrissat, 1991). Family 18 chitinases are widely distributed across the kingdoms and are present in bacteria, fungi, viruses, plants and animals, whereas family 19 chitinases are almost exclusively found in plants. The chitinases of both families differ in their biochemical features; family 18 chitinases employ a retention mechanism (catalysis product has the same configuration as the substrate) and family 19 chitinases use an inversion mechanism changing the configuration form of the catalysis product (Brameld and Goddard, 1998; van Aalten et al., 2001). The protein structure of chitinases contains a signal sequence for secretion, a glycosyl hydrolase domain and sometimes an additional chitin-binding domain (Passarinho and De Vries, 2002).

1.3 Peptidoglycan - A bacterial cell wall constituent

Virtually all bacteria have a cell wall, which upholds the cell shape and provides a rigid exoskeleton, a so-called sacculus, protecting the bacterial cell against mechanical and osmotic lysis (Nanninga, 1998; Navarre and Schneewind, 1999). Besides its protective function and shape formation, the cell wall also provides an interface for interactions with the surrounding environment and possible hosts. The most important component of the bacterial cell wall conferring strength and rigidity is the heteropolymeric macromolecule peptidoglycan (PGN). It consists of firm glycan chains that are interlinked either directly or via short peptide bridges (Glauner et al., 1988; Schleifer and Kandler, 1972). The disaccharide building block establishing the glycan chains is made up of N-acetylglucosamine (GlcNAc) and N-acetyl muramic acid (MurNAc) in \( \beta(1\rightarrow4) \) linkage (Figure 1-2). The D-lactyl group of MurNAc provides a possibility for the attachment of the stem peptides via amide linkage. Whereas the glycan chain displays surprisingly little divergence among different bacterial species, the amino acid composition of the peptide bridges can vary. Depending on the presence of L-lysine (Lys) or meso-diaminopimelic acid (Dap) at the third position in the stem peptide...
(Figure 1-2) the peptidoglycan is termed either Lys-type or Dap-type. Lys-type PGN peptides are usually interconnected by a peptide bridge and the Dap-type PGN peptides are directly crosslinked (Navarre and Schneewind, 1999; Schleifer and Kandler, 1972).

![Figure 1-2: The schematic structure of peptidoglycan](image)

Bacterial peptidoglycan is composed of alternating N-acetylglucosamine (NAG, GlcNAc) and N-acetylmuramic acid (NAM, MurNAc) molecules, which form the glycan strands, and stem peptide units which connect neighboring glycan strands in some bacteria, via a peptide crossbridge, with each other. The NAG and NAM sugars are linked with each other by β-1,4 linkage. DA stands for diamino acid (generally diaminopimelic acid or L-lysine) and n for the number of amino acids in the cross-bridge (n=0 to 5 depending on the bacteria). Based on van Heijenoort (2001).

Most Gram-positive bacteria contain the Lys-type peptidoglycan, whereas the Dap-type peptidoglycan is typical for Gram-negative bacteria like *Pseudomonas syringae* and *Escherichia coli*. Gram-positive and Gram-negative bacteria differ not only in the type of peptidoglycan they harbor, but also in the PGN amount. Gram-positive cell walls contain a thick multilayered PGN coat (20 - 80 nm), which is embedded with teichoic and lipoteichoid acids and proteins (see Figure 1-3). Instead, Gram-negative cell walls consist of only few layers of peptidoglycan (1-7 nm) and an additional membrane, the outer membrane (Cabeen
and Jacobs-Wagner, 2005). These two cell wall components are connected with each other via lipoproteins and contain also lipopolysaccharide (LPS) molecules (Figure 1-3).

![Illustration of Gram-positive and Gram-negative cell walls](image)

**Figure 1-3: Gram-positive and Gram-negative cell walls**

A schematic picture shows the components of the cell wall of Gram-positive (left) and Gram-negative (right) bacteria. Left: The Gram-positive cell wall is composed of a thick PGN layer outside of the cytoplasmic membrane. Teichoic and lipoteichoic acids are embedded into the peptidoglycan. Right: The cell wall of Gram-negative bacteria consists of an outer membrane linked by lipoproteins to thin, mostly single-layered PGN in the periplasmic space. In addition to lipoproteins, also porins and lipopolysaccharides are present in the cell wall. Modified after Cabeen and Jacobs-Wagner (2005).

The growth of the bacterial peptidoglycan sacculus takes place by insertion of disaccharide pentapeptide subunits into the existent peptidoglycan and is mediated by the penicillin-binding proteins (PBPs) (Höltje, 1998; Macheboeuf et al., 2006; Nanninga, 1998). To allow the extension of this covalent structure peptidoglycan hydrolase activity is required. This is accomplished by peptidoglycan (murein) hydrolases, which are divided into groups according to their enzymatic properties (Shockman and Höltje, 1994). N-acetylmuramidases (EC 3.2.1.17) and N-acetylglucosaminidases (EC 3.2.1.96) cleave the glycosidic bond behind either the MurNAc or GlcNAc sugar, respectively (Tipper et al., 1964). N-Acetylmuramoyl-L-Ala amidases (amidases, EC 3.5.1.28) are specialised in the hydrolysis of the peptide bond between the polysaccharide chain and the stem peptide and glycyl-glycine endopeptidases (e.g. lysostaphin, EC 3.4.24.75) cleave the attachment site of the peptidoglycan cross-bridges (Jayaswal et al., 1990; Schindler and Schuhardt, 1964). The peptidoglycan hydrolysis by bacterial enzymes is essential not only during bacterial growth but also for cell division, peptidoglycan turnover and other biological processes, hence being highly controlled to avoid autolysis of the cells. During the reconstruction of the cell wall peptidoglycan fragments are released from the murein sacculus and despite an effective recycling system a portion of these turnover products is lost in the surroundings (Boothby et
al., 1973; Goodell, 1985). Other microorganisms, animals and plants also contain peptidoglycan degrading activities. However, these enzymes are a part of the antibacterial defense machinery (see chapters 1.2.3 and 1.4).

Due to its essential function within the bacterial cell wall peptidoglycan is strongly conserved. In addition, PGN is exposed at the cell surface, hence fulfilling important requirements for an optimal PAMP.

1.4 PGN processing and perception in animals

The bacterial peptidoglycan acts immunostimulatory in metazoans and several components of the PGN detection machinery have been elucidated in the past years both in insects and mammals (Dziarski and Gupta, 2010; Girardin and Philpott, 2004; Royet and Dziarski, 2007).

In the fruit fly, *Drosophila melanogaster*, two distinct systems of peptidoglycan detection exist. The Toll pathway senses preferentially Lys-type peptidoglycan derived from Gram-positive bacteria, whereas the IMD (immune deficiency) pathway recognises mainly Gram-negative bacteria and their Dap-type PGN (Leulier et al., 2003). Interestingly, the LRR-domain containing cell surface receptor Toll is not directly activated by peptidoglycan, but by the cytokine Spätzle (Figure 1-4), a signaling intermediate, of which proteolytic cleavage is triggered by infection with Gram-positive bacteria (Weber et al., 2003). The intracellular domain of the Toll receptor, the TIR (Toll/Interleukin-1 receptor) domain is then mediating the signal transduction via the TIR-containing adapter protein dMyD88 (Belvin and Anderson, 1996; Horng and Medzhitov, 2001; Kopp and Medzhitov, 1999).

The genuine PGN receptors belong to the Peptidoglycan Recognition Protein (PGRP) family. In total, 19 PGRP proteins are present in *Drosophila*, some of them being differential splicing products of a single PGRP gene (Werner et al., 2000). The PGRP domain resembles the N-acetylmuramoyl-L-alanine amidase domain of bacterial enzymes and indeed some of the PGRPs have amidase activity and are able to digest PGN (Mellroth et al., 2003). The PGN receptor PGRP-SA binds specifically to Lys-type PGN and thereby triggers the Toll pathway culminating in the nuclear factor κB (NF-κB)-dependent induction of the antimicrobial peptide (AMP) gene *Drosomycin* (Michel et al., 2001). In addition, The Gram-negative binding protein 1 (GNBP1) is needed for the PGRP-SA-mediated activation of the Toll pathway (Figure 1-4). Experimental data suggest that both the degradation of polymeric PGN into smaller fragments and the generation of reducing MurNAc ends by GNBP1, and the physical interaction of GNBP1 with the receptor protein PGRP-SA in a PGN-dependent manner are crucial steps for the perception (Filipe et al., 2005; Wang et al., 2006). Whereas *M. luteus*
PGN is strictly recognized by PGRP-SA, some other Lys-type PGNs can be sensed by the partially redundant receptors PGRP-SA and PGRP-SD (Bischoff et al., 2004). PGRPs with amidase activity have been suggested to have either immunostimulatory or scavenger properties (Garver et al., 2006; Mellroth et al., 2003). The paradigm of receptors and scavengers belonging to the same family is logic; the scavenger PGRP cleaves in the middle of the binding site of the PGRP receptor dampening thereby the immune response (Mellroth et al., 2003).

The IMD pathway is named after the signaling intermediate protein IMD (a Death Domain protein), which upon stimulation of the PGN receptors activates several downstream pathways resulting in the activation of genes encoding for antimicrobial peptides and other defense responses. In contrast to the Toll pathway, the IMD pathway preferentially induces the expression of Dipterican, Attacin and Drosocin genes (Royet and Dziarski, 2007). The main PGN receptor stimulating the IMD pathway upon infection with Gram-negative bacteria is the membrane-bound PGRP-LC (Choe et al., 2002; Gottar et al., 2002; Ramet et al., 2002). However, also the soluble receptor protein PGRP-LE can activate the IMD pathway and for instance upon E.coli infection both PGRP-LC and PGRP-LE are required to effectively produce resistance (Takehana et al., 2004). The minimal motif of the Dap-type PGN stimulating PGRP-LC is a GlcNAc-MurNAc monomer attached to a tetrapeptide containing m-DAP, the so-called tracheal cytotoxin (TCT) (Stenbak et al., 2004). TCT is perceived by a heterodimeric complex containing the isomers PGRP-LCx (which has affinity to PGN) and PGRP-LCa (an isomer with no PGN affinity), and the PGRP-LCx homodimer complex acts as a receptor for the polymeric Dap-type PGN in vitro (Kaneko et al., 2004; Lim et al., 2006; Mellroth et al., 2005; Stenbak et al., 2004). Also the dimerisation of PGRP-LE upon ligand binding has been suggested (Lim et al., 2006). Additionally to the extracellular role described above, PGRP-LE has an additional role within the cytosol. The Drosophila PGRP-LE mutant is susceptible to infection by the intracellular bacterium Listeria monocytogenes with Dap-type PGN (Yano et al., 2008). The finding suggests that a specific sensoring system depending on PGRP-LE is present detecting invading pathogenic bacteria able to escape the cell-surface receptors. Beside the PGRP receptors, also enzymatically active PGRPs are involved in the regulation of the IMD pathway. For example, PGRP-LB, PGRP-SC1 and PGRP-SC2 have all been implicated as negative regulators of the IMD pathway by cleaving peptides from the glycan chains (Figure 1-5), thus reducing or eliminating the biological activity of PGN (Bischoff et al., 2006; Zaidman-Rémy et al., 2006). These proteins are mainly expressed in the gut and maintain the fine balance of the immune response to commensal and pathogenic bacteria (Royet and Dziarski, 2007). A recent publication added the PGRP-LF protein to the list of negative regulators of the IMD pathway.
(Basbous et al., 2011). However, unlike the other PGRPs PGRP-LF contains no PGN-docking groove and thus cannot bind PGN. The downregulation of the IMD pathway is taking place via interaction with the PGN receptor PGRP-LC (Basbous et al., 2011).

Figure 1-4: Peptidoglycan sensing in animals

Recognition of peptidoglycan in *Drosophila* (left) and mammals (right) by the innate immune system relies on specific detection of Dap-type or Lys-type peptidoglycan (PGN). In *Drosophila*, Dap-type PGN is sensed by PGRP-LC and PGRP-LE leading to the stimulation of the IMD pathway and finally production of antimicrobial peptides. PGRP-LB and PGRP-SC1/2 act via their amidase activity as modulators of PGN-induced inflammatory responses. Lys-type PGN is mainly recognised by the PGRP-SA, whereas it’s complexed to GNBP1, leading to Spätzle-dependent activation of the Toll receptor. The signal transduction is mediated via TIR domain-containing dMyD88 resulting in immune responses, including gene expression of antimicrobial peptides. Additionally, also PGRP-SD mediates the perception of certain types of Gram-positive PGN and PGRP-SB1 acts bactericidal. In mammals, the intracellular LRR-domain containing proteins Nod1 and Nod2 recognise Dap- and/or Lys-type PGN. The TLR2/CD14 receptor complex mediates perception of Lys-type PGN. The mammalian PGRPs, PGLYRPs, function either directly bactericidal or harbor amidase activity. PGRP, peptidoglycan recognition protein; GNBP1, Gram-negative binding protein 1; IMD, Immune deficiency; TLR, Toll-like receptor; Nod, nucleotide oligomerisation domain; LRR, leucine-rich repeat; TIR, Toll/Interleukin 1 receptor; CARD, caspase recruitment domain. Based on Girardin and Philpott (2004).

In mammals, both extracellular and intracellular receptors detecting peptidoglycan are present. The membrane-bound Toll-like receptor 2 (TLR2) contains an extracellular LRR-domain similar to its *Drosophila* homologue Toll. TLR2 has been shown to colocalise with
peptidoglycan and mediate together with another cell-surface receptor protein, CD14, the extracellular recognition of PGN (Dziarski et al., 1998; Gupta et al., 1996; Müller-Anstett et al., 2010; Schandner et al., 1999; Takeuchi et al., 1999; Yoshimura et al., 1999). The role of TLR2 as a specific PGN receptor has been heavily debated though (Dziarski and Gupta, 2005b; Travassos et al., 2004; Zähringer et al., 2008), and experimental data have also suggested other agonists for the TLR2-CD14 receptor complex, such as lipoproteins and chitin (Bubeck Wardenburg et al., 2006; Da Silva et al., 2008; Müller et al., 2010). The intracellular PGN sensing system containing members of the Nod-like receptor (NLR) family is widely accepted. The two main receptors, Nod1 and Nod2, contain a C-terminal LRR-domain for ligand sensing, an N-terminal caspase recruitment domain (CARD) and a central nucleotide oligomerisation domain (NOD) (Figure 1-4) (Inohara et al., 2005; Tanabe et al., 2004). Interestingly, these mammalian cytosolic receptors have striking structural similarities to the plant NBS-LRR-type resistance proteins (Ausubel, 2005; Nürnberger et al., 2004). However, the mammalian Nod proteins and plant NBS-LRR resistance proteins perceive different pathogenic signatures and activate completely different downstream signaling cascades giving little evidence for a common evolutionary origin (Ausubel, 2005). The PGN binding specificities of Nod1 and Nod2 vary greatly. Nod1 is a specific receptor for Dap-type PGN and its minimal motif is L-Ala-D-Glu-DAP (iE-DAP) (Chamaillard et al., 2003; Girardin et al., 2003a). In contrast, the Nod2 is a more universal PGN receptor recognising muramyl dipeptide (MurNAc-L-Ala-D-Glu; MDP), which is present in all peptidoglycans (Girardin et al., 2003b; Inohara et al., 2003). Upon PGN-dependent stimulation Nod1 and Nod2 activate a NF-κB-mediated pro-inflammatory response (Strober et al., 2006). The TLR- and Nod-dependent defense pathways also crosstalk with each other and can act in a synergistic manner (Petterson et al., 2011; Tada et al., 2005).

In addition to the receptor-based surveillance systems, mammals also possess PGRPs orthologous to the insect PGRPs. Whereas the Drosophila PGRPs carry out various tasks (recognition of PGN and activation of defense pathways, negative regulation of PGN-triggered inflammation or direct antimicrobial activity), the mammalian PGRPs have somewhat limited functions (Figure 1-4). The four Peptidoglycan Recognition Proteins (PGLYRP1-4) present in mammals are all secreted proteins, three of them (PGLYRP1, PGLYRP3 and PGLYRP4) are bactericidal and PGLYRP2 is an amidase (Figure 1-5) (Royer and Dziarski, 2007). The mammalian PGRP domain binds similar to the insect PGRP with high affinity to muramyl-tripeptide, but it does not bind muramyl-dipeptide or a peptide without MurNAc (Guan et al., 2004; Kumar et al., 2005; Swaminathan et al., 2006). Mammalian PGRPs bind both Dap- and Lys-type PGN (Liu et al., 2000; Lu et al., 2006) and thus do not display the same specificity and differential PGN-mediated responses as D.
melanogaster PGRPs (Royet and Dziarski, 2007). Furthermore, it has been reported that some PGRPs also bind to other polymeric structures, like lipoteichoic acid (LTA) and LPS and to some fungi (Liu et al., 2000; Lu et al., 2006; Tydell et al., 2002). In the case of human and mouse PGRPs the highest affinity is for peptidoglycan (Liu et al., 2000; Lu et al., 2006). The preferred substrates for PGLYRP2 are soluble PGN fragments, which may be generated either by bacterial PGN hydrolases or other host enzymes, like lysozyme (Gelius et al., 2003; Wang et al., 2003). The minimal PGN fragment hydrolysed by PGLYRP2 is muramyl tripeptide, which is also the minimal binding motif for PGRPs (Wang et al., 2003). It has been suggested that PGLYRP2 acts as a scavenger dampening PGN-triggered pro-inflammatory responses similar to the amidase-active fruit fly PGRPs (Hojier et al., 1997). Interaction with the bacterial cell wall peptidoglycan is essential for the bactericidal activity of PGLYRP1, PGLYRP3 and PGLYRP4, and they also require N-glycosylation and divalent cations for their antimicrobial properties (Wang et al., 2007). Although the bactericidal activity of PGLYRPs differs from the activity of other antimicrobial peptides, they are present in the same sites in the body and combat bacteria synergistically with each other (Wang et al., 2007).

Figure 1-5: Metazoan PGN hydrolytic activities

The cleavage of glycan and peptide bonds of bacterial peptidoglycan by Drosophila and mammalian PGRPs and lysozyme with N-Acetylmuramoyl-L-alanine amidase or N-Acetylglucosaminidase activities are shown. Lysozyme and other N-Acetylglucosaminidases hydrolyse the glycosidic bond between MurNAc and GlcNAc, whereas the N-Acetylmuramidases cleave between GlcNAc and MurNAc. PGRP, peptidoglycan recognition protein.

Lysozymes also belong to the antimicrobial defense armory in metazoans. They are divided into three subgroups: c-type (chicken or conventional type), g-type (goose-type) and i-type (invertebrate type) lysozymes (Callewaert and Michiels, 2010). Lysozymes are present in tissues and body secretions that are in contact with the environment or involved in bacterial
neutralization. The muramidase activity of lysozyme hydrolyses the glycosidic bond between MurNAc and GlcNAc (Figure 1-5) resulting in loss of peptidoglycan integrity and bacterial lysis (Jollès, 1996). Thereby soluble immunoactive PGN fragments are released (Zaidman-Rémy et al., 2006). Some pathogenic bacteria have found ways to acquire lysozyme resistance. For instance, peptidoglycan modification by O-acetylation and N-glycolylation or production of lysozyme inhibitors can ward off the PGN-hydrolytic activity of lysozyme (Clarke and Dupont, 1992; Monchois et al., 2001; Raymond et al., 2005).

The detection of bacterial PGN in animals is a highly complex system involving PGN sensing, binding and hydrolysing molecules. Similarities but also many differences occur between the surveillance systems of insects and higher animals.

1.5 Lysin motif (LysM) as mediator in carbohydrate signaling in plants

The lysin motif (LysM), usually about 40 amino acids in length, is a ubiquitous protein domain found in all living organisms except for Archaea (Bateman and Bycroft, 2000; Zhang et al., 2007). The three dimensional $\beta\alpha\alpha\beta$ structure of the LysM contains two $\alpha$-helices stacking onto one side of a two-stranded antiparallel $\beta$-sheet (Bateman and Bycroft, 2000; Bielnicki et al., 2006; Mulder et al., 2006). In prokaryotes, many LysM-containing proteins are bacterial lysins or chitinases, which hydrolyse glycosidic bonds present in peptidoglycan and chitin, a homopolymer of N-acetylglucosamine, respectively (Buist et al., 2008; Ponting et al., 1999). Thus, the LysM is generally thought to mediate (peptido)glycan binding. In fact, also the plant LysM-domain containing proteins characterised so far are implicated in recognition of carbohydrates, like chitin and lipochitooligosaccharides, containing N-Acetylglucosamine moieties. The plant LysM proteins are divided into several subgroups depending on their domain architecture (Zhang et al., 2007). The Nodulation factor receptor 1 and 5 (NFR1 and NFR5) from *Lotus japonicus* and *Glycine max* (Indrasumunar et al., 2010; Madsen et al., 2003; Radutoiu et al., 2003) and the LysM-type receptor-like kinase 3 (LYK3) and LYK4 from *Medicago truncatula* (Limpens et al., 2003) belong to the LysM receptor-like kinase (LYK) subgroup. These legume receptor proteins have been shown to be essential for sensing lipochitooligosaccharides (Nod factors) derived from rhizobacteria and hence for the formation of nitrogen-fixing nodules and the establishment of endosymbiosis (Indrasumunar et al., 2010; Limpens et al., 2003; Madsen et al., 2003; Radutoiu et al., 2003). The perception of the fungal PAMP chitin in rice is mediated by two LysM-containing PRRs, a LysM receptor-like protein (LYP) called CEBiP (Chitin elicitor binding protein) and a LysM receptor kinase, OsCERK1 (Kaku et al., 2006; Shimizu et al., 2010). In *Arabidopsis*, the receptor kinase CERK1, which contains three extracellular LysM motifs, has been reported to play an
important role in chitin perception and basal fungal resistance (Miya et al., 2007; Wan et al., 2008). The combination of LysM and kinase domains is a unique feature of plant LysM proteins (Bateman and Bycroft, 2000). Likewise, the presence of a specific intervening sequence between LysM domains containing a conserved CxC motif is exclusively present in the plant lineage (Arrighi et al., 2006; Madsen et al., 2003; Radutoiu et al., 2003). Although the plant LysM motifs display similarities, they are evolutionarily specialised in the perception of distinct oligosaccharide structures. Accordingly, *Lotus japonicus* NFR1 and NFR5 did not participate in the activation of chitin-induced responses (Wan et al., 2008). Not only alterations within the LysM domain, but also adaptations of the kinase domain renders specificity for the LysM receptor kinases as demonstrated by Shimizu et al. (2010).

### 1.6 Aims of the thesis

The main goal of this work was to gain insights into the plant perception mechanism for the bacterial PAMP peptidoglycan. The activation of the immune response by peptidoglycan in *Arabidopsis thaliana* was only recently revealed and experimental data suggested a receptor-dependent mode of action (Erbs et al., 2008; Gust et al., 2007). Reverse genetic approaches, bacterial infection assays and peptidoglycan elicitation assays were employed to find possible peptidoglycan receptor proteins in *Arabidopsis*.

Moreover, the possible degradation of the large biopolymeric structure of PGN by plant enzymes prior to the recognition process was analysed. Such processing of complex microbial surface molecules into more accessible fragments has been reported to take place *in planta* (Fliegmann et al., 2004; Mithöfer et al., 2000). The role of a putative *Arabidopsis* PGN hydrolase in bacterial cell wall degradation and bacterial resistance was studied both *in vitro* and *in vivo*. 
2 Materials and methods

2.1 Materials

2.1.1 Chemicals

All used standard chemicals were of standard purity and purchased from Sigma-Aldrich (Taufkirchen), Carl Roth (Karlsruhe), Merck (Darmstadt), Qiagen (Hilden), Invitrogen (Karlsruhe), Duchefa (Haarlem, Niederlande), Molecular Probes (Leiden, Niederlande), Fluka (Buchs, Schweiz) und BD (Sparks, USA), unless noted otherwise in the text. Restriction enzymes, ligase and DNA modification enzymes were purchased from Fermentas (St. Leon-Rot) and New England Biolabs (Beverly, USA). Oligonucleotides were received from Eurofins MWG Operon (Ebersberg) and antibodies from the companies Sigma-Aldrich (Taufkirchen), New England Biolabs (Beverly, USA) and Acris Antibodies GmbH (Herford). The antibody against tobacco class III chitinases from rabbit and the antibody against YFP from rabbit were kind gifts from Frédéric Brunner and Sara Mazzotta, respectively.

*Xanthomonas campestris pv. campestris* PGN and muro-peptides derived thereof were kindly provided by Mari-Anne Newman. The synthetically generated Flg22 peptide was a kind gift from Georg Felix. *Pto* DC3000 PGN was prepared as described in 2.4.1. *Staphylococcus aureus* PGN was obtained from Fritz Götz (Microbiology Department, University of Tübingen) or prepared as described in 2.4.2. *Bacillus subtilis, Escherichia coli* and *Micrococcus luteus* PGN and chitin were commercially available (Invivogen, Sigma). All described PGNs and chitin were dissolved in water at a concentration of 10mg/ml and stored at -20°C.

2.1.2 Media

Table 2-1 summarizes the media used in this work. All media were prepared using deionized water and sterilized by autoclaving for 20 minutes at 121°C. For solid media 15g/l Bacto-agar (BD) or 8g/l Select-agar for MS plates (Sigma-Aldrich) was added to the medium. If necessary, antibiotics were added to the sterilized medium in appropriate final concentrations as listed in Table 2-2.
### Materials and Methods

<table>
<thead>
<tr>
<th>Medium</th>
<th>Ingredients per 1 liter</th>
<th>Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>LB</td>
<td>10 g Bacto-Tryptone, 5 g NaCl, 5 g Yeast extract (YE)</td>
<td><em>Escherichia coli</em></td>
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<tr>
<td>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>
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<tr>
<td>MD</td>
<td>1.34 % (w/v) YNB, 4x10^-5 % (w/v) Biotin, 2 % (w/v) Glucose</td>
<td><em>Pichia pastoris</em></td>
</tr>
<tr>
<td>YPD</td>
<td>20 g Peptone, 20 g Glucose, 10 g YE</td>
<td><em>Pichia pastoris</em></td>
</tr>
<tr>
<td>BMGY and BMMY</td>
<td>1 % (w/v) YE, 2 % (w/v) Peptone, 1.34 % (w/v) YNB, 100 mM potassium phosphate, pH 6.0, 4x10^-5 % (w/v) Biotin, 1 % (v/v) glycerol or 2 % (v/v) MeOH</td>
<td><em>Pichia pastoris</em></td>
</tr>
<tr>
<td>CSM-LT</td>
<td>6.7 g YNB, 20 g glucose, 1546 mg -Leu, -Trp Kaiser-dropout (20 g Oxoid agar for plates)</td>
<td><em>Saccharomyces cerevisiae</em></td>
</tr>
<tr>
<td>CSM-LTA</td>
<td>6.7 g YNB, 20 g glucose, 1546 mg -Ade, -His, -Leu, -Trp Kaiser-dropout, 76 mg L-Histidin (20 g Oxoid agar for plates)</td>
<td><em>Saccharomyces cerevisiae</em></td>
</tr>
<tr>
<td>½ PDB</td>
<td>12 g PDB (Potato Dextrose Broth, Duchefa), pH 5.8 (NaOH)</td>
<td><em>Botrytis cinerea</em></td>
</tr>
<tr>
<td>½ MS</td>
<td>2.2 g MS (Duchefa), pH 5.7 (KOH)</td>
<td><em>Arabidopsis thaliana</em></td>
</tr>
</tbody>
</table>

**Table 2-1: Used media**

<table>
<thead>
<tr>
<th>Antibiotics</th>
<th>c (µg/µl)</th>
<th>Solvent</th>
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<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>Tetracyclin</td>
<td>50</td>
<td>Ethanol</td>
</tr>
</tbody>
</table>

**Table 2-2: Used antibiotics**
2.1.3 Vectors

All used vectors are listed in Table 2-3.

<table>
<thead>
<tr>
<th>Vector</th>
<th>Characteristics</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>pDONR201</td>
<td>Ori Puc, rrnB, T2, rrnB,T1, attP1, attP2, ccdB, Cm', Kan'</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pDONR207</td>
<td>Ori Puc, rrnB, T2, rrnB,T1, attP1, attP2, ccdB, Cm', Gent'</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pK7FWG2.0</td>
<td>$P_{35S}$, $T_{35S}$, eGFP, attR1, attR2, ccdB, Cm', Kan'</td>
<td>VIB</td>
</tr>
<tr>
<td>pK7WGF2.0</td>
<td>$P_{35S}$, $T_{35S}$, eGFP, attR1, attR2, ccdB, Cm', Kan'</td>
<td>VIB</td>
</tr>
<tr>
<td>pDEST17</td>
<td>PT7, RBS, His$_6$-tag, attR1, attR2, ccdB, Cm', PT7, bla, Promotor, Amp', pBR322 origin, ROP, orf</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pBGWFS7</td>
<td>attR1, attR2, ccdB, Ba', Sm/Sp', GUS, eGFP</td>
<td>Karimi et al. 2005</td>
</tr>
<tr>
<td>pBGW</td>
<td>attR1, attR2, ccdB, Cm',Sm/Sp', Ba'</td>
<td>Karimi et al. 2005</td>
</tr>
<tr>
<td>miR319a pBSK</td>
<td>B reverse, T3 promoter, miR319a, T7 promoter, A forward, Amp'</td>
<td>Weigelworld.org</td>
</tr>
<tr>
<td>(pRS300)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pBluescript</td>
<td>pUCori, P Lac, MCS, lac Z', f1+ori, Amp'</td>
<td>Stratagene</td>
</tr>
<tr>
<td>pGREEN0229</td>
<td>pSa-ORI, ColE1 ori, MCS, lac Z', Kan', Ba'</td>
<td>Hellens et al. 2000</td>
</tr>
<tr>
<td>pPIC9K</td>
<td>5'AOX1, Ampr, pBR322, 3'AOX1, Kan', HiS4 ORF, TT, secretion signal</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pPICZoC</td>
<td>5'AOX1, Ori Puc, CYC1 TT, Zeocin', $P_{EM7}$, $P_{TEF1}$, AOX1 TT, $\alpha$-Factor, cmyc-epitope, His$_6$-tag</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pGADT7-GW</td>
<td>MCS, Amp', GAL4-AD, HA-tag, T7 promoter, gateway cassette (attR1, attR2, ccdB, Cm') introduced into MCS-site</td>
<td>Clontech Laboratories (modified by Sandra Postel)</td>
</tr>
<tr>
<td>pGBK7-GW</td>
<td>MCS, Amp', GAL4-DNABD, Myc-tag, T7 promoter, gateway cassette (attR1, attR2, ccdB, Cm') introduced into MCS-site</td>
<td>Clontech Laboratories (modified by Sandra Postel)</td>
</tr>
</tbody>
</table>

Table 2-3: Used vectors

2.1.4 Primers

The primers used in this work for cloning, genotyping, transcript analysis and sequencing are listed in the Appendix Table 8-1.
2.2 Organisms

2.2.1 Bacteria and fungi

The bacterial strains used in the frame of this work are listed in Table 2-4.

<table>
<thead>
<tr>
<th>Species</th>
<th>Strain</th>
<th>Genotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>Escherichia coli</td>
<td>DH5α</td>
<td>supE44 ΔlacU169 (Φ80 lacZM15) hsdR17 recA1 endA1 gyrA96 thi-1 relA1</td>
</tr>
<tr>
<td></td>
<td>TOP10</td>
<td>mcrA, delta (mrr-hsdRMS-mcrBC), phi 80delta lac delta M15, delta lacX74, deoR, recA1, araD139 delta (ara, leu), 7697, galU, galK, lambda, rpsL, endA1, mupG</td>
</tr>
<tr>
<td></td>
<td>DB3.1</td>
<td>RR1 gyrA endA recA Spec′</td>
</tr>
<tr>
<td></td>
<td>BL21AI</td>
<td>F-ompT hsdSb(rb-mb-) gal dcm araB::T7RNAP-tetA</td>
</tr>
<tr>
<td>Pseudomonas syringae</td>
<td>Pto DC3000</td>
<td>Rif′</td>
</tr>
<tr>
<td></td>
<td>Pto DC3000 ΔavrPto/PtoB</td>
<td>Rif′, Kan′, ΔavrPto, ΔavrPtoB</td>
</tr>
<tr>
<td></td>
<td>Pto DC3000 hrcC′</td>
<td>Rif′ Kan′ (nptII)</td>
</tr>
<tr>
<td>Agrobacterium tumefaciens</td>
<td>GV3103::pMP90</td>
<td>T-DNA vir′ rif′, pMP90 gen′</td>
</tr>
</tbody>
</table>

Table 2-4: Used bacterial strains

Additionally, the necrotrophic fungi Alternaria brassicicola (MUCL 20297) and Botrytis cinerea (B05-10) were used for fungal infections, the yeast Pichia pastoris (GS115) for protein expression and Saccharomyces cerevisiae strain AH109 for yeast two-hybrid experiments.

2.2.2 Arabidopsis thaliana lines

All experiments were conducted using the Arabidopsis thaliana ecotypes Columbia-0 (Col-0), Columbia-2 (Col-2) or Landsberg erecta (Ler) and transgenic lines generated in these ecotypes. The knock-out (KO) lines mainly used in this work are listed in Table 2-5 and were purchased from the Nottingham Arabidopsis Stock Centre (NASC) or received from other
research groups. The transgenic \textit{p35S::secGFP} (\textit{secGFP}) line has been described previously (Teh and Moore, 2007).

<table>
<thead>
<tr>
<th>Stock number</th>
<th>T-DNA/KO line</th>
<th>\textit{AGI}</th>
<th>Position of the T-DNA</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>WiscDsLox387C11</td>
<td>\textit{chia-1}</td>
<td>At5g24090</td>
<td>promoter</td>
<td>this work</td>
</tr>
<tr>
<td>SALK_095362</td>
<td>\textit{chia-2}</td>
<td>At5g24090</td>
<td>3. exon</td>
<td>this work</td>
</tr>
<tr>
<td>CSHL_ET14179</td>
<td>\textit{chia-3}</td>
<td>At5g24090</td>
<td>1. intron</td>
<td>this work</td>
</tr>
<tr>
<td>GABI_096F09</td>
<td>\textit{cerk1-2}</td>
<td>At3g21630</td>
<td>10. exon</td>
<td>Miya et al. 2007</td>
</tr>
<tr>
<td>SALK_012441</td>
<td>\textit{lyk2-1}</td>
<td>At3g01840</td>
<td>1. intron</td>
<td>Volker Lipka</td>
</tr>
<tr>
<td>SALK_140374</td>
<td>\textit{lyk3-1}</td>
<td>At1g51940</td>
<td>10. exon</td>
<td>Volker Lipka</td>
</tr>
<tr>
<td>SALK_030271</td>
<td>\textit{lyk3-2}</td>
<td>At1g51940</td>
<td>10. exon</td>
<td>Volker Lipka</td>
</tr>
<tr>
<td>GABI_857A10</td>
<td>\textit{lyk4-1}</td>
<td>At2g23770</td>
<td>exon at 1615 bp</td>
<td></td>
</tr>
<tr>
<td>CSHL_GT7089</td>
<td>\textit{lyk5-1}</td>
<td>At2g33580</td>
<td>exon at 1280 bp</td>
<td>Volker Lipka</td>
</tr>
<tr>
<td>SALK_131911</td>
<td>\textit{lyk5-2}</td>
<td>At2g33580</td>
<td>exon at 870 bp</td>
<td></td>
</tr>
<tr>
<td>SALK_140374/CSHL_GT7089</td>
<td>\textit{lyk3-1/lyk5-1} (LGK2-6-1-2)</td>
<td>At1g51940/At2g33580</td>
<td></td>
<td>Volker Lipka</td>
</tr>
<tr>
<td>GABI_096F09/SALK_140374</td>
<td>\textit{cerk1-2/lyk3-1}</td>
<td>At3g21630/At1g51940</td>
<td></td>
<td>this work</td>
</tr>
<tr>
<td>GABI_096F09/SALK_140374/CSHL_GT7089</td>
<td>\textit{cerk1-2/lyk3-1/lyk5-1}</td>
<td>At3g21630/At1g51940/At2g33580</td>
<td></td>
<td>this work</td>
</tr>
<tr>
<td>SALK_111212</td>
<td>\textit{lym3-1}</td>
<td>At1g77630</td>
<td>1. intron</td>
<td>Roland Willmann 2011</td>
</tr>
</tbody>
</table>

Table 2-5: Knock-out and complementation lines used in this work

### 2.3 Cultivation conditions of the organisms

#### 2.3.1 Growth of \textit{Escherichia coli}

\textit{E.coli} strains were cultivated overnight at 37°C either on LB-plates or in liquid LB medium at 230 rpm. Antibiotics were added into the media according to the resistance cassettes the strains were harboring.
2.3.2 Growth of *Pseudomonas syringae*

*P. syringae* strains were grown for 24-48 hours at 28°C either on King's B-plates or in liquid King's B medium at 180 rpm. For the determination of bacterial growth in infection assays the *Pseudomonas* strains were re-isolated from plant material (see 2.6.1) and plated on LB-plates containing cycloheximide in addition to other antibiotics.

2.3.3 Growth of *Agrobacterium tumefaciens*

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

2.3.4 Growth of *Pichia pastoris*

*P. pastoris* strains were grown in liquid BMGY medium overnight at 30°C and 230 rpm to an OD600 of 2-6. The cells were harvested and resuspended in BMMY medium (OD600~1) to induce expression. To maintain the expression the yeast cells were fed with 0.5% (v/v) MeOH every 24 hours.

2.3.5 Growth of *Alternaria brassicicola* and *Botrytis cinerea*

The cultivation of *A. Brassicicola* and *B. cinerea* and the preparation of the spores for the infection assays were performed as published previously (Kemmerling et al., 2007).

2.3.6 Growth of *Arabidopsis thaliana* and *Nicotiana benthamiana*

*A. thaliana* 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 and in the dark the plants were grown in environmental chambers either in long-day (16 h light, 8 h darkness) or short-day (8 h light, 16 h darkness) 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).
2.4 Methods

2.4.1 Isolation of peptidoglycan from Gram-negative bacteria

Peptidoglycan was isolated from the Gram-negative phytopathogenic bacteria *Pseudomonas syringae* (Pto DC3000) using a modified protocol of Glauner (1988). The fermentation of the bacteria was performed in the fermentation unit of the Microbiology department by Andreas Kulik (RG Fiedler). The preparation of the 10L fermentation culture was done in two steps. First, a small preculture (10 ml) was inoculated with bacteria freshly grown on King’s B agar plate and grown at 28°C overnight. Secondly, a large preculture (1L) was inoculated with the small preculture and cultivated at 28°C for 8 hours up to a density of 3 - 3.5 (OD600nm). The 1 liter culture was used for the inoculation of the 10 liter fermenter of the type Biostat E. The fermentation was performed at 28°C for 16 hours using free pH and pO2 conditions. The bacteria were harvested by flow-through centrifugation (45 minutes at 5500 x g and 4°C) and immediately frozen in liquid nitrogen. After lyophilization the dry bacterial pellet (25 - 30 g) was homogenized by grinding and resuspended in ice-cold water in small portions (~5 g in 40 ml water). Immediately after resuspension the bacteria were added dropwise into boiling 4 % (w/s) SDS solution (~1g dry bacteria/10 ml SDS). The bacterial suspension was boiled for 1 hour under continuous stirring and then cooled down overnight at RT. The heat-stable peptidoglycan was harvested by ultracentrifugation for 1 hour at 43000 x g and at RT (Sorvall WX Ultra 80, Thermo Scientific). The PGN pellet was washed until it was free of SDS. For detection of residual SDS 335 µl washing supernatant was mixed with 170 µl 0.7 M sodium phosphate buffer, pH 7.2, 7µl 0.5 % (w/v) methylene blue and 1 ml chloroform. An SDS-free solution appeared light red, whereas a SDS-containing sample showed a light blue colour. The SDS-free peptidoglycan sample was then treated with several enzymes for degradation and removal of PGN-associated molecules. Degradation of high molecular weight glycogen was facilitated using 100µg/ml α-amylase (Fluka) in 10 mM Tris-HCl buffer pH 7 (2 hours at 37°C). In a subsequent second step the peptidoglycan was treated with 10 µg/ml D Nasel (Sigma) and 50 µg/ml RNase A (Qiagen) for additional two hours at 37°C, following an overnight digestion with 100 µg/ml Trypsin (Sigma) at 37°C (degradation and removal of RNA and DNA and covalently bound lipoproteins). Afterwards the PGN pellet was resuspended in Proteinase K buffer (50 mM Tris pH 7.5, 5 mM CaCl2, 0.5 % (w/v) SDS) and digested for one hour at 65°C with 50 µg/ml Proteinase K (Carl Roth) for removal of residual proteinous impurities. After the digestion Proteinase K was inactivated for 15 minutes at 75°C. After washing the PGN pellet with water and harvesting by ultracentrifugation it was incubated at RT for one hour with shaking in 8 M LiCl solution. After washing the pellet for two times, it was further incubated in 100 mM EDTA, pH 8 for one hour. Then the PGN pellet was washed again and finally treated with 100 % (v/v) acetone at RT for two hours (removal
of lipoteichoid acids and traces of LPS). After final washing steps (3 times with water) the
PGN pellet was lyophilized and stored either as powder or resuspended in water at -20°C. In
the case of the presence of residual impurities in the PGN preparation after the isolation
procedure, the Proteinase K digestion was repeated.

2.4.2 Isolation of peptidoglycan from Gram-positive bacteria
The isolation procedure for peptidoglycan from Gram-positive bacteria, such as
Staphylococcus aureus, was performed as published previously (Bera et al., 2005). In
comparison to the isolation protocol of PGN from Gram-negative bacteria, some steps of the
basic scheme are either added or removed. The starting material was less (0.5 - 2 liter) and
the bacterial cells were disrupted after boiling in SDS by vigorous homogenization. The
digestion step with Proteinase K was omitted, but instead the PGN pellet was treated with
hydrofluoric acid (HFA) before the LiCl incubation step to remove residual covalently bound
polysaccharides, e.g. teichoic acids.

2.4.3 General molecular biology methods
Standard protocols were used for PCR, agarose gel electrophoresis, restriction digestion,
ligation, transformation of bacteria and yeast and plasmid isolation (Ausubel, 1993;
Sambrook and Russell, 2001). The transformation of TOP10 cells was performed according
to the manufacturer’s protocols (Invitrogen). The enzymes were used according the
manufacturer’s protocols (Fermentas and NEB). Deglycosylation of proteins was carried out
using the deglycosylation Kit and according to manufacturer’s recommendations (NEB). For
the generation of PCR fragments either the Taq DNA-Polymerase or the Pfu DNA-
polymerase (cloning purposes; Fermentas) were used. GeneRuler™ DNA Ladder Mix
(Fermentas) was used as size marker for the agarose gel electrophoresis. DNA fragments
were extracted out of agarose gels or purified out of PCR reactions by using the Qiagen Gel
Extraction Kit and Qiagen PCR Purification Kit (Qiagen).

2.4.4 Cloning
The constructs were generated either by traditional cloning techniques (via introduced
restriction sites) or by the Gateway™ Technology (Invitrogen). For the traditional cloning the
digested vector was treated with the antarctic phosphatase (NEB) prior ligation to inhibit self-
ligation. Gateway™-cloning was performed according to the manufacturer’s
recommendations (Invitrogen). To obtain Gateway-compatible inserts gene-specific adaptor
primers were used in the first PCR. The essential recombination sites were then completed in a second PCR using the Gateway-primers attB1 and attB2 (see Appendix Table 8-1). The resulting inserts were then subcloned into pDONR201 or pDONR207 (Invitrogen) by using the BP clonase reaction and afterwards inserted into the expression vectors by using the LR clonase reaction following the manufacturer’s specifications (Invitrogen).

2.4.5 DNA isolation
Genomic DNA from plant tissue for genotyping purposes was isolated as outlined in Edwards et al. (1991). For sequencing purposes plasmid-DNA were isolated and column-purified using the QIAprep Spin MiniPrep Kit (Qiagen). Sequencing of the generated constructs was performed by the companies Eurofins MWG Operon (Ebersberg) and GATC Biotech AG (Konstanz). The sequence analysis was performed using the Lasergene DNA*STAR software.

2.4.6 RNA isolation
Total RNA from leaves or seedlings was isolated using the Trizol method according to the standard protocol (Chomczynski and Sacchi, 1987). For seedling samples the standard volumes were reduced up to one-third. In the end of the isolation procedure the RNA pellet was eluted in ddH₂O (leaf RNA in 20-40 µL and seedling RNA in 10 µL) and stored at -20°C.

2.4.7 Semi-quantitative RT-PCR
1 µg of total leaf RNA was used for the first strand cDNA synthesis using RevertAidTM M-MuLV Reverse Transcriptase according to manufacturer’s recommendations (Fermentas). The analysis of residual transcript in KO lines was performed using semi-quantitative RT-PCR. 1 µL cDNA was used for a standard PCR reaction with primers specific for the analyzed transcript. In a control PCR primers specific for the house-keeping gene elongation factor 1α (EF1α) were used (Table 8-1).

2.4.8 Quantitative Real-time PCR
2.5 µL seedling RNA (amounts not adjusted) 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 Multicolour Real Time PCR detection system
from Bio-Rad. RT-qPCR amplifications were monitored using the ABsolute 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 4-6 seedlings.

2.4.9 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). After the isolation procedure
protoplasts were resuspended in W5 solution and incubated overnight at RT and in the dark
(2x10^5 protoplasts in 1ml W5 solution).

2.4.10 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 preculture
of selected agrobacteria and cultivated for further 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 (OD_{600nm}). 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.

2.4.11 Transient transformation of Nicotiana benthamiana

Agrobacterium tumefaciens-mediated transient transformation was used for the transient
expression of proteins in tobacco. The bacterial strain carrying the appropriate expression
vector was cultured as described in 2.3.3. After harvesting the cells at 4°C for 10 minutes at
2000 x g they were washed for two times with 10mM MgCl2. The density of the culture was
diluted to 5 x 10^4 cfu/ml and 150 μM acetosyringone is added. The bacterial suspension was
then incubated 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 the leaves of 3 week-old tobacco leaves. The leaf tissue was
analyzed 2-4 days post infection for the presence of the protein.
2.4.12 Crossing *Arabidopsis thaliana* plants

The sepals, petals and anthers of the young flowers of the mother plant were carefully removed. Then a mature flower from the father plant was gently opened by pinching with the forceps and rubbed onto the stigma of the emasculated mother plant until some pollen were visible on the surface of the stigma. The procedure was repeated with 4 to 6 flowers per inflorescence and after labeling of the crossed flowers they were kept at 100 % humidity under a glas bell for two days to increase the chance of fertilisation. After a day or two a successful fertilisation was observable by obvious elongation of the stigma to generate a silique. The siliques were harvested before the seeds fell out and let ripen in a reaction tube. Fully ripened seeds were sown on soil and the offspring seedlings were analysed by genotyping as described in 2.4.19.

2.4.13 Generation of knock-down lines

Artificial microRNA-mediated gene silencing was used to specifically knock-down *CHIA* in Col-0 background. The Web microRNA Designer (WMD; http://wmd.weigelworld.org) was used to select the primers (see Appendix Table 8-1) for the generation of an artificial 21mer microRNA (Schwab et al., 2005). The insert was generated in four PCR-steps (see Figure 2-1). In PCR 1 the template pRS300 and the primers A and At5g24090miR*a were used. In PCR 2 the primers At5g24090miR*-s and At5g24090miR-a were used to amplify the second product from the same template. The third product was generated using the same template and the primer pair At5g24090miR-s and B. In the fourth PCR the final insert was generated using the overlapping products from the PCRs 1-3 as template. The *CHIA*-specific amiRNA was introduced into the EcoRV site of pBSK(+) (Stratagene) via T/A cloning and subsequently cloned into the EcoRI/XbaI site of pGREEN0229 (Hellens et al., 2000). Additionally, the p35S from pK7FWG2.0 (VIB) was introduced into the HindIII/EcoRI site and the 35S-terminator from pAeq-Hyg (a kind gift from Magdalena Krzymowska) into XbaI/Sacl site of the binary vector pGREEN0229. The transformation of the resulting vector into agrobacteria was mediated using the accessory plasmid pSOUP (Hellens et al., 2000). The stable transformation of the construct into the *Arabidopsis* genome was performed using the floral-dip method (see 2.4.10). Analysis of the *CHIA* transcript level in the *CHIA* knock-down line (*chia-kd*) was performed by quantitative RT-PCR using primers listed in Table 8-1.
2.4.14 Generation of overexpression lines

For the p35S::CHIA-GFP fusion construct, a 906bp fragment of the CHIA coding sequence was cloned using the primers At5g24090gatF and At5g24090gatR (Table 8-1). For the p35S::CHIAΔSP-GFP fusion construct, which was lacking the predicted N-terminal signal peptide, a 849bp fragment of the CHIA coding sequence lacking amino acids 2-22 was cloned using the primers FP_5g24090d(2-22)gat and RP_5g24090-STOPgat (Table 8-1). In a second PCR the recombination sites of the inserts were completed using the Gateway adaptor primers (Invitrogen, see Table 8-1). The resulting inserts were first introduced into an entry vector pDONR201 and then finally into the binary expression vector pK7WGF2.0 (Karimi et al., 2005) as described in 2.4.4. The p35S::CHIA-GFP construct was stably transformed in WT A.thaliana plants (see 2.4.10) and offspring was screened for phosphinothricin (Basta) resistance.

2.4.15 Generation of pCHIA::GUS lines

For the pCHIA::GUS reporter construct, a 1948 bp fragment of the CHIA promoter sequence was cloned using the primers At5g24090_gatF and At5g24090_gatR (Table 8-1). In a second PCR the recombination sites of the inserts were completed using the Gateway adaptor primers (Invitrogen). The resulting insert was first introduced into an entry vector pDONR207 and then finally inserted into the binary expression vector pBGWFS7 (Karimi et al., 2005) by using the Gateway™ Technology (Invitrogen).
2.4.16 Generation of constructs for expression in *E. coli*

A 909 bp fragment of the coding sequence of CHIA gene was cloned using the primers At5g24090gatF and At5g24090gatR-STOP (Table 8-1). After completion of the recombination sites the resulting fragment was first introduced into an entry vector and afterwards into the *E. coli* expression vector pDEST17 (Invitrogen) using the Gateway™ Technology.

2.4.17 Generation of constructs for expression in *P. pastoris*

For the expression of the CHIA protein in the *Pichia* system both untagged and His₆-tagged versions were generated. The primers 5g24090EcoRIF and 5g24090NotI_mstopR were used to create a construct with the CHIA insert without the stop codon in pPICZαC (untagged), whereas the primers 5g24090EcoRIF and 5g24090NotI_ostopR enabled the generation of a CHIA construct with a C-terminal His₆-tag in pPICZαC. To generate an untagged version of CHIA in the pPIC9K vector the primers FP_EcoRI_pPIC9K and RP_20bp_tag_stop_NotI were taken. Each pair of used cloning primers (listed in Table 8-1) introduced the restriction sites EcoRI and NotI into the inserts, which then allowed the site-directed insertion of the inserts into the expression vectors pPICZαC and pPIC9K (both from Invitrogen).

2.4.18 Generation of constructs for the yeast two-hybrid system

The entry clone for CHIA∆SP (lacking the sequence for signal peptide; see 2.4.14) was used for the generation of the Y2H-constructs. The insert was introduced into bait and prey vectors pBGKT7-GW and pGADT7-GW, respectively. The addition of the Gateway recombination cassette into the original vectors pBGKT7 and pGADT7 (Clontech Laboratories) enabled the usage of the Gateway™ Technology (Invitrogen) (the modified vectors were a kind gift from Sandra Postel).

2.4.19 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 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 (see Figure 2-2). In the WT-PCR, primers were used, which bind a region flanking the T-DNA insertion (product amplified only in the WT plants, the large size of the T-DNA insertion inhibits the amplification in mutants).
In the second PCR a T-DNA specific left border a 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.

**Figure 2-2: Genotyping analysis of T-DNA lines by gene- and insertion-specific primer**

In the WT-PCR gene-specific primers (5’ and 3’ primer) amplify a product in WT plants, whereas in homozygous T-DNA insertion lines the large size of the T-DNA blocks the amplification. In the Lba-PCR reaction the usage of a gene-specific and a T-DNA-specific primer allows product amplification only in the T-DNA insertion lines. In heterozygous plants both PCR reactions produce an amplicon.

**2.5 Biochemical methods**

**2.5.1 Protein expression in E.coli**

BL21AI cells were used for expression of recombinant proteins. An overnight culture was used to inoculate a day culture with the density of 0.05 (OD$_{600nm}$), which was grown at 37°C up to an OD$_{600nm}$ of 0.4-0.6. Then the culture was split in two; in the one culture flask the expression of the protein was induced by the addition of 0.01 % (w/v) L-arabinose and the other culture was left uninduced. Both cultures were further incubated at 18°C and harvested at different time points. The bacterial cells were pelleted at 2000 x g for 5 minutes and frozen in liquid N$_2$. The samples were afterwards treated with lysis buffer (500 mM potassium phosphate buffer, pH 7.8, 400 mM NaCl, 100 mM KCl, 10 % (v/v) glycerol, 0.5 % (v/v) Triton X-100, 10 mM Imidazole and freshly added 100 µg/ml lysozyme) and incubated for 30 minutes at 4°C on a stirrer. After 3 freeze-thaw cycles the samples were sonicated for 2 x 10 seconds (70% intensity, Bandelin UW 2070 sonicator) on ice. In a centrifugation step the unsoluble proteins and cell fragments were pelleted and the soluble proteins were further characterized by SDS-PAGE (2.5.7) following Coomassie blue stain (2.5.9) or immunoblot (2.5.8).
2.5.2 Ni\textsuperscript{2+}-NTA affinity purification

Ni\textsuperscript{2+}-NTA agarose from Qiagen was used for the affinity purification of His\textsubscript{6}-tagged \textit{E.coli} proteins. Therefore, 1.5 ml column material was first washed 5 times with one column volume of water and then 5 times with lysis buffer (50 mM KH\textsubscript{2}PO\textsubscript{4}, 300 mM KCl, 10 mM imidazole, pH 8), elution buffer (50 mM KH\textsubscript{2}PO\textsubscript{4}, 300 mM KCl, 250 mM imidazole, pH 8) and washing buffer (50 mM KH\textsubscript{2}PO\textsubscript{4}, 300 mM KCl, 20 mM imidazole, pH 8) each. Then the crude supernatant (see 2.5.1) derived from a 500 ml bacterial culture was loaded on the column, the flow-through was collected and the column was washed two times with 6 ml washing buffer. Afterwards the His\textsubscript{6}-tagged protein was eluted from the column with 750 µl elution buffer in 6 elution steps. The protein content in the elution fractions was then analyzed by SDS-PAGE followed by Coomassie blue stain or immunoblotting.

2.5.3 Protein expression in \textit{P.pastoris}

For protein expression in \textit{Pichia pastoris} a clone carrying the desired construct was cultivated in BMGY medium while shaking at 30°C until the OD\textsubscript{600nm} reached 2-6. Then the cells were pelleted for 5 minutes at 1500 x g, resuspended in BMMY to a density of 1 (OD\textsubscript{600nm}) and further grown in the induction medium. Every 24 hours 0.5 % (v/v) MeOH was added to maintain induction conditions. After different time points 1 ml of culture was harvested, the cell pellet was concentrated using the sodium deoxycholate-trichloroacetic acid (DOC-TCA) precipitation method (Bensadoun and Weinstein, 1976) and then analysed for the presence of expressed protein.

2.5.4 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, pH5.2/15mM 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 N\textsubscript{2} and after addition of the extraction buffer the sample was incubated for 30 minutes at 4°C. Afterwards the soluble proteins were separated from the insoluble ones in a centrifugation step (15 minutes 20800 x g at 4°C) and used for further analysis.

For the extraction of protein from the protoplast samples the protoplast pellet was first separated from the medium by centrifugation (20sec 800rpm 4°C). The secreted protein in the medium was concentrated using Vivaspin 2 columns with a 10kD cut-off (GE
Healthcare). Protein from the harvested protoplast pellet was extracted using 20mM sodium acetate, pH5.2/15mM β-mercaptoethanol supplemented with 1x proteinase inhibitor cocktail (Roche).

2.5.5 Immunoprecipitation
Leaf protein was extracted from the CHIA overexpression plants (see 2.5.4) and approximately 200 µg total protein was used for the immunoprecipitation of CHIA-GFP and therefore 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), once with 800 µl buffer A (50 mM Tris-HCl, pH 7.5, 5 mM EDTA, pH 8, 5 mM EGTA, pH 8, 2 mM DTT, 10 mM AEBSF, 2 µg/ml Aprotinin, 2µg/ml Antipain). 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 further 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.

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

2.5.7 SDS-PAGE
SDS polyacrylamide gel electrophoresis was performed as described in Sambrook et. al (2001) using the gel chamber system of BioRad. 12 % SDS-PA gels were used as separating gels (with 5 % stacking gels) for the discontinuous SDS-PAGE by the method of Laemmli (1970) if not mentioned otherwise. The Prestained Protein Ladder Mix (Fermentas) was used as a protein marker.

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
Materials and Methods

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 peroxidase-coupled secondary antibody was detected using the Enhanced Chemiluminescence Kit (GE Healthcare) according to the manufacturers’ 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 the staining the membrane was washed with water.

2.5.9 Coomassie blue stain
For non-specific staining of proteins after SDS-PAGE a Coomassie blue R-250 stain (0.125 % (w/v) Coomassie blue R-250, 50 % (v/v) MeOH, 10 % (v/v) acetic acid) was used. After incubation for 30 minutes at RT the superfluous stain was removed by 10 % (v/v) acetic acid.

2.5.10 Silver stain
Silver staining was used to control the purity of peptidoglycan preparations (higher sensitivity in comparison to a Coomassie blue stain). 100 µg PGN was loaded on a 15 % SDS-PA-gel and the proteinous impurities were separated by SDS-PAGE. Then the gel was incubated for at least one hour in a fixing solution (50 % (v/v) MeOH, 12 % (v/v) acetic acid and 0.0185 % (v/v) formaldehyde), washed with 50 % (v/v) EtOH for 3 x 20 minutes and treated with fresh 0.02 % (w/v) Na2S2O3 solution for one minute. Subsequently, the gel was washed with water (3 x 20 seconds), incubated for one hour with an impregnation solution (0.2 % /w/v) AgNO3 and 0.028 % (v/v) formaldehyde) and then repeatedly washed with water. In a final step the gel was treated for 10-15 minutes with a developer solution containing 6 % (w/v) Na2CO3, 0.0185 % (v/v) formaldehyde and 0.4 % (w/v) Na2S2O3. The staining of the proteins was stopped with washing the gel with water for 2 x 2 min and then treating it with 50 % (v/v) MeOH and 12 % (v/v) acetic acid for 10 minutes.
2.5.11 Turbidity assay (PGN-hydrolysis assay)
The turbidity assay was performed as described in Park et al. (2002). Shortly, 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 (2.5.4) and 15 µg total protein of the protoplast samples (2.4.9) were added to the
reaction solutions.

2.5.12 4-MUCT assay (Chitin-hydrolysis assay)
The 4-MUCT assay was performed as described in Brunner et al. (Brunner et al., 1998).
Shortly, 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). Same protein amounts were used as for
the turbidity assay (see 2.5.11).

2.5.13 Colloidal chitin hydrolysis assay
The chitinase activity (chitodextrinase, EC 3.2.1.14) was measured using a method
published by Reissig et al. (1955). 500 µl suspension of colloidal chitin (10 mg/ml chitin in
100 mM Sodium acetate, pH 5.2) was incubated with approximately 60 µg total leaf protein
(see 2.5.4) and incubated for 2 hours at 37°C. After centrifugation (20 min 10600 x g 4°C) the
supernatant containing soluble hydrolyzed chitin fragments was collected and further
incubated with a cytohelicase for 20 minutes at 37°C to further digest the chitin fragments
into monomeric N-acetylglucosamines. Then 100 µl 0.6 M potassium tetraborate was added
to the sample and incubated for 3 minutes at 100°C. After cooling down to RT 1 ml DMAB
solution was added (1% (w/v) p-dimethylaminobenzaldehyde, 1% (v/v) 12 M HCl in 100 ml
acetic acid). After an incubation of 60 minutes at 60°C the absorbance was measured at 585
nm. An N-acetylglucosamine (GlcNAc, from Sigma) standard curve was used for the
calculation of the chitinase activity.
2.5.14 Yeast two-hybrid

Yeast two-hybrid experiments were performed using the Matchmaker System (Clontech). CHIA cDNA fragment lacking the signal peptide sequence (849 bp) was cloned into pGBK7 or pGADT7 (Clontech). Plasmids were transformed into S. cerevisiae using a lithium acetate/single-stranded carrier DNA/polyethylene glycol method (Gietz and Woods, 2002). The cultivation of the yeast cells was performed as described previously (Bergman, 2001; Kaiser, 1994). After 4 to 5 d of growth on vector-selective medium (CSM-LT), 12 independent clones in pools of four clones each were propagated in liquid vector selective medium and subsequently diluted to the same optical density. Of the three pooled cultures, 7.5 µl of a serial dilution was dropped on vector- and interaction-selective medium (CSM-LTA) and incubated at 28°C. At day 3, the growth of the clones was monitored.

2.6 Bioassays

2.6.1 Infection with Pseudomonas syringae

For the bacterial infection assay each Pseudomonas strain was diluted with 10mM MgCl₂ to a density of 1 x 10⁴ 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, 2 and 4 days post infection. For the quantification infected leaves were harvested (2 leaves at 0 dpi and 3 leaves at 2 or 4 dpi) and washed for one minute in both 70 % (v/v) EtOH and water. Afterwards 2 leaf discs/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.

2.6.2 Infection with Alternaria brassicicola

Alternaria brassicicola spores used for the infection assays were obtained as published previously (Thomma et al., 1999b). Leaves to be tested were drop-inoculated with six 5µl droplets of aqueous spore solution (5 x 10⁵-1 x 10⁶ spores/ml) if not mentioned otherwise. Two leaves per plant and a minimum of 8 plants per line were infected. To avoid positional effects plants of different lines were randomly distributed in the tray and incubated at 100% relative humidity. Fungal growth was scored after 7-14 days by symptom classification: 1 (no symptoms), 2 (light necrotic lesions), 3 (severe necrotic lesions), 4 (spreading of lesions beyond infection site), 5 (whole leaf affected) and 6 (sporulation of the fungus) and a disease index (Kemmerling et al., 2007) was calculated.
2.6.3 Infection with *Botrytis cinerea*

Cultivation of *Botrytis* and the preparation of spores was performed as described in Thomma et al. (1999a). *Botrytis cinerea* spores were diluted with PDB medium to a final density of 5 x 10^5 spores/ml. The leaf surface was drop-inoculated with two 5µl droplets of spore solution if not mentioned otherwise. Two leaves per plant and a minimum of 8 plants per line were used for infection. The development of symptoms was monitored 2-3 days post infection.

2.6.4 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 seedlings 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 200µl medium/well, 48er 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 following concentrations: 1 µM flg22 and 100 µg/ml chitin, PGNs and muropeptides derived from PGN.

2.6.5 Microarray analysis

Microarray experiments were performed on *Arabidopsis thaliana* Col-0 plants and on the T-DNA insertion line *cerk1-2*. Leaves of adult plants were infiltrated with 100 µg/ml Xcc PGN or water as a control and analyzed after 6 h. RNA was extracted as described in 2.4.6 and profiled using the NimbleGen DNA microarray (*A. thaliana* Gene Expression 12x135K Array TAIR 9.0) according to the manufacturer’s protocol (Roche). Three independent experiments (biological replicates) were performed. Probe signal values were subjected to the quantile normalization (Bolstad et al., 2003) using all the arrays. Normalized probe signal values were subjected to the robust multi-array average (RMA) summarization algorithm (Irizarry et al., 2003) to obtain the expression level values of the genes. Results were analyzed by the following linear model using the lmFit function in the limma package in the R environment:

\[
\text{log2 (expression level value)} \sim \text{genotype:treatment + replicate}
\]

The eBayes function in the limma package was used for variance shrinkage in calculation of the p-values and the Storey’s q-values were calculated using the q-value function in the q-value package (Storey and Tibshirani, 2003) from the p-values.
2.6.6 pH assay
Medium alkalization in At cell culture upon PAMP treatment was performed as described previously (Gust et al., 2007). The changes in pH were monitored and recorded by the Observer II program (Brainchild Electronics Co., Ltd, Taipei, Taiwan).

2.7 Microscopy and Histochemistry

2.7.1 Confocal 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 PlanApo H$_2$O objective. The Software LCS Lite Version 2.61 was used for the processing of the images. As membrane-selective dye, 2 nM FM4-64 (diluted in ddH$_2$O, Invitrogen), was infiltrated into the leaves shortly before the microscopical analysis. Concave plasmolysis was induced by infiltrating the leaves with 850 mM NaCl, 0.01 % (v/v) Silwet L-77 prior microscopy.

2.7.2 Aniline blue stain
The induction of callose deposits upon PAMP treatment was analyzed by aniline blue (water blue) staining (Gómez-Gómez et al., 1999). Leaves were infiltrated with the different PAMPs and incubated for 24 hours and subsequently incubated with a fixing solution (1% (v/v) glutaraldehyde; 5mM citric acid; 90mM Na$_2$HPO$_4$; pH7.4) again for 24 hours at RT. After fixation the leaf tissue was bleached with 100% (v/v) EtOH for 1-2 days. The leaves were then transferred to 50% (v/v) EtOH and afterwards equilibrated in 67 mM K$_2$HPO$_4$ (pH 12.0) and finally stained for 1 h at RT in 0.1% (w/v) aniline blue dissolved in 67 mM K$_2$HPO$_4$ (pH 12.0). The stained leaves were transferred to a microscopic slide in 70% (v/v) glycerol and 30% (v/v) staining solution and examined under UV epifluorescence.

2.7.3 Trypan blue stain
The trypan blue stain was used to visualize dead cells and fungal structures after infections with necrotrophic fungi. Infected leaves were treated with for 1 minute with trypan blue stain (10 ml lactic acid, 10 ml 100% glycerol, 10ml Aqua-Phenol, 10 ml ddH2O, 80 ml EtOH and 300 mg Trypan blue) and afterwards bleached with a 1mg/ml chloral hydrate solution.
2.7.4 GUS stain
The transgenic plants bear a cassette containing the uidA (gusA) gene under the control of a promoter of interest (Jefferson et al., 1987). For the histochemical detection of the enzymatic activity of the reporter gene β-glucuronidase (GUS) leaves were vacuum infiltrated with a staining solution containing 50 mM sodium phosphate buffer pH 7.0, 0.021% (w/v) K₄Fe(CN)₆, 0.016% (w/v) K₃Fe(CN)₆, 1 mM EDTA pH 8.0, 0.5% Triton X-100 (v/v) and 0.05% (w/v) 5-bromo-4-chloro-3-indolyl β-D-glucuronide (X-Gluc). The leaves were incubated overnight at 37°C and subsequently chlorophyll was removed by shaking in 100% EtOH. The stain was examined under the microscope.

2.8 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). The significance for the differences of a data pair was calculated using the t-test.
3 Results

The aim of this work was to gain more insights into the perception system of the bacterial PAMP peptidoglycan in the model plant *Arabidopsis thaliana*. In vertebrates and mammals the action of PGN as PAMP and the corresponding perception systems have been extensively analysed (Dziarski and Gupta, 2005a; Dziarski and Gupta, 2006; Girardin and Philpott, 2004; Royet and Dziarski, 2007), whereas *in planta* peptidoglycan was only lately described as a novel elicitor of the innate immunity (Erbs et al., 2008; Gust et al., 2007).

3.1 Isolation and analysis of PGN from *P. syringae* and other bacteria

In previous studies peptidoglycan from the Gram-positive bacteria *Staphylococcus aureus*, as well as the Gram-negative bacteria *Escherichia coli* were shown to induce typical plant defense reactions, such as induction of defense-related gene expression, medium alkalization and accumulation of phytoalexins (Gust et al., 2007). Gram-positive bacteria are underrepresented among the plant-pathogenic bacteria, however it has been reported that *Streptomyces* spp. and also *S. aureus* are able to infect plants, including *Arabidopsis thaliana* (Joshi et al., 2007; Prithiviraj et al., 2005). The Gram-negative bacteria attacking plants include *Xanthomonas* spp., *Ralstonia solanacearum*, *Erwinia* spp. and the model plant pathogenic bacterium *Pseudomonas syringae* pv. *tomato* DC3000 (*Pto* DC3000) (Boch and Bonas, 2001). Since *Pto* DC3000 is routinely used in bacterial infection assays for monitoring effects of *Arabidopsis* mutants lines in bacterial resistance, it was logical to analyse the properties of its peptidoglycan as elicitor of the plant innate immunity. Therefore, a protocol for the isolation of highly purified PGN from Gram-negative bacteria was established and preparations of *Pseudomonas syringae* PGN (*Pto* PGN) along with other PGNs were tested in different bioassays. First, the purity of *Staphylococcus aureus* PGN (*Sa* PGN), *Pto* PGN, *Bacillus subtilis* PGN (*Bs* PGN), *Xanthomonas campestris* pv. *campestris* PGN (*Xcc* PGN) and muropeptides derived of *Xcc* PGN (Erbs et al., 2008) was analysed for protein contaminations using the sensitive silver staining method (Figure 3-1A). In all preparations no high molecular weight impurities could be observed. To monitor the immunogenic properties of the different PGNs in *Arabidopsis* plants, Col-0 seedlings were treated with 100 μg/ml PGN and the transcript accumulation of the immune marker gene *Flagellin-induced receptor kinase 1* (*FRK1*) was measured 6 hours post elicitation using quantitative RT-PCR (RT-qPCR, Figure 3-1B). *FRK1* is transcriptionally up-regulated relatively early upon treatment with several MAMPs, such as flg22, HrpZ (Harpin) and NPP1 (necrosis-inducing
Results

Phytophthora protein 1) (Boudsocq et al., 2010; He et al., 2006). Treatment of seedlings with the tested peptidoglycans led to a four to 10-fold higher FRK1 gene expression in comparison to water treatment. In addition to FRK1 expression, the expression of the defense-related gene pathogenesis related 1 (PR1) was also analysed in adult leaves upon PGN treatment using a transgenic pPR1::GUS reporter line (Shapiro and Zhang, 2001). All tested PGNs and also the Xcc muropeptide induced the PR1 promoter (Figure 3-1C). Water and flg22 treatments were used as negative and positive control, respectively. Pto PGN also triggers other early plant immune responses, as was observed in the medium alkalization assay with plant cell cultures (Figure 3-1D). In comparison to the rapid and transient flg22-induced pH shift, the PGN reaction is slower but still clearly measurable, as reported earlier (Gust et al., 2007). The deposition of the polysaccharide callose (β-1,3-glucan) between the plasma membrane and the plant cell wall as a rather late cellular response is triggered upon different abiotic or biotic stresses, such as wounding and microbial attack (Stone and Clarke, 1992), but also upon treatment with PAMPs (Gómez-Gómez et al., 1999). Upon infiltration of adult leaves with Pto PGN, Xcc PGN and Xcc muropeptides a weak but visible callose accumulation was observed (Figure 3-1E). As expected, the positive control flg22 induced a strong deposition of callose, whereas water treatment led to no callose accumulation.

The immunogenic properties of all tested Lys- and Dap-type peptidoglycans were generic suggesting that either there are several PGN recognition machineries present in Arabidopsis or that there is no discrimination between the different peptidoglycan types. The modification of the peptidoglycan either within the sugar backbone (acetylation) or peptide moiety (amino acid change) can change the immunogenic properties as reported by Erbs et al. (2008), but this did not seem to be the case for the examined PGNs. All in all, the PGN-mediated immune responses showed weaker levels when compared to the peptide PAMP flagellin, what might result from the polymeric structure of peptidoglycan and hence poor or delayed accessibility to the plant receptors at the plasma membrane.

Efforts of isolating peptidoglycan from the model phytopathogen Pseudomonas syringae pv. tomato DC3000 were profitable. The purified Dap-type Pto PGN was able to trigger both early and late defense responses (Figure 3-1). In addition, the usage of the highly pure Pto PGN in elicitation assays contributed to the identification of AtLYM3, a membrane-tethered LysM protein, as a putative PGN receptor protein (Willmann, 2011). Thus, the collection of plant immunity stimulating peptidoglycans is increased by the Dap-type PGN from P. syringae.
Results

Figure 3-1: Peptidoglycan induces defense responses in *Arabidopsis thaliana*

(A) The purity of peptidoglycan and muropeptides was analysed by silver staining. 100 µg of different PGNs or muropeptides (*Staphylococcus aureus*, *Sa*; *Pseudomonas syringae* pv. tomato DC3000, *Pto*; *Bacillus subtilis*, *Bs*; *Xanthomonas campestris* pv. *campestris*, *Xcc*; muropeptides from *Xanthomonas campestris* pv. *campestris* PGN, *murop Xcc*;) were separated by SDS-PAGE and the proteinous contaminations in the samples monitored by silver staining. (B) PGN-induced *FRK1* gene expression in Col-0. Seedlings were treated with water or 100 µg/ml PGN and subjected to RT-qPCR 6 hours post treatment. *EF1α* transcripts served for normalization, corresponding water controls were set to 1. Data represent means ± SD of three independent experiments with 4-6 seedlings/sample. (C) Induction of the reporter gene *PR1* upon PGN treatment. Adult leaves of transgenic *pPR1::GUS* plants were infiltrated with water, 100 µg/ml PGN or 1 µM flg22. Leaves were harvested 24 hours post treatment and stained for GUS activity. (D) Medium alkalinisation in Col-0 cell culture upon treatment with water, 100µg/ml PGN *Pto* or 1µM flg22. (E) Accumulation of callose deposition upon PGN treatment. Adult Col-0 leaves were infiltrated with water, 100 µg/ml PGN or 1 µM flg22 and stained with aniline blue 24 hours post treatment. Under UV light the callose deposits appear as light blue dots (upper panel). Lower panels show a light image of the leaf tissue.
3.2 Identification of putative PGN receptor(s) among the LysM-RLKs

Specific LysM domain proteins have been characterised in the binding and recognition of carbohydrate molecules structurally similar to peptidoglycan. In rice and Arabidopsis, the fungal carbohydrate elicitor molecule chitin is perceived by the receptor proteins OsCEBiP, OsCERK1 and AtCERK1, respectively (Kaku et al., 2006; Miya et al., 2007; Wan et al., 2008). Furthermore, LysM receptor proteins in the legumes Medicago truncatula and Lotus japonicus recognize bacterial lipochitooligosaccharides thereby establishing plant root symbioses with soil-borne rhizobacteria (Limpens et al., 2003; Radutoiu et al., 2003). In Arabidopsis thaliana, there are five genes encoding LysM receptor-like kinases (LYKs, Figure 3-2A). Due to the similarity between the LysM receptor proteins in the different plant species and the structural similarity between peptidoglycan, chitin and lipochitooligosaccharide, a targeted reverse-genetics approach was chosen to analyse the members of the Arabidopsis LysM receptor kinase family more in detail regarding peptidoglycan recognition and bacterial resistance. The predicted protein domain structures of the five LYKs harbour a signal peptide for secretion (SP), a transmembrane domain (TM), a serine/threonine protein kinase domain (PK) and one or several lysin motifs, with the exception of LYK2 (Figure 3-2B). The analysis of the absolute expression of the LYK2 gene in Col-0 leaf tissue revealed extremely low values in comparison to the other members of the LYK family (data not shown, ATGenExpress Initiative). LYK2 displayed relatively high expression levels in the 1st node and the pedicel of the flower, whereas CERK1 (LYK1), LYK3, LYK4 and LYK5 were strongly expressed in leaves, suggesting that the four other LYK members possibly play a more important role within innate immunity to leaf pathogens such as Pto. Changes in the expression profiles of single genes upon different stimuli can give indications for an involvement of the gene products in corresponding signaling pathways or cellular mechanisms. Therefore, the ATGenExpress microarray data for CERK1, LYK3, LYK4 and LYK5 was analysed upon infection with different strains of Pseudomonas syringae (Figure 3-2C). CERK1 expression was first slightly induced upon infection with the virulent Pto DC3000 (2h sample) but was already suppressed after 6 hours likely through effector-triggered suppression. After 24 hours CERK1 expression showed control levels again. Infection with the type III secretion system-deficient strain Pto DC3000 hrcC− or the non-host Pseudomonas syringae pv. phaseolicola (Pph) led to enhanced CERK1 transcript levels at all investigated data points. In contrast, LYK3 transcript levels were strongly suppressed upon infection with the three tested Pseudomonas strains and at all data points. The expression profile of LYK4 resembled the one of LYK3 upon infection with Pto DC3000 and Pph, and the one of CERK1 upon Pto DC3000 hrcC− infiltration. The expression of the LYK5 gene showed a similar pattern than CERK1 gene upon treatment with the three tested bacteria, however with a lower overall induction.
Alterations in the gene expression upon infection with bacterial pathogens suggest a role for the *Arabidopsis* LysM receptor kinases in the plant innate immunity. Hence, selected LYK family members were in the following characterised more in detail.

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**Figure 3-2: Arabidopsis LysM receptor-like kinase family**

(A) A multiple sequence alignment of the full-length protein sequences of the five LysM-RLK (LYK) members using the ClustalW2 algorithm. (B) The predicted domain structures of the LYK proteins as annotated in the databases PFAM and SMART. Signal peptide, SP; LysM, Lysin motif; Transmembrane domain, TM; Protein kinase domain, PK. (C) Leaves of adult *Arabidopsis* Col-0 plants were infiltrated with the virulent *Pto* DC3000, the T3SS-deficient *Pto* DC3000 hrcC- or the avirulent *Pseudomonas syringae pv. phaseolicola* (Pph) strain (10^8 cfu/ml) or 10 mM MgCl_2 as control. Leaves were harvested at indicated time points and total leaf RNA was used for microarray analysis. Data for *CERK1* (At3g21630), *LYK3* (At1g51940), *LYK4* (At2g23770) and *LYK5* (At2g33580) result from experiments performed within the ATGenExpress initiative (http://www.arabidopsis.org/info/expression/ATGenExpress.jsp). The gene transcription in the control treatment was set to 1.
3.2.1 Analysis of the LYK T-DNA insertion lines

For the functional analysis of the LYKs T-DNA insertion lines were obtained from Volker Lipka (Göttingen) and NASC (Nottingham Arabidopsis Seed Collection, UK). In the beginning of this work only one of the LYK members, CERK1, was characterised having a biological role as a chitin elicitor receptor kinase. The knock-out line for CERK1, cerk1-2 was published by Miya et al. (2007). The exon-intron structure of the CERK1 gene and the position of the T-DNA insertion in the 11th intron are depicted in Figure 3-3A. The homozygous genotype of the cerk1-2 line and the absence of full-length CERK1 transcript was confirmed by genotyping PCR and semi-quantitative RT-PCR, respectively (Figure 3-3A). Moreover, protein expression was absent in cerk1-2 (Gimenez-Ibanez et al., 2009). For both LYK3 and LYK5 two independent T-DNA lines and double mutants (lyk3-1 lyk5-1; lyk5-1 lyk3-1) were available (Volker Lipka). The gene models and the T-DNA insertions for LYK3 and LYK5 are illustrated in Figure 3-3B and C. As for CERK1, also LYK3 and LYK5 T-DNA insertion lines were homozygous and showed no residual transcript as controlled by genotyping and transcript analysis (Figure 3-3B, C).
Figure 3-3: Characterisation of CERK1, LYK3 and LYK5 and their mutants

Gene models of CERK1 (A), LYK3 (B) and LYK5 (C) including the positions of the T-DNA insertions. Exons and introns are indicated by black bars and black lines, respectively. 5’- and 3’-UTR regions are represented by grey bars and the T-DNA insertions by grey triangles. Leaf genomic DNA was isolated and genotyping PCRs were performed. Total RNA was isolated from leaves and transcribed into cDNA for transcript analysis using semi-quantitative RT-PCR. EF1α-s and EF1α-as primers were used to amplify the transcript of the house-keeping gene EF1α. (A) Genotyping of the cerk1-2 mutant was done with the primers 580H03LP and 580H03RP (WT-PCR) and 580H03LP and Gabi-Kat-Lba (Lba-PCR). CERK1 transcript was amplified with the primers 580H03LP and 580H03RP. (B) Lyk3 mutants were genotyped with the primers 640374LP and 640374RP (WT-PCR) and 640374LP and SALK-Lba (Lba-PCR). LYK3 transcript was amplified with primers 640374LP and 640374RP (lyk3-1) and N654015F2 and N660797R2 (lyk3-2). (C) Lyk5 mutants were genotyped using the primers GT7089LP and GT7089RP2 (WT-PCR, lyk5-1), 631911LP and 631911RP (WT-PCR, lyk5-2), Ds3-1 + GT7089RP2 (Lba-PCR,
Results

lyk5-1) and Salk-Lba and 631911RP (Lba-PCR, lyk5-2). LYK5 transcript analysis was done with the primers GT7089LP and GT7089RP2.

T-DNA insertion lines for the LYK2 and LYK4 genes were also analysed with respect to homozygosity. The T-DNA insertion in the lyk2-1 line is positioned in the first exon and the T-DNA insertion in the lyk4-1 at position of 1615 bp (Figure 3-4). However, both lines turned out to be heterozygous as for some individual plants still WT-PCR product could be observed (Figure 3-4A, B). Candidate plants (lyk2-1, lane 3; lyk4-1, lanes 2 and 3) were chosen for production of seeds and the next generation has to be genotyped again to be subsequently used for further analyses.

Figure 3-4: Characterisation of LYK2 and LYK4 and their mutants

Gene models of LYK2 (A) and LYK4 (B) including the positions of the T-DNA insertions. Exons and introns are indicated by black bars and black lines, respectively. 5' and 3' UTR regions are represented by grey bars and the T-DNA insertions by grey triangles. Genotyping analysis was performed as described in Figure 3-3. (A) Genotyping of the lyk2-1 mutant was done with the primers 512441LP and 512441RP (WT-PCR) and Salk-Lba and 512441RP (Lba-PCR). (B) The lyk4-1 mutant was genotyped with the primers 2g23770F2 and 2g23770R1 (WT-PCR) and 2g23770F2 and Gabi-Kat-Lba (Lba-PCR).

3.2.1.1 Phenotypic analysis of lyk3 and lyk5 single and double mutants

The T-DNA insertion mutant cerk1-2 was published to have no visible phenotypic alteration from wild type (Col-0) plants (Gimenez-Ibanez et al., 2009). The phenotypes of the T-DNA insertion line lyk5-1 and the double mutants lyk3-1 lyk5-1 and lyk5-1 lyk3-1, however, differ from the phenotype of the corresponding ecotype. The lyk5-1 (GT7089) mutant has the same
leaf morphology than the WT plant Ler, but shows an early flowering transition, whereas the \textit{lyk3-1} (N654015) mutant resembles the Col-0 ecotype both in its morphology and flowering behavior (Figure 3-5B). Whether the observed developmental differences in the \textit{lyk5} mutant line result from \textit{LYK5} deletion or additional 2\textsuperscript{nd} site insertions should be further investigated in future experiments. The double mutant lines \textit{lyk3-1 lyk5-1} (LGK2-6-1-2, in Col-0) and \textit{lyk5-1 lyk3-1} (LGK2-8-1-2, in Ler) display upward curled leaves, what might indicate defects in polarity and cell division (Liu et al., 2010). The production of seeds was unaltered in all T-DNA lines (data not shown). However, since changes in the reproduction process or leaf morphology should not negatively affect plant innate immunity, the lines were considered as suitable for further experiments. The \textit{lyk2} and \textit{lyk4} mutant lines were so far not included in the study.

![Figure 3-5: Phenotypes of \textit{lyk3} and \textit{lyk5} single and double mutants](image)

Representative five week-old mutant plants of \textit{lyk3-1} (N654015), \textit{lyk5-1} (GT7089), \textit{lyk3-1 lyk5-1} (LGK2-6-1-2, Col-0), \textit{lyk5-1 lyk3-1} (LGK2-8-1-2, Ler) and the corresponding ecotypes (Col-0 and Ler) were photographed from the top (A) and the side (B) view.
3.2.2 Role of LYKs in fungal resistance

The importance of one of the LysM receptor-like kinases, CERK1, in the fungal resistance has been reported. Knock-out mutants displayed more fungal growth than WT plants upon infection with the biotrophic powdery mildew fungal pathogen *Erysiphe cichoracearum* (Wan et al., 2008) or larger lesions upon infection with the necrotrophic fungus *Alternaria brassicicola* (Miya et al., 2007). Microarray data upon infection with the necrotrophic fungal pathogen *Botrytis cinerea* (ATGenExpress initiative) suggest weak upregulation of the CERK1 and LYK5 gene expression, whereas the expression of LYK3 is suppressed (Figure 3-6E). To gain additional information about the possible influence of the CERK1, LYK3 and LYK5 genes in resistance to fungal pathogens, the aggressive necrotrophic fungus, *Botrytis cinerea*, was employed.

*Botrytis cinerea* spore solution was droplet inoculated on the one half of the leaf and two leaves per plant were infected (2.6.3). The infection symptoms were monitored 2 and 3 days post infection. The infection site was already visible after two days showing necrotic tissue around the spore drop (data not shown) and after 3 days the necrotised leaf area was expanded (Figure 3-6A). All lyk mutant lines exhibited similar symptom development in comparison to WT plants. For disease analysis on a cellular level, dead plant cells and fungal hyphae were visualised using Trypan blue staining (2.7.3). Two days post infection all tested lines including Col-0 showed relatively loose fungal hyphae within the infection site, which was surrounded by an intensively stained ring, the cell death zone (Figure 3-6B). One day later (3 dpi) the cellular symptoms of the infection were more dramatic, i.e. the observed fungal hyphae were denser, however there were no differences between the lines (Figure 3-6C). Also the measurement of the lesion size three days post infection delivered no differential data, all tested lyk mutants behaved like the wild type (Figure 3-6D). Although the cerk1-2 mutant showed a tendency of larger lesions and the lyk3 and lyk5 mutants of smaller lesions than the WT plants, these changes were not significant.

In summary, upon infection with *Botrytis cinerea* no significant mutant phenotype for the tested LYK genes was observed.
Results

Figure 3-6: Infection of lyk mutants with *Botrytis cinerea*

Five week-old plants were infected with the necrotrophic fungus *Botrytis cinerea*. 5 µl spore suspension of 5 x 10⁵ spores/ml was drop-inoculated on the one half of the leaf; two leaves per plant were infected and analysed for symptom development after 2 or 3 days post infection. (A) Visible symptoms after 3 dpi. Microscopic analysis of the infection site and fungal hyphae visualised by Trypan blue stain 2 dpi (B) and 3 dpi (C). (D) Measurement of the lesion size 3 days post infection. Shown are means and standard errors (n=16). (E) Leaves of adult Arabidopsis Col-0 plants were infected with *Botrytis cinerea* (10⁵ spores/ml) or PDB medium as control. Leaves were harvested at indicated time points and total leaf RNA was used for microarray analysis. Data for CERK1 (At3g21630), LYK3 (At1g51940) and LYK5 (At2g33580) genes result from experiments performed within the ATGenExpress initiative (http://www.arabidopsis.org/info/expression/ATGenExpress.jsp). The gene transcription in the control treatment was set to 1.
3.2.3 Influence of LYKs on bacterial resistance

Defects in the recognition process of pathogenic microbes can lead to enhanced virulence of the pathogens. For instance, FLS2-deficient plants, which no longer sense the bacterial PAMP flagellin, are more vulnerable to phytopathogenic bacteria (Zipfel et al., 2004). Also the deletion of the co-receptor of FLS2, BAK1, leads to enhanced disease symptoms upon bacterial infection in the mutant background (Kemmerling et al., 2007). In the following the homozygous T-DNA lines of the LYK family members were assayed for their bacterial resistance properties using different strains of the hemibiotrophic bacterium *Pseudomonas syringae*. Interestingly, a bacterial susceptibility phenotype of two independent *cerk1* mutant alleles reported recently implicated a role for *CERK1* in the perception of a bacteria-derived MAMP (Gimenez-Ibanez et al., 2009). Similar hypersusceptibility was observed for the tested *cerk1-2* mutant upon infection with the virulent *Pto* DC3000 4 days upon infiltration of the bacteria at a dose of $10^4$ cfu/ml (Figure 3-7A, left diagram). Also the less-virulent mutant strain *Pto* DC000 ΔavrPto/PtoB and the type III secretion system (TTSS)-deficient mutant *Pto* DC3000 hrcC- showed more growth in the mutant than in the WT plants as depicted in Figure 3-7A (middle and right diagram). Two independent *LYK3* knock-out mutants, *lyk3-1* and *lyk3-2*, were also subjected to infection assays with the same bacterial strains. Both mutant lines were more susceptible to *Pto* DC3000, whereas only *lyk3-1* allowed significantly more growth of the *Pto* DC3000 ΔavrPto/PtoB strain (Figure 3-7B, left and middle diagram). Despite of repetitive experiments, so far no significant differences in the growth of *Pto* DC3000 hrcC- strain could be observed for the *lyk3* mutants in comparison to the WT (Figure 3-7B, right diagram). The two *lyk5* mutants, *lyk5-1* and *lyk5-2*, displayed no measurable changes to the corresponding WT upon infection with either *Pto* DC3000 or *Pto* DC3000 ΔavrPto/PtoB (Figure 3-7C). Infection with the *Pto* DC3000 hrcC- strain was up to date only performed with the *lyk5-2* mutant, but it showed similar levels of bacterial growth than the WT as shown in Figure 3-7C (right diagram). Although single *lyk3-1* mutant did show a bacterial growth phenotype, the double mutants for the *LYK3* and *LYK5* genes behaved like WT plants upon infection with either virulent and or less-aggressive mutant strains (Figure 3-7D). However, crosses of two different ecotypes might be problematic and lead to such ambiguous results. Nevertheless, the bacterial infection experiments on the single mutants suggest that additionally to *CERK1* also *LYK3* as a member of the LYK family might have a role in resistance to bacterial pathogens.
Figure 3-7: Infection of lyk mutants with Pseudomonas syringae strains

Bacterial infections of Pto DC3000 (left diagram), Pto DC3000 ΔavrPto/PtoB (middle diagram) and Pto DC3000 hrcC⁻ (right diagram) in the cerk1-2 mutant (A), lyk3 mutants (B), lyk5 mutants (C) and lyk3 lyk5 double mutants (D). Growth of bacteria was determined 4 days post infiltration with 10⁴ cfu/ml. Data represent means and standard errors (n=6/genotype/data point). Statistical significance compared to wild type (p ≤ 0.05, Student’s test) is indicated by asterisks. Shown are representative of at least 3 independent experiments.
3.2.4 Analysis of peptidoglycan responsiveness in *lyk* mutants

The structure of the PAMP chitin resembles the glycan backbone structure of peptidoglycan and CERK1 or the ectodomain of CERK1 containing all three LysM domains have been shown to bind to the ligand chitin *in vitro* (Iizasa et al., 2010; Petutschnig et al., 2010). In parallel, a LysM domain containing GPI-anchored protein, LYM3, was identified as a putative PGN-binding protein whereby the *lym3* mutant displays defects in PGN-triggered defense responses (Willmann, 2011). Since LYM3 lacks an intracellular signaling domain, a signaling partner might be found within the LYK family. The bacterial susceptibility phenotype observed for *cerk1* and *lyk3* mutants suggested that one or both of these receptor kinases might fulfill the missing part in the PGN recognition machinery. First, the responsiveness of the *cerk1*-2 mutant to PGN was tested. Adult leaves were treated with 100 µg/ml *Xcc* PGN, 1 µM flg22 or 100 µg/ml chitin and subjected to RT-qPCR three hours post infiltration to monitor induction of the defense-related gene *FRK1* (Figure 3-8A). Similar transcript accumulation was observed upon flg22-treatment for Col-0, *cerk1*-2 and the *lym3*-1 mutant, which was included as a control. In PGN-treated leaves the transcript level of *FRK1* was reduced in the *lym3* and *cerk1* mutant backgrounds when compared to WT. As expected, the chitin-response was strongly reduced in the *cerk1*-2 but not in the *lym3*-1 mutant (Figure 3-8A). The PGN responsivity of *cerk1*-2 was also analysed in seedlings and here the *lyk3*-1 mutant was also included into the assay. The *FRK1* transcript accumulation was monitored 6 hours post treatment with 100 µg/ml PGN *Pto* or chitin (Figure 3-8B). Whereas Col-0 and *lyk3*-1 seedlings were sensitive towards PGN treatment, in *lym3*-1 and *cerk1*-2 mutant seedlings PGN failed to induce the expression of *FRK1*. The chitin-response was defective only in the *cerk1*-2 mutant.

The elicitation experiments with peptidoglycan from two different Gram-negative bacteria indicate that of the so far tested *lyk* mutants only *cerk1*-2 mutant is defective in the peptidoglycan-triggered defense response.
3.2.4.1 The LysM-receptor kinase CERK1 mediates sensitivity to PGN

Microarray analyses were performed using the cerk1-2 mutant and the corresponding wild type to survey the effects of PGN elicitation on the global gene expression in Arabidopsis thaliana (in collaboration with Fumiaki Katagiri and Kenichi Tsuda, University of Minnesota). Leaves of five-week-old plants were infiltrated with either water or 100 µg/ml Xcc PGN and analysed 6 hours post treatment. In total 682 genes were either significantly induced or repressed by peptidoglycan in WT plants. Interestingly, the gene ontology (GO) analyses showed among the PGN-inducible genes an over-representation of those known to be induced upon stress (12 %) and biotic stimulus including bacterial infection (9.5 %) as depicted in Figure 3-9. The genes repressed by PGN-treatment showed a highly similar functional classification (data not shown). Among the affected genes were many genes implicated in plant defense including FRK1, ADR1 (Activated disease resistance 1, NBS-LRR protein), At1g51890 (LRR receptor-like kinase), PAD3 (Phytoalexin deficient 3, phytoalexin biosynthesis), NPR1 (Nonexpresser of PR genes 1, key regulator of SA-mediated SAR pathway), MYB7 (MYB domain protein 7, transcription factor) and MLO12 (Mildew resistance locus 12, disease resistance protein).
The global PGN-inducible gene expression in wild type plants was analysed using microarray experiments. PGN treatment was performed as described in Figure 3-8A and harvested after 6 h and extracted RNA was used for microarray analyses. Significantly induced or repressed genes by either treatment were chosen (682 genes; Storey’s q-value < 0.15) and functionally analysed using GO slim Classification for plants (GO Biological Process, TAIR). The genes were organized into sets according to broad GO ontology categories and percentage of the genes belonging to a specific biological process was calculated ([number of annotations to terms in this GOslim category x 100]/number of total annotations in this ontology) = %).

A strong deregulation of both up- and down-regulated genes was revealed in the cerk1-2 mutant when compared to Col-0 plants as visualised by a heat map in Figure 3-10A. The Michael-Eisen cluster analysis was facilitated for clustering of the affected genes. Similarly striking deregulation pattern of the gene expression in cerk1-2 mutant is visible when the log2 ratios of the PGN-treated transcript levels are plotted against water samples (WT, Figure 3-10B upper diagramm; cerk1-2, Figure 3-10B; lower diagram, red line indicates the regression line for WT). To verify the microarray data the expression levels of randomly selected genes (At1g51890, MLO12 and PAD3) were monitored using quantitative RT-PCR (Figure 3-10C, performed by Roland Willmann). All three tested genes show a strong reduction in PGN-induced accumulation of their transcripts in the cerk1-2 mutant in comparison to the WT. Additionally, to confirm that the PGN-insensitivity is not restricted to the one tested T-DNA insertion mutant, a second independent allele, cerk1-3 (Gimenez-Ibanez et al., 2009), which is in Ws-4 background was also tested and found also to be insensitive towards peptidoglycan (data not shown).
Results

Figure 3-10: CERK1 mediates PGN sensitivity

Microarray analysis of global PGN-inducible gene expression in wild type and cerk1-2 mutant plants. PGN treatment was performed as described in Figure 3-8A and harvested after 6 h and extracted RNA was used for microarray analyses. Significantly induced or repressed genes by either treatment were chosen (682 genes; Storey's q-value < 0.15). (A) Heat map comparing the changes in gene expression in the tested lines. Legend indicates values that correspond to the color scale; \( \log_{2} \) ratio (PGN/water). The responsive genes were clustered using Michael-Eisen cluster (uncentered correlation) and visualized by tree view. (B) The \( \log_{2} \) ratios of transcript levels observed PGN-treated Col-0 and cerk1-2 plants were plotted vs. water controls. The linear regression line indicates gene expression levels in Col-0 (black line in the upper diagram, red in the lower diagram) or cerk1-2 (black line in the lower diagram). (C) Transcript accumulation of Atig51890, MLO12 and PAD3 genes in PGN treated cerk1-2 and Col-0 are shown relative to those detectable in water-treated plants. Seedlings were treated with 100 µg/ml PGN Xcc and subjected to RT-qPCR as described in Figure 3-1B.
3.2.5 Analysis of potential redundancy among LYK genes

Gene families with several members are common in plants (Vandepoele and van de Peer, 2005). Individual genes may exhibit differential temporal expression patterns or they can be expressed in different tissues. However, these genes may also have redundant functions, with differences in dose requirements. Regarding the LYK gene family, the expression patterns indicate that LYK2 might have a very tissue-specific task in Arabidopsis. The other LYK members are strongly expressed in the leaf, one of the main infection sites of the plant, suggesting potentially redundant roles or completely different functions. The generation of double, triple or quadruple lyk mutants could help elucidating the roles of the single genes.

The data presented so far supported the importance of CERK1 and possibly also LYK3 in the bacterial resistance and in the first round of crossing the cerk1-2 mutant was crossed with both the lyk3-1 single mutant and also the lyk3-1 lyk5-1 double mutant (both mutants in Col-0 background). An additional benefit from the crossing of the cerk1-2 with the double mutant was to get an additional back-crossing in Col-0 background for minimising the Ler portion still left in the genotype of the double mutant. The crossing was successful for both combinations and the obtained offspring was monitored for heterozygous individuals by genotyping PCR analysis (data not shown). The F2 generation of the generated lyk triple mutant cerk1-2 lyk3-1 lyk5-1 contained some candidate individuals showing amplification products only in the Lba-PCR reactions, indicating that these plants could be fully homozygous for all three T-DNA insertions within the CERK1, LYK3 and LYK5 genes in the following generation (Figure 3-11; lane 1-2, 4-6). The phenotype of this newly generated triple mutant can in future be analysed with respect to PGN sensitivity and bacterial resistance. WT plants and the mutant lines cerk1-2, lyk3-1 and lyk3-1 lyk5-1 were included as controls in the genotyping analysis (Figure 3-11). As soon as the lyk4-1 mutant is homozygous it can also be used for a further crossing with the cerk1-2 lyk3-1 lyk5-1 triple mutant to generate a quadruple lyk mutant.

![Figure 3-11: Generation of triple lyk knock-out mutants](image)

Genotyping analysis of the triple mutant cerk1-2 lyk3-1 lyk5-1 in Col-0 background. Genomic DNA was isolated from leaves and genotyping PCR reactions were performed with the primer combinations described in Figure 3-3A-C.
3.3 The role of PGN hydrolysis in the PGN sensing process

Peptidoglycan recognition proteins (PGRPs) mediate beside the LRR-receptor proteins TLR2 and NOD1/2 recognition of PGN, and are conserved between insects and higher animals, including humans (Bischoff et al., 2006; Cho et al., 2005; Dziarski and Gupta, 2010; Kurata, 2010). Among these PGRPs are both PRRs as perception and signaling molecules and proteins with PGN hydrolysing properties (Bischoff et al., 2006; Dziarski and Gupta, 2010; Gelius et al., 2003; Kurata, 2010; Wang et al., 2003). PGRPs act together with other innate immunity proteins to effectively combat pathogens. One example for such innate immunity protein is lysozyme, which is able to lyse bacteria by massive PGN hydrolysis and thus boosts the antibacterial defense (Callewaert and Michiels, 2010). It has been also shown to synergistically function together with the human PGLYRP1 and to contribute to the innate immunity by killing Gram-negative bacteria trapped by the neutrophils (Cho et al., 2005). Besides the direct bacteriolytic activity, it has been suggested that the PGN fragments released by the lytic activity of lysozyme have an impact on bacteria-host interactions by modulating the activation of the immune response and inflammation pathways (Chaput and Boneca, 2007). Such processing of polymeric carbohydrates derived of pathogenic cell walls and release of immunogenic fragments by host enzymes has also been reported in plants. In the legume plant soybean a β-glucan-binding protein (GBP) harbours two carbohydrate-active protein domains; on the one hand a binding site for a β-glucoside ligand derived from the cell wall of the oomycete Phytophthora sojae and on the other hand an endoglucanase activity (Fliegmann et al., 2004; Mithöfer et al., 2000). Hence, during contact with Phytophthora hyphae the intrinsic 1,3-β-glucanase activity of the GBP produces soluble oligoglucoside fragments enriched in motifs that are ligands for the high-affinity binding site present in the same protein (Fliegmann et al., 2004). However, GBP does not contain any signal transmitter domains leaving the question open how the defense mechanism triggered by β-glucan perception is transduced across the plasma membrane.

In order to find a connecting link between PGN processing and receptor-mediated PGN recognition in planta the general plant PGN hydrolase properties as well as the immunogenic nature of PGN fragments were examined. Previous studies demonstrated that soluble oligomeric PGN fragments derived from peptidoglycan of the Gram-negative phytopathogen Xanthomonas campestris pv. campestris (Xcc muropeptides) were able to induce defense reactions in Arabidopsis thaliana (Erbs et al., 2008). This muropeptide mixture was used to infiltrate adult leaves of the cerci1-2 and lym3-1 mutants and Col-0 plants. The following RT-qPCR analysis revealed that the muropeptide-induced expression of the immune marker genes FRK1 and MLO12 was clearly reduced in both mutant lines (Figure 3-12A, B). The
results indicate that soluble muropeptides derived from polymeric PGN are sensed in a receptor-mediated manner via LYM3 and CERK1.

![Figure 3-12: Muropeptide-induced gene expression requires LYM3 and CERK1](image)

Leaves were infiltrated with 100 μg/ml muropeptides derived from PGN Xcc and analysed 3 hours post treatment for the transcript accumulation of the defense-related genes FRK1 (A) and MLO12 (B) using RT-qPCR as described in Figure 3-8A.

### 3.4 Identification of a putative PGN hydrolase in *Arabidopsis thaliana*

Lysozyme-encoding sequences are restricted to animal genomes. Nevertheless, a subset of plant enzymes (hevamine-like chitinases) have been reported to possess PGN hydrolase activity *in vitro* (Bokma et al., 1997; Park et al., 2002). Interestingly, also some lysozymes are able to hydrolyse N-acetyl glucosamine substrates and have thus chitinase activity (Miyauchi et al., 2006; Nilsen et al., 1999; Xue et al., 2004). Based on these informations it was likely to find a putative PGN hydrolase among the *Arabidopsis* chitinases. In *Arabidopsis*, 24 chitinases are annotated and they are subdivided into 5 classes according to their sequence and structural features (Passarinho and De Vries, 2002). The arrangement of the chitinases into classes I to V is illustrated by the alignment of the full-length amino acid sequences using the ClustalW2 algorithm (Figure 3-13). The class III is represented by only one member, CHIA (At5g24090).
Results

Figure 3-13: Sequence alignment of the *Arabidopsis* chitinases

A multiple sequence alignment of the 24 annotated members of the chitinase family in *Arabidopsis thaliana* using the ClustalW2 algorithm. Full-length amino acid sequences were aligned and subgroups (I-V) were classified according to Passarinho and De Vries (2002). *Arabidopsis CHIA* (At5g24090) represents the only member of class III.

### 3.4.1 Analysis of the class III chitinase CHIA

The rubber tree hevamine is a class III chitinase and has been reported to display additionally to chitinolytic activity also PGN hydrolysing properties (Bokma et al., 1997). A BLAST (Basic Local Alignment Search Tool) search using the protein sequence of *Hevea brasiliensis* hevamine revealed only one *Arabidopsis* chitinase, the putative acidic endochitinase CHIA, as homologous to the rubber tree enzyme. Alignment of these two protein sequences using the ClustalV algorithm resulted in approximately 70 % identity (see Figure 3-14A). The CHIA protein contains the conserved catalytic core with an glutamic acid residue (E-157; Figure 3-14A, blue box) and almost all residues essential for substrate-binding (Figure 3-14A, blue asterisks) as shown for the rubber tree hevamine (Terwisscha van Scheltinga et al., 1996). The protein sequence of CHIA was also analysed for the
presence of a signal peptide and subcellular protein localisation using the SignalP 3.0 program and Cell EFP Browser (BAR, The Bio-Array Resource for Plant Biology) and a high probability was calculated for a putative secretion signal (amino acids 2-22) targeting CHIA to the extracellular space (Figure 3-14A, B). The calculated isoelectric point of CHIA (pI 9.3) (The Arabidopsis Information Resource, TAIR) is similar to that of *Hevea brasiliensis* hevamine (Tata et al., 1983).

Due to the presence of all essential enzymatic residues known to date to be required for chitinase activity and the predicted secretion into the plant apoplast, the endochitinase CHIA makes a perfect candidate for a putative PGN hydrolase.

Figure 3-14: CHIA protein sequence and annotated features
(A) Alignment of the full-length amino acid sequences of the Arabidopsis thaliana CHIA (At5g24090) and its homolog Hevea brasiliensis HevamineA using the ClustalV algorithm. The sequences share 70 % identity, grey boxes underline differences and the red box depicts the predicted secretion signal. The amino acid sequence after the secretion signal is annotated as acidic endochitinase domain (glycoside hydrolase family 18, EC 3.2.1.14). The blue box indicates the conserved catalytic site with a glutamic acid residue and the blue asterisks the substrate-binding sites (Terwisscha van Scheltinga et al., 1996). (B) Secretion signal prediction for the CHIA protein sequence performed with the SignalP 3.0 program using neural networks (NN) and hidden Markov models (HMM) trained for eukaryotes. The most likely cleavage site is between aa pos. 22 and 23 (SLS-KP).

3.4.2 Expression pattern of the CHIA gene upon biotic stress

According to the publicly available microarray data from the ATGenExpress initiative the CHIA gene is expressed at relatively low levels in most plant tissues, including leaves of different stages (data not shown). In certain stages of the flower development the CHIA transcript is abundantly present in anthers. In addition, studies using promoter-GUS reporter lines showed that CHIA was also expressed in hydathodes and stomatal guard cells (Samac and Shah, 1991). Upon infection with the fungal pathogen Rhizoctonia solani the expression of CHIA was induced around the lesion sites (Samac and Shah, 1991). Furthermore, CHIA expression was inducible by different strains of the plant phytopathogenic bacterium Pseudomonas syringae (ATGenExpress, Figure 3-15A). Whereas the induction of CHIA expression was less than two-fold after 24 hours upon infection with the virulent Pto DC3000 strain in comparison to control treatment, infection with the non-virulent mutant strain Pto DC3000 hrcC yielded nearly 2.5-fold induction (Figure 3-15A). The infection of Arabidopsis leaves with the non-host pathogen Pph resulted in a relatively fast and strong induction of CHIA transcription with already 2.2-fold induction after 6 hours and 2.4-fold induction upon 24 hours when compared to the mock treatment (Figure 3-15). The expression of CHIA was also induced 48 hours after inoculation with the necrotrophic fungi Botrytis cinerea as depicted in Figure 3-15B.
Microarray analyses were performed within the ATGenExpress initiative using Col-0 plants infected with different *Pseudomonas syringae* strains (A) and *Botrytis cinerea* (B) as described in detail in Figure 3-2C (*P.syringae* infection) and Figure 3-6E (*B.cinerea* infection).

To analyse the mode of *CHIA* expression in more detail transgenic promoter-GUS reporter lines were generated containing the *β-glucuronidase* (GUS) reporter gene under the control of a 2 kb long fragment of the *CHIA* promoter (see 2.4.15). One half of the leaves of homozygous plants was infiltrated with either water or 100 µg/ml PGN Bs, chitin or 1µM flg22 and stained for GUS activity 24 hours post treatment (Figure 3-16A). Transgenic *pPR1::GUS* reporter plants served as control (Shapiro and Zhang, 2001). No GUS staining was observed in the water-treated leaves of both reporter lines. Also peptidoglycan from *Bacillus subtilis* failed to induce the *CHIA* promoter and the treatment with chitin resulted in very weak activation of the promoter (Figure 3-16A, second and third upper panel from left). However, elicitation with the proteinaceous bacterial PAMP flg22 led to a visible GUS staining indicating activation of the *CHIA* promoter (Figure 3-16A, upper right panel). The *pPR1::GUS* line showed similar GUS staining pattern as the *pCHIA::GUS* line but the *PR1* promoter was slightly more sensitive showing visible activation also upon PGN and chitin treatments (Figure 3-16A, second and third lower panel from left). The flg22-treatment induced a strong GUS staining of infiltrated half of the leaf (Figure 3-16A, lower right panel). The PAMP treatment led in general to a local induction of the promoters, thus GUS staining was only detected within the infiltrated leaf tissue (Figure 3-16A). The ability of the nectrotrophic fungi *Alternaria brassicicola* and *Botrytis cinerea* to induce the *CHIA* promoter was also explored. 48 hours after infection of *pCHIA::GUS* leaves with *Alternaria* weak GUS stain at the site of the drop-inoculation was observed (Figure 3-16B, second upper panel from right).
activation of the CHIA promoter was visible upon Botrytis infection, but also the PDB medium used as control induced the promoter in a weak manner (Figure 3-16B, third and fourth upper panel from left). The PR1 promoter showed stronger activation upon Alternaria and Botrytis infections than the CHIA promoter and especially the infection with Botrytis resulted in GUS activity beyond the inoculation site (Figure 3-16B, lower panel). Finally, the promoter GUS reporter plants were also infected with Pseudomonas syringae strains to monitor effects of bacterial phytopathogens on the activity of the CHIA promoter. Here, infection with the virulent Pto DC3000 showed no induction of the CHIA promoter, when leaves were observed 24 hours post infection (Figure 3-16C, second upper panel from left). Pto DC3000 hrcC− and Pph strains both activated the CHIA promoter, however the GUS stain upon Pph-infection was clearly stronger than upon infection with Pto DC3000 hrcC− (Figure 3-16C, third and fourth upper panel from left). The pPR1::GUS line responded to all three Pseudomonas strains (Figure 3-16C, lower panel). Also, the bacterial infections resulted similar to the PAMP treatments only in a local promoter activation.

The expression of the CHIA gene is differentially regulated upon infection with fungal or bacterial pathogens. Whereas virulent Pseudomonas strains seem to suppress the CHIA expression, non-host bacterial pathogens like Pph and and also the bacterial PAMP flg22 induce it, thus indicating a possible relevance for CHIA within plant defense against biotic stress.
**Results**

Figure 3-16: CHIA expression analysis using a pCHIA::GUS reporter line

The expression of CHIA in transgenic pCHIA::GUS reporter plants was monitored 24 hours post treatment with indicated PAMPs (A), 48 hours upon infection with the necrotrophic fungi *Alternaria brassicicola* or *Botrytis cinerea* (B) and 24 hours post infection with different *Pseudomonas syringae* strains (C). The treatment of the leaves and used concentrations were described in Figure 3-1D (PAMP infiltration), Figure 3-2C (infection with *Pseudomonas*) and Figure 3-6E (Infection with *Botrytis*). The leaves were harvested after indicated time points and stained for GUS activity. The pPR1::GUS line containing the promoter of the defense-inducible gene *PR1* (Shapiro and Zhang, 2001) was used as control.
3.4.3 Analysis of the transgenic CHIA lines

Three independent T-DNA insertion lines for the CHIA gene were obtained from NASC and CSHL (Cold Spring Harbor Laboratory, NY, USA) for the characterisation of the importance of CHIA in resistance to bacteria and PGN perception. The chia-1 mutant line (WiscDsLox387C11) contains a T-DNA insertion ~500 bp upstream of the start codon in the promoter region, the T-DNA insertion in the chia-2 mutant (SALK_095362) is in the end of the third exon and the third mutant line, chia-3 (CSHL_ET14179), carries an Enhancer Trap (ET) transposon insertion within the first intron. The gene structure of CHIA and the positions of the insertions are depicted in Figure 3-17A. The genotyping analysis using gene- and insertion-specific primers could verify that all three chia mutant lines were homozygous for the corresponding insertions (Figure 3-17B). However, the subsequent transcript analysis using semi-quantitative RT-PCR revealed that these mutant lines still had CHIA transcript even to similar extent than the corresponding wild types (Figure 3-17C). Thus, chia-1, chia-2 and chia-3 were not suitable for analysis of the CHIA mutant phenotype.

Figure 3-17: Characterisation of CHIA T-DNA insertion lines

(A) Gene model of CHIA with the T-DNA insertions indicated by grey triangles. The exons, introns and the untranslated regions (3'-UTR, 5'-UTR) are depicted as described in Figure 3-3. (B) The T-DNA insertion lines and the corresponding ecotypes were genotyped using following primer combinations: LP_853931 and RP_853931 (WT-PCR, chia-1), Wisc-Lba and RP_853931 (Lba-PCR, chia-1), LP_N595362 and RP_N595362 (WT-PCR, chia-2), Salk-Lba and RP_N595362 (Lba-PCR, chia-2), At5g24090F1 and At5g24090R1 (WT-PCR, chia-3) and Ds5-1 and At5g24090R1 (Lba-PCR, chia-3). (C) The CHIA transcript analysis was done using following primer combinations: At5g24090F and At5g24090R (chia-1 and chia-2) and At5g24090F and At5g24090RP2 (chia-3).
In a parallel approach, transgenic CHIA knock-down (chia-kd) lines were generated using the artificial microRNA technology (2.4.13), which can be used to effectively silence specific genes in plants (Ossowski et al., 2008). The sequence region used for the generation of the CHIA-specific amiRNA is indicated by a grey triangle in Figure 3-18A. Also CHIA overexpression (CHIA-oe) lines carrying a p35S::CHIA-GFP cassette were created in Col-0 background as described in chapter 2.4.14. Two independent lines of both CHIA knock-down and overexpression plants were further analysed. The genotyping PCRs showed that both CHIA-oe lines contained additionally to the genomic CHIA fragment (gDNA) also the introduced cDNA using primers that amplify the whole gene, and also the 35S promoter was detectable (Figure 3-18B, WT- and 35S-PCR). AmiRNA-specific primers were used to detect the amiRNA-cassette in the chia-kd-1 and chia-kd-2 mutant lines, which was absent in Col-0 plants (Figure 3-18B, amiRNA-PCR). Since the effects of the amiRNA-mediated gene silencing act at a posttranscriptional level, the full-length genomic CHIA gene product can be amplified also in the knock-down lines (Figure 3-18B, WT-PCR). The CHIA transcript levels were monitored via RT-qPCR. Whereas CHIA overexpression lines displayed massively elevated transcription of CHIA (250 to 400-fold transcript levels in comparison to WT; Figure 3-18C, left diagram), the CHIA knock-down lines contained only approximately 5 - 10 % of the WT CHIA transcript amounts (Figure 3-18C, right diagram). Hence, the CHIA amiRNA and overexpression lines displayed the expected genetic properties and could be used for further experiments.
Figure 3-18: Characterisation of CHIA knock-down and overexpression lines

(A) Gene model of CHIA with the 20 bp region targeted for artificial microRNA-mediated gene silencing (grey triangle). Exons and introns are indicated by black boxes and black lines, respectively. The 3’- and 5’-UTR regions are shown as grey boxes. (B) Leaves of two independent plants of the CHIA overexpression lines (CHIA-oe) and chia knock-down lines (chia-kd) were genotyped using two sets of PCR reactions with the primer combinations: At5g24090F and At5g24090R (WT-PCR, CHIA-oe and chia-kd), GC248 and At5g24090R (3SS-PCR, CHIA-oe), A-PRS300 and B-PRS300 (amiRNA-PCR, chia-kd). (C) CHIA transcript accumulation was monitored using RT-qPCR and the primers At5h24090Fq and At5g24090Rq in the CHIA-oe lines (left diagram) and the chia-kd lines (right diagram).

3.4.3.1 Phenotypic analysis of the CHIA overexpression and knock-down lines

Under control cultivation conditions in short day both the CHIA knock-down and overexpression plants exhibited normal phenotypes comparable to Col-0 (Figure 3-19A, upper row from left: Col-0, CHIA-oe-1, CHIA-oe-2, lower row from left: Col-0, chia-kd-1, chia-kd-2). Under low light intensity the leaves of the CHIA overexpression lines were somewhat stunted in comparison to WT leaves (Figure 3-19B, upper row from left: Col-0, CHIA-oe-1, CHIA-oe-2, lower row from left: Col-0, chia-kd-1, chia-kd-2). But as all the bio assays were
always performed in short day and under controlled cultivation conditions the “low-light” aberrance of the CHIA-oe lines was of no consequence.

Figure 3-19: Phenotypes of CHIA overexpression and knock-down lines

CHIA-oe-1, CHIA-oe-2, chia-kd-1, chia-kd-2 and the corresponding ecotype were cultivated in the short day either under normal conditions (A) or under low light intensity (B) and photographed in the age of 5 weeks from the top view.

3.4.4 Subcellular localisation of CHIA

According to the Bio-Array resource analysis tool (BAR) the CHIA protein is localised outside the plasma membrane in the apoplastic space. To visualise the localisation of the GFP-tagged CHIA protein leaves of two transgenic Arabidopsis CHIA overexpression lines were investigated using the confocal laser-scanning microscope. The GFP fluorescence was localised in both CHIA-oe lines in the cell periphery showing a slightly patchy and irregular pattern (Figure 3-20). As negative control leaves of Col-0 plants were monitored using the same excitation setup and laser intensity.
Figure 3-20: Localisation of CHIA-GFP in *Arabidopsis*

GFP fluorescence in the leaf epidermal cells of the two independent transgenic lines stably expressing a 35S::CHIA-GFP construct (CHIA-oe-1, CHIA-oe-2) was analysed using confocal laser-scanning microscopy. Col-0 leaves served as control. Argon/Krypton laser was used to excite the fluorescence at 488 nm, the light emission was monitored between 500 and 600 nm. Left panel (GFP channel), middle panel (light) and right panel (merge of GFP and light).
To gain more insights into the subcellular localisation of CHIA the transient expression system in *Nicotiana benthamiana* was used (see chapter 2.4.11). It allowed, beside the expression of a WT CHIA-GFP construct, also the expression and rapid analysis of a mutated version of CHIA lacking the putative secretion signal peptide (CHIAΔSP-GFP). The WT CHIA-GFP displayed a similar patchy localisation pattern in tobacco as in *Arabidopsis* (compare Figure 3-20 and Figure 3-21, CHIA-GFP). The plasma membranes of two adjacent plant cells appeared as one thick layer as stained by FM4-64 and the GFP fluorescence was detected in distinct areas around this layer as shown in the close-up picture (Figure 3-21, CHIA-GFP, lower panel). The deletion of the signal peptide led to a dramatic change in the localisation of the fluorescence signal. In leaves expressing CHIAΔSP-GFP the GFP fluorescence was present in the cytosol, showing an evenly spread signal completely different to the patchy localisation pattern of the WT CHIA (Figure 3-21, compare close-ups for CHIA-GFP and CHIAΔSP-GFP). Unfortunately, the membrane stain FM4-64 was no longer detectable at the plasma membrane but was weakly visible in distinct cytosolic structures. FM4-64 is rapidly taken up by endocytosis into the cytoplasm and thus does not stably stain the plasma membrane (Bolte et al., 2004; Ueda et al., 2001). GFP alone was expressed as a positive control for cytoplasmic localisation and mock-infected tobacco leaves served as negative control (Figure 3-21, GFP and negative control). Approaches to induce plasmolysis in leaf epidermal cells and to observe whether fluorescence signal is truly between the cells turned out to be technically challenging. The plasmolysis did not take place evenly in many surrounding cells, so that the possible fluorescence signal outside of an affected cell could not be distinguished from the normal signal of a neighboring unaffected cell (data not shown). The direct microscopical evidence for apoplastic localisation of CHIA is not trivial, also because the low apoplastic pH additionally quenches the fluorescence signal (Scott et al., 1999). Nonetheless, the microscopical localisation studies suggest that the CHIA protein is indeed secreted into the plant apoplast and that the signal peptide is essential for this targeted localisation. Moreover, this prediction is supported by a proteomic approach using *Arabidopsis thaliana* cell cultures that revealed the CHIA chitinase among the secreted, apoplastic proteins (Kwon et al., 2005).
The 35S::CHIA-GFP and 35S::CHIAΔSP-GFP constructs were transiently expressed in Nicotiana benthamiana leaves using Agrobacterium tumefaciens-mediated transformation. GFP fluorescence in the leaf epidermal cells was analysed 3 days post infection. As positive GFP control leaves were transfected with the pK7WGF2.0 vector. As negative control uninfected leaves were used. FM4-64 was used to stain the plasma membrane. Argon/krypton laser was used for excitation of GFP at 488 nm and the 543 nm line of helium/neon laser for the excitation of FM4-64. Detection wavelengths of emitted light were 500 nm to 600 nm (GFP) and 560 nm to 615 nm (FM4-64). Panels from left: GFP, FM4-64, merge of GFP and FM4-64 and light.

Figure 3-21: Localisation of CHIA-GFP and CHIAΔSP-GFP in tobacco
3.4.5 Detection of CHIA protein in the transgenic CHIA lines

To monitor the CHIA protein levels in the generated transgenic CHIA lines two antibodies were available. First, antibodies raised against GFP could be used to detect the GFP-tagged CHIA in the CHIA-oe plants. Second, an antibody raised against the tobacco class III chitinases (α-class III chitinase antibody from rabbit, provided by Frédéric Brunner) could be tested for recognition of the Arabidopsis class III chitinase homolog CHIA. The α-GFP antibody detected specific bands in leaf extracts of CHIA-oe lines in the sizes of approximately 60 kDa and 30 kDa, which correlated with the expected sizes of CHIA-GFP (60.1 kDa) and GFP alone (27 kDa) (Figure 3-22A). The detection of CHIA using the α-class III chitinase revealed a more complex pattern of protein bands. In the CHIA overexpression plant extracts a clear band was visible below the 35 kDa marker. This protein band possibly represents the CHIA protein, which has the calculated size of 33.1 kDa, without the GFP tag (Figure 3-22B). In the WT lane no protein band was visible between 25 and 35 kDa (Figure 3-22B). Weak bands in the size of the CHIA-GFP fusion protein were also present in the overexpressor extracts (Figure 3-22B). It is possible that the large GFP tag is disturbing the recognition of the CHIA protein by the α-class III chitinase antibody leading to weaker detection of the tagged protein or indeed most of the protein is cleaved. The additional bands visible on the immunoblot might present cross-reacting bands. The presence of the secreted form of GFP protein in the transgenic p35S::secGFP line was also confirmed by immunoblot analysis (Teh and Moore, 2007). SecGFP could be detected using α-GFP antibody in the expected size of approximately 30 kDa (Figure 3-22C). The exact size of secGFP is due to an additional c-myc tag 31 kDa (Teh and Moore, 2007). In the same immunoblot analysis also the absence of CHIA-GFP in the chia-kd line was investigated (Figure 3-22C).

In order to purify and enrich the CHIA-GFP fusion protein from the CHIA-oe total leaf extracts an immunoprecipitation approach was exploited. The purified CHIA-GFP sample could then be used for further experiments, such as activity assays. Agarose A beads coupled with the antibody against GFP were used to pull down the GFP-tagged CHIA protein. The immunoprecipitation functioned well and CHIA-GFP could be detected at the expected size of approximately 60 kDa as shown in Figure 3-22D. Interestingly, upon immunoprecipitation specific high-molecular weight bands appeared in both CHIA-oe samples, but were not present in present in WT samples possibly representing CHIA-GFP multimers (Figure 3-22D, asterisk). Also here degradation of CHIA-GFP was observed since GFP was visible as a separate band at about 30 kDa.
Results

Figure 3-22: Detection of CHIA protein in Arabidopsis

Immunoblot analysis of protein extracts from leaves of transgenic CHIA lines and Col-0 plants. Total leaf protein was separated by SDS-PAGE and blotted onto a nitrocellulose membrane. The immunodetection was carried out using α-GFP rabbit (A), α-class III chitinase rabbit (B) or α-GFP goat antibody (C). The α-mouse-HRP and α-goat-HRP were used as secondary antibodies and the protein bands were visualized by chemiluminescence. Ponceau staining of the large subunit of RuBisCO served as loading control. Transgenic plants expressing a p35S:secGFP construct was used as control for GFP. (D) Total leaf protein of CHIA overexpression lines was subjected to immunoprecipitation using α-GFP goat antibody and agarose A beads. The immunodetection was performed with rabbit α-GFP and α-rabbit-HRP antibodies. IN, protein input; IP immunoprecipitated sample; * indicates a specific high-molecular weight band present in IP samples; ** indicates the band of the heavy chain of the α-GFP goat antibody.

3.4.6 Posttranslational modification of the CHIA protein

Posttranslational modifications of plant proteins include glycosylation (Tekoah, 2004). As in mammals both N- and O- linked glycosylation takes place in plants, although the mechanisms vary. The attachment of N-linked glycans to proteins begins in eukaryotes in the
endoplasmic reticulum (ER), where glycosylation sites, tripeptides N-X-S/T (X can be any amino acid except proline and aspartic acid) are recognised (Lerouge et al., 1998). The CHIA protein sequence contains two putative N-glycosylation sites (N44 and N113). As many secretory proteins undergo this type of posttranslational modification, it is possible that also the CHIA chitinase is chemically altered in this way. Depending on the amount and size of attached N-glycans the modification leads to an increased size of the glycosylated protein. This glycosylation-dependent size shift can be exploited in the analysis of posttranslational modification. Thus, total leaf protein extracts of either Arabidopsis CHIA-oe plants or tobacco transiently expressing p35S::CHIA-GFP or p35S::CHIAΔSP-GFP, were subjected to deglycosylation and afterwards the possible reduction in the size of the CHIA-GFP fusion protein due to the action of the glucosidases was detected by immunoblot analysis (Figure 3-23). For both Arabidopsis CHIA-GFP and in Nicotiana expressed CHIA-GFP a small size shift was observable in the deglycosylated sample in comparison to the untreated protein sample. In contrast, the CHIAΔSP-GFP fusion protein lacking the secretion signal displayed no reduction in the protein size after deglycosylation treatment (Figure 3-23), probably because of missing glycosylation due to misregulated targeting.

The deglycosylation experiments revealed that the CHIA-GFP fusion protein was glycosylated both in Arabidopsis thaliana and Nicotiana benthamiana and that this modification was dependent on the presence of the secretion signal.

![Figure 3-23: Deglycosylation of CHIA-GFP and CHIAΔSP-GFP](image)

Protein extracts from leaves of transgenic Arabidopsis CHIA-oe-1 plants (A) and Nicotiana expressing either p35S::CHIA-GFP or p35S::CHIAΔSP-GFP were subjected to deglycosylation with a mixture of deglycosylases. The negative control (-) was treated as the deglycosylation sample (+) but without addition of the deglycosylation enzyme mix. The immunoblot analysis was carried out as described in Figure 3-22.

3.4.7 Expression of CHIA protein using heterologous expression systems
To analyse the enzymatic properties of the CHIA chitinase in vitro it was expressed heterologously in E.coli. The expression of the His₆-CHIA fusion protein in BL21AI cells was induced by L-arabinose for 3 hours at 18 °C and after lysis of the cells the soluble protein fraction was subjected to affinity-purification using a Ni²⁺-NTA column. The His₆-CHIA could be purified to high extent and most protein was eluted from the column in the first two elution
steps as visualised by Coomassie stain and immunoblot analysis (Figure 3-24A, B; lanes 3 and 4). The majority of the protein was visible at the expected size of approximately 35 kDa, but also an additional double band was running at the size of about 65 kDa (Figure 3-24B). Afterwards, the PGN-hydrolytic activity of the fusion protein was assessed in a turbidity assay using Micrococcus luteus cells as substrate. The cell walls of the Gram-positive bacteria Micrococcus luteus are widely used as crude peptidoglycan substrate for enzymatic assays (Biswas et al., 2006; Brunner et al., 1998; de Azambuja et al., 1991; Park et al., 2002). The reduction in the turbidity of the suspension due to PGN-hydrolysis is thereby monitored and the absolute reduction of the OD or the relative activity (comparison to control PGN-hydrolysing enzymes like lysozyme) can be measured. The purified His6-CHIA protein or the corresponding uninduced sample (negative control) were incubated together with the M.luteus substrate and the hydrolytic activity was monitored after 2 and 20 hours (Figure 3-24C). After 2 hours of incubation the relative activity of the His6-CHIA protein was approximately 10 % of the lysozyme activity and 20 hours after incubation a relative activity of 40 % was measured. Unluckily, the negative control showed similar activities than the His6-CHIA sample, indicating that the background activity resulting from bacterial enzymes was very high. It is also possible that traces of lysozyme, used for the lysis of the BL21AI cells, were present in the samples despite the affinity purification. All in all, the measured enzyme activity after 2 hours of incubation was very weak implicating that the prokaryotic E.coli system might be suboptimal for the expression of CHIA, which seems to be posttranslationally modified (Figure 3-23).
His6-tagged CHIA expression in the *E. coli* strain BL21AI was induced by L-arabinose and the bacteria were harvested after incubation for 3 hours at 18 °C. The soluble His6-CHIA protein was purified using Ni²⁺-NTA affinity column. The protein amounts in the input (lane 1), flow-through fraction (lane 2) and 5 elution fractions (lanes 3-7) were analysed by Coomassie blue stain (A) or immunoblot using the α-His6 mouse and α-mouse-HRP antibodies (B). Arrows indicate purified proteins and the asterisk putative CHIA oligomers. (C) The PGN-hydrolysis activity of the His6-CHIA protein was monitored in a turbidity assay. 20 µg of the purified His6-CHIA protein was incubated with *Micrococcus luteus* cells and the reduction in turbidity (OD₅₇₀nm) was monitored over time. The relative activity was calculated using the hen egg-white lysozyme as standard (lysozyme activity was set as 100 %). As negative control the bacterial protein extract of the uninduced sample and elution buffer were used.

Fungal expression systems can be used for expressing active plant enzymes (Park et al., 2002; Petersen et al., 2009). Therefore, for the second approach of expressing active CHIA protein the methylotrophic yeast *Pichia pastoris* was chosen. Two different vectors enabling secretion of the expressed protein in the culture medium were used (pPICZαC and pPIC9K, Invitrogen). The protein expression was induced by methanol and monitored after 1 to 4 days of cultivation of the yeast cells at 30 °C (Figure 3-25A, B). However, the expression of the CHIA protein was not induced in any of the tested clones. The protein patterns in the cells transformed with the CHIA constructs resembled the patterns in the empty vector (ev) controls.
The approaches to heterologously express an active CHIA chitinase failed so far. Nonetheless, especially the eukaryotic *Pichia* system has still potential and new expression constructs and the usage of other tags could solve the induction problems.

Figure 3-25: Expression of CHIA in *Pichia pastoris*

Expression of His_{6}-tagged or untagged CHIA in the yeast *Pichia pastoris* was induced by methanol. The cells were harvested after 1 to 4 days incubation at 30 °C. The secreted protein in the medium was precipitated by TCA and analysed by Coomassie blue stain. (A) Expression of CHIA-His_{6} and untagged CHIA using the pPICZαC expression vector. (B) Expression of untagged CHIA using the pPIC9K vector. The empty vector sample is indicated by ev.

3.4.8 Analysis of homo-oligomerisation properties of CHIA

Since some CHIA-specific high-molecular weight protein bands were detected both in *Arabidopsis* (Figure 3-22D) and *E.coli* (Figure 3-24A, B) protein extracts, the homo-oligomerisation properties of CHIA were analysed more in detail in a yeast two-hybrid (Y2H) assay (Figure 3-26). The Y2H system is based on the ability of two hybrid proteins to interact with each other inducing thereby the induction of GAL4-regulated transcription of reporter genes. One of the hybrid proteins contains the DNA binding domain (DB) of the GAL4 transcription factor and the other the GAL4 activation domain (AD). The two constructs are combined with different selection markers (BD: tryptophan, W; AD: leucine, L). An interaction between the CHIA-BD and CHIA-AD due to homo-oligomer formation would then lead to the
reconstitution of the GAL4 transcription factor and induction of the transcription of reporter genes. The gene product of the reporter gene *Ade2* encodes an enzyme enabling the biosynthesis of adenine and the usage of specific media lacking adenine can be used to assay the interaction of the constructs. First, the transformation of the yeast cells was monitored using a drop test and medium lacking the amino acids tryptophan and leucine (Figure 3-26, -LT). Here all the tested combinations showed growth. The interaction medium contained no tryptophan, leucine and adenine (Figure 3-26, -LTA) and allowed growth of yeast cells only in the positive control. The positive control was the interaction between the SV40 large T-antigen and the murine p53 (pGAD-T + PGBK-53). The protein expression in the different combinations was verified using immunoblot analysis (data not shown). Thus, an ability of CHIA to form homo-oligomers was not detectable in the yeast two-hybrid assay. However, since Y2H assay is an artificial system further studies *in planta* are necessary.

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3.4.9 Assessment of the chitin- and PGN-hydrolase activity of CHIA

To learn more about the function of the *Arabidopsis* CHIA chitinase its enzymatic properties in hydrolysis of chitin and peptidoglycan were assessed (chapters 2.5.11, 2.5.12 and 2.5.13). First, the chitinolytic activity of leaf extracts from transgenic CHIA lines and Col-0 plants towards the chitin derivate 4-methylumbelliferyl-β-D-N, N’, N” triacetylchitotriose (4-MUCT)
was measured after 4 hours of incubation (Figure 3-27A). The relative enzymatic activity was calculated using the activity of *Streptomyces griseus* chitinase as standard (*S. griseus* activity was set as 100 %). The transgenic secGFP line expressing secreted GFP was used as a control for external GFP. The background activity observed in the buffer control was approximately 5 % and the relative activities measured in the WT, chia-kd and secGFP samples were roughly 10 % of the *S. griseus* chitinase activity. The most active sample hydrolysing 4-MUCT was the CHIA-oie leaf protein extract, which showed a relative activity of almost 250 %. The strong activity of the CHIA-oie sample was already visible after 2 hours (data not shown). As another chitin substrate, colloidal chitin was used in a hydrolysis assay. In this assay the Col-0 leaf protein sample displayed approximately 27 % relative activity, whereas the two tested CHIA-oie samples showed a relative activity between 38 - 40 % (Figure 3-27B). However, these increased relative activities in the CHIA-oie samples were not significantly different from the Col-0 sample. Possibly the general chitinolytic activity against colloidal chitin resulting from other *Arabidopsis* chitinases present in the crude leaf extract add up to the resulting enzymatic activity. In contrast, 4-MUCT, might represent a specific substrate for class III chitinases. To explore the peptidoglycan hydrolysis capacity of the transgenic CHIA lines complex Lys-type peptidoglycan from intact Gram-positive *M.luteus* cells were used as substrate in a turbidity assay (Figure 3-27C). Here, similar activity patterns were observed as for the 4-MUCT substrate; Col-0, chia-kd and secGFP leaf samples showed similar activity levels (around 20 % of lysozyme activity which was set to 100%), whereas the CHIA-oie sample displayed the clearly strongest hydrolytic activity (120 %). In parallel, also the immunoprecipitated CHIA-GFP samples (see chapter 3.4.5 for details) were subjected to turbidity assay with *M.luteus* cells as substrate, but no activity was measured (data not shown). The lack of enzymatic activity in the IP samples might be due to sterical hindrance by the beads or the absence of essential co-factors, which were not immunoprecipitated. Ultimately, also purified insoluble Dap-type PGN from *Bacillus subtilis* was subjected to hydrolysis by the total leaf protein samples (Figure 3-27). Once again the leaf sample of the transgenic line overexpressing CHIA was most active (40 %) and the other tested leaf samples showed slightly higher activity than the buffer control (buffer: 3 %; Col-0, chia-kd and secGFP: 8 %).
Figure 3-27: Chitin- and PGN-hydrolysis activity of CHIA leaf protein

Protein extracts from leaves of adult homozygous lines (CHIA-oE, chia-kd) were assayed for chitinolytic activity with 4-MUCT substrate (A) and colloidal chitin (B) or for PGN-hydrolytic activity against M. luteus cells in a turbidity assay (C). Relative activities (4 hrs post treatment) were calculated using Streptomyces griseus chitinase (A, B) or hen egg-white lysozyme (C) as standards. Plants expressing secreted GFP (secGFP) served to control the effect of external GFP. (D) Bs PGN was subjected to hydrolysis by leaf extracts for 2 hours, and PGN solubilization was calculated as in (C). 60 µg total leaf protein was used per sample. Means and standard errors of two replicates per sample are given. Significant differences in enzyme activities relative to those in Col-0 are indicated (asterisks, p ≤ 0.05 for A and C and p ≤ 0.09 for D; Student’s t-test).

The activity assays performed with the crude leaf protein extracts give no additional information about the localisation of the enzyme activity. To address the question, whether CHIA is really secreted and acts as an enzyme in the apoplast an approach using protoplasts was exploited. Protoplasts were isolated from WT and transgenic plants and incubated...
overnight in the dark. The protoplasts were then gently separated from the medium and the secreted proteins in the medium were concentrated before usage in hydrolytic activity assays. In the 4-MUCT assay the majority of the chitinolytic activity was present in the secreted (supernatant) fraction of the CHIA-oe protoplast sample (Figure 3-28, 70 %) but a lot of activity was also present in the CHIA-oe protoplast pellet (40 %) indicating possible overexpression-related artefacts. The relative activity measured in the other samples was either very low (chia-kd and secGFP samples, 5%) or not detected at all (control samples and Col-0 samples). The PGN hydrolysis assay using M. luteus cells as substrate revealed that the secreted protein present in the CHIA-oe protoplast sample had the strongest PGN-hydrolysing properties (CHIA-oe-S, 50 %) compared to lysozyme activity. The CHIA-oe protoplast pellet displayed together with the Col-0 and secGFP samples relative activity levels between 10 - 15 %. Notably, the secreted protein fraction of the chia-kd protoplast sample was significantly less active than the corresponding WT sample. The presence of CHIA protein in the protoplast samples was confirmed by immunoblot analysis (Figure 3-28C). CHIA protein was detectable both in the CHIA-oe supernatant and protoplast sample. Interesting was the finding, that whereas in the pellet sample only CHIA-GFP could be visualised using the α-GFP antibody, in the supernatant sample exclusively processed CHIA protein lacking the GFP tag was detected with the α-class III chitinase antibody. A possible explanation for the differences in the protein pattern in the supernatant and pellet samples could be of biological but also technical nature. It is conceivable that the recombinant CHIA-GFP is proteolytically processed during the secretion pathway or that the additional enrichment step of the secreted protein samples led to enhanced protein degradation (2.5.4).

In conclusion, CHIA overexpression plants can hydrolyse both peptidoglycan and substrates derived from chitin indicating that the Arabidopsis CHIA protein represents an enzyme harboring both chitinolytic and PGN-hydrolysis activity. Furthermore, CHIA was able to degrade both Lys-type and Dap-type peptidoglycan substrates, presented by M. luteus and B. subtilis respectively, suggesting similarly broad PGN-specificity than the mammalian peptidoglycanolytic PGLYRP2 (Gelius et al., 2003). Detailed analysis using protoplasts showed also that the majority of the CHIA activity was indeed present in the secreted protein fraction of CHIA-oe protoplasts. The CHIA protein levels in uninduced wild type leaves are probably too low to be able to result in differences between WT and chia-kd protein extracts. Removal of the plant cell wall during protoplast preparation might lead to stress-induced induction of CHIA in the WT but not chia-kd sample. Hence, differences were visible only using protoplast samples (Figure 3-27), but not with leaf extracts (Figure 3-28). The inducibility of CHIA-derived activity in wild type plants and also the possibility of using
immunoprecipitated CHIA-GFP samples for the activity assays have to be further investigated.

Figure 3-28: Chitin- and PGN-hydrolysis activity of CHIA protoplast samples

Protoplasts of transgenic lines were pelleted, and protein extracts of the pellet (P) or supernatant (S) was subjected to either hydrolysis of 4-MUCT (A) or *M. luteus* cells (B). The relative activities were calculated 3 hours post treatment as described in Figure 3-27. 15 µg of total protein was used per sample. Means and standard errors of two replicates per sample are given. Significant differences in enzyme activities relative to those in Col-0 are indicated (asterisks, p ≤ 0.05; Student’s t-test). Samples with no measurable activity are indicated accordingly (not detected, n.d.). (C) CHIA protein levels in the supernatant and pellet samples were analysed by immunoblotting. The immunoblot analysis using α-GFP rabbit, α-class III chitinase rabbit and α-rabbit-HRP antibodies was carried out as described in Figure 3-22.
3.4.10 Role of CHIA in fungal resistance
The CHIA protein belongs to the family of chitinases, which are generally considered as pathogenesis-related (PR) proteins. Members of this family have been shown to degrade fungal cell walls and inhibit fungal growth in vitro (Arlorio et al., 1992; Mauch et al., 1988; Schlumbaum et al., 1986). Since CHIA also exhibited chitinolytic activity its influence in defending the plant from fungal infections was analysed.

3.4.10.1 Infection of the transgenic CHIA lines with Botrytis cinerea
Leaves of transgenic CHIA lines and WT plants were infected by drop-inoculation with the necrotrophic fungus *Botrytis cinerea* and kept under high humidity to provide optimal infection conditions for the pathogen. The disease symptoms were monitored two and three days post infection. A Trypan blue stain was used to visualise both fungal hyphae and dead plant cells (Figure 3-29A, B). The infection site after 2 days post infection appeared similar in all tested lines as shown in Figure 3-29A. Also the analysis of disease symptoms at a cellular level revealed no differences between the analysed lines (Figure 3-29B). The hyphal outgrowth and the cell death zones observed in the transgenic lines resembled the ones in WT plants. In addition to the infection symptoms also the size of the lesions was measured three days post infection (see Figure 3-29C). However, no significant differences were seen in the lesion sizes between transgenic CHIA lines and Col-0 plants. In total, the infection of the aggressive fungus *B. cinerea* proceeded in the CHIA overexpression and knock-down lines similarly as in the WT plants.
Figure 3-29: Infection of the transgenic CHIA lines with *Botrytis cinerea*

Five week-old plants were infected with the necrotrophic fungus *Botrytis cinerea*. 5 µl spore suspension of $5 \times 10^5$ spores/ml was drop-inoculated on the one half of the leaf, two leaves per plant were infected. The plants were analysed for symptom development after 2 and 3 days post infection. (A) Trypan blue stain showing visible symptoms after 2 dpi. (B) Microscopic analysis of the infection site and fungal hyphae 2 dpi visualised by Trypan blue stain. (C) Measurement of the lesion size 3 days post infection. Shown are means and standard errors (n=16).

3.4.10.2 Infection of the transgenic CHIA lines with *Alternaria brassicicola*

*Botrytis cinerea* is an aggressive pathogen leading to massive maceration and cell death in the host tissue already at very early stages in the infection (Glazebrook, 2005). Thus, a weaker fungal pathogen causing disease symptoms after a longer time period might evoke differences in the progression of infection in the CHIA-oe or chia-kd lines when compared to WT plants. Therefore, fungal infection assays with the weaker necrotroph, *Alternaria*
brassicicola, were performed. Like Botrytis, Alternaria spores were also drop-inoculated on the leaf surface of the different lines and the infected plants were cultivated under high humidity. After 14 days post infection the leaves showed chlorosis and also necrosis spreading from the inoculation sites (Figure 3-30A, B). All the tested lines showed these disease symptoms, although the necrotic lesions displayed by the two CHIA overexpression lines appeared smaller and the lesions in the knock-down lines more drastic than in the WT plants. The microscopical analysis of the infection sites could not reveal major differences in the fungal hyphal growth in the tested lines as visualised by Trypan blue stain (Figure 3-30C). Additionally, also the course of the infection was monitored over a 14-day period and the degree of symptoms was determined as described in 2.6.2 (Figure 3-30D). After 7 days of infection the disease index was roughly 200 for all included lines. After 11 and 14 days the infection was more advanced and a slight tendency was visible; the CHIA-oe lines showed less and the chia-kd lines slightly more symptoms than the corresponding WT plants. However, these subtle differences could not be verified as being significant.

The results of the fungal infections cannot completely rule out the possibility that the CHIA chitinase is involved in fungal resistance. It is possible that it has a supportive role acting together with other chitinases or defense-related enzymes, like β-1,3-glucanases. In that case the generation of multiple KO mutants is most likely necessary to be able to get clear phenotypes in fungal defense.
Figure 3-30: Infection of the transgenic CHIA lines with *Alternaria brassicicola*

Five week-old plants were infected with the necrotrophic fungus *Alternaria brassicicola*. Six 5 µl droplets of spore suspension of 5 x 10⁵ spores/ml were inoculated on the leaf, two leaves per plant were infected. The plants were analysed for symptom development after 7, 11 and 14 days post infection. (A) Visible symptoms of four independent leaves after 14 dpi. (B) Disease symptoms after 14 dpi visualised by Trypan blue stain. (C) Microscopic analysis of the infection site and fungal hyphae 14 dpi visualised by Trypan blue stain. (D) Calculation of the disease index 7, 11 and 14 days post infection. Shown are means and standard errors (n=16).
3.4.11 Impact of CHIA on bacterial resistance

As CHIA also had PGN-lytic activity, its importance in defending the plant against bacterial pathogens was assessed in bacterial infection assays (2.6.1). *Pseudomonas syringae* strains with different virulence properties were tested. First, leaves of transgenic CHIA lines and Col-0 plants were infiltrated with virulent *Pto* DC3000 bacteria and the growth of the bacteria was measured 4 days post infection (Figure 3-31A). Importantly, directly after the infiltration similar amounts of bacteria were counted in the leaves of the different lines enabling the comparison between the lines at later time points (0 days post infection). During the 4 days of infection the bacterial cells had propagated strongly, in the wild type leaves $10^7$ colony forming units per cm$^2$ leaf tissue were measured (Figure 3-31A, Col-0). The two independent CHIA knock-down lines were significantly more susceptible to the virulent *Pseudomonas* strain than Col-0 (Figure 3-31A, *chia-kd-1* and *chia-kd-2*) indicating that CHIA is essential for the bacterial resistance. Unexpectedly, also the CHIA-oe plants allowed *Pto* DC3000 to grow better (nearly $10^9$ cfu/cm$^2$ leaf tissue) than WT plants. The susceptibility phenotype of the CHIA-oe lines was already visible 2 days post infection (data not shown). Next, the hypovirulent *Pto* DC3000 strain lacking avrPto and PtoB ($\Delta$avrPto/PtoB) was tested. The growth of *Pto* DC3000 $\Delta$avrPto/PtoB was monitored 2 days after infiltration; both the two *chia-kd* lines and the two CHIA-oe lines showed significantly enhanced bacterial growth rates compared to WT plants (Figure 3-31B). 4 days post infection the susceptibility phenotype of the transgenic CHIA lines was no longer detectable (data not shown). The infection assays with the type III secretion system-deficient strain *Pto* DC3000 hrcC$^{-}$ could so far deliver no significant differences between the transgenic CHIA lines and wild type plants (data not shown). Additionally, also the infection of the CHIA lines with the non-host strain *Pseudomonas syringae* pv. *phaseolicola* should be investigated.

The absence of CHIA led to a susceptibility phenotype upon infection with both the hypervirulent phytopathogenic bacteria *Pto* DC3000 and the less virulent mutant strain *Pto* DC3000 $\Delta$avrPto/PtoB. These data suggest that lack of PGN-degrading activity dampens plant immunity. The results showing more bacterial growth also in the CHIA overexpressing plants was unexpected. It seems that changes in the PGN-lytic activity provided by CHIA no matter in what direction lead to distorted bacterial resistance behavior. Follow-up experiments are needed to clarify the role of CHIA and its PGN-hydrolase activity in the protection against bacterial attack.
Figure 3-31: Infection of the transgenic CHIA lines with *Pseudomonas syringae*

Transgenic CHIA plants are hypersusceptible to bacterial infection. Growth of *Pto* DC3000 (A) and *Pto* DC3000 ΔavrPto/PtoB (B) was determined 2 or 4 days post infiltration of $10^5$ colony forming units ml$^{-1}$ (cfu/ml). Data represent means ± SD of six replicate measurements/genotype/data point. Statistical significance compared to wild-type ($p \leq 0.05$, Student's t-test) is indicated by asterisks. Representative data of at least four independent experiments are shown.
4 Discussion

Although preliminary knowledge has been gathered in the past few years regarding bacterial peptidoglycans as bioactive stimuli of the plant immune response (Erbs et al., 2008; Gust et al., 2007), the molecular details of the perception mechanisms were still unresolved. In animals, the PGN sensing machinery is widely understood and distinct pathways encompassing a large variety of membrane-tethered and cytosolic receptors, PGN recognition proteins and PGN hydrolytic enzymes have been unraveled in the past decades (Royet and Dziarski, 2007). Several plant LysM-domain containing receptor-like kinases (LysM-RLKs or LYKs) are implicated in the recognition of glycan moieties, either during pathogenic host-fungi or symbiotic host-rhizobacteria interactions (Limpens et al., 2003; Miya et al., 2007; Radutoiu et al., 2003). Hence, a targeted reverse genetics approach was undertaken to search for peptidoglycan receptor(s) among the *Arabidopsis* family of LysM-RLKs.

Complex polymeric carbohydrates derived from pathogenic cell walls, such as fungal chitin and oomycete β-glucan, are preferably bound by corresponding plant high-affinity binding proteins in specific oligomeric fragment lengths (Fliegmann et al., 2004; Ito et al., 1997). Also polymeric peptidoglycan is processed by metazoan enzymes into smaller, yet still immunostimulatory fragments, which can then activate the corresponding signaling pathways (Chaput and Boneca, 2007; Wang et al., 2006; Zaidman-Rémy et al., 2006). As it could be shown that *Arabidopsis* is able to perceive not only complex but also fragmented peptidoglycan (Figure 3-12), it was also assayed whether PGN processing enzymes, which show similar activity than the animal PGN hydrolases, are present in *Arabidopsis thaliana* and contribute to bacterial immunity.

4.1 LYKs contribute to plant innate immunity

4.1.1 Effects of LYK gene deletions in plant fungal and bacterial resistance

Many proteins are controlled already at the level of gene transcription allowing stimulus-dependent increase or reduction of the gene products (Singh, 1998). Hence, changes in the gene expression pattern upon different developmental or environmental cues can give indications for possible protein functions. Interestingly, the expression of all investigated LYK genes showed a clear suppression either 6 or 24 hours after infection with the virulent *Pseudomonas syringae pv. tomato* DC3000 (Figure 3-2). Among the genes encoding *Arabidopsis* LysM proteins, also *LYM3* was similarly suppressed upon *Pto* DC3000 treatment (Willmann, 2011). Such infection-mediated transcriptional reduction has been shown to result
from the effector activity of plant pathogens (He et al., 2006). In contrast, the application of hypovirulent \((Pto\ hrcC^-)\) or non-host \((Pph)\) strains led to an upregulation of the expression of \(CERK1\), \(LYK4\) and \(LYK5\) genes (Figure 3-2). The expression profile of the \(LYK3\) gene differed most from the other \(LYKs\) upon bacterial infection displaying mainly different degrees of repression (Figure 3-2), however whether this \(P.syringae\)-dependent repression of \(LYK3\) also has a biological role remains to be clarified.

\(LYK2\) gene was omitted from the analysis since the extremely low expression levels in leaf tissue make it the poorest receptor candidate during pathogen attack. In addition, the prediction of protein domains for \(LYK2\) gave no clear results regarding a LysM domain, however this might be due to some sequence variability within the lysin motif (Figure 3-2).

The expression profile analysis of the \(LYK\) gene family members is suggestive of a possible participation of \(CERK1\), \(LYK3\), \(LYK4\) and/or \(LYK5\) in bacterial resistance. \(CERK1\) was the first \(Arabidopsis\) \(LYK\) to be ascribed in pathogen resistance. Transgenic plants containing T-DNA insertions within the \(CERK1\) gene displayed severe reduced chitin-mediated defense responses and resulted in enhanced susceptibility against \(Alternaria brassicicola\) and \(Erysiphe cichoracearum\) (Miya et al., 2007; Wan et al., 2008). However, neither the tested \(cerk1-2\) mutant nor two other \(lyk\) mutants, \(lyk3\) and \(lyk5\), showed more disease symptoms against the necrotrophic fungus \(Botrytis cinerea\) than wild type plants (Figure 3-6). This result suggests that either the loss of plant resistance to fungal pathogens is species-specific or that the degree of susceptibility is mild, hence the determination of disease symptoms requires more sophisticated detection methods. For instance, the amount of fungal growth in the infected leaves could be measured using quantitative RT-PCR to be able to discriminate between subtle differences in the WT and mutant plants (Gachon and Saindrenan, 2004). Additionally, the deletion of several \(LYK\) genes might result in a stronger fungal phenotype than observed for the single \(cerk1\) mutant (Miya et al., 2007; Wan et al., 2008). Therefore, it would be intriguing to analyse the generated triple mutant \(cerk1\ lyk3\ lyk5\) (Figure 3-11).

The impact of \(LYK\) mutations in the resistance against phytopathogenic bacteria was assessed using the Gram-negative hemibiotroph \(Pseudomonas syringae\). Besides the hypervirulent WT strain \(Pto\ DC3000\), also strains with diminished virulence due to deletion of certain effectors or the essential type III secretion system (no secretion of effectors at all) can be used to monitor the importance of host genes within bacterial immunity. Gimenez-Ibanez et al. (2009) reported that \(CERK1\) is additionally to its role in chitin perception a determinant of bacterial resistance. The \(CERK1\) gene depletion led to loss of bacterial growth restriction, thus allowing the TTSS-mutant strain \(Pto\ hrcC^-\) but also the virulent \(Pto\ DC3000\) to propagate in an enhanced manner in the two independent mutant alleles in comparison to
WT plants. This susceptibility phenotype for CERK1 depletion could also be observed in the infection assays performed in the frame of this work. Increased bacterial propagation was visible in the cerk1-2 mutant post infection with Pto DC3000, Pto ΔavrPto/PtoB and Pto hrcC' (Figure 3-7). Interesting was the result that also lyk3 mutants displayed a susceptibility phenotype towards the virulent Pseudomonas syringae strain, Pto DC3000, indicating that LYK3 might, together with CERK1, contribute to plant immunity. The lyk5 single mutants, but surprisingly also the lyk3 lyk5 double mutants behaved similarly to wild type plants in bacterial infection assays (Figure 3-7). Possibly, the lyk3 susceptibility phenotype is masked by the Landsberg erecta ecotype portion present in the double mutants. The bacterial susceptibility phenotype mediated by the cerk1 and lyk3 gene depletions and the repression of CERK1 and LYK3 expression upon the hypervirulent Pto DC3000 treatment together with the finding that the Pseudomonas effector protein AvrPtoB specifically targets CERK1 for degradation to promote bacterial virulence (Gimenez-Ibanez et al., 2009) deliver strong evidence for essential participation of CERK1, but maybe also of LYK3, in the formation and maintenance of bacterial immunity.

4.1.2 CERK1 serves together with LYM3 peptidoglycan recognition

Bacteria harbor a whole battery of known but also so far uncharacterised PAMPs (Nürnberger et al., 2004; Zipfel, 2009). To find out whether the recognition of the cell wall-derived PAMP peptidoglycan is connected to the discovered cerk1- or lyk3-dependent limitation of bacterial resistance, PGN-induced expression of defense-related genes was analysed in the cerk1 and lyk3 mutants. Interestingly, the depletion of the CERK1 gene resulted in a similarly diminished response upon treatment of leaves or seedlings with Gram-negative PGN as seen for the lym3 mutant (Figure 3-8 and Willmann (2011)). LYM3 was shown to bind both Gram-negative and Gram-positive peptidoglycan in a reversible and ligand-specific manner and transgenic Arabidopsis plants lacking the functional LYM3 gene were not only PGN-insensitive but allowed also more growth of Pseudomonas syringae (Willmann, 2011). The assessment of the effects of PGN treatment on the global gene expression revealed a dramatic PGN-insensitivity in the cerk1 mutant when compared to WT plants (Figure 3-10). Also mutants lacking LYM3 were not responsive to PGN treatment and showed a massive deregulation of the global gene expression (Willmann, 2011). The reduction of PGN-mediated defense gene expression both in the cerk1 and lym3 mutants was not only measured for Dap-type but also for Lys-type PGN derived from Staphylococcus aureus (Willmann, 2011), suggesting that the identified LysM receptor proteins are part of a peptidoglycan sensing system, which is responsible for the perception of different peptidoglycan subtypes. In addition, also soluble PGN structures activated the defense gene
expression in a LYM3/CERK1-dependent manner (Figure 3-12 and Willmann (2011)). The obvious lack of discrimination between differences in complexity and peptide bond of peptidoglycan resembles the broad ligand specificity of the mammalian PGN receptor Nod2 (Girardin et al., 2003b; Inohara et al., 2003), however the minimal PGN motif recognised by Nod2, the muramyl dipeptide (MDP), is not able to activate the plant immune response (Gust et al., 2007). Instead, the results obtained so far substantiate the significance of the glycan moiety for the Arabidopsis perception system. Mutanolysin-mediated complete digestion of the glycosyl bonds of peptidoglycan but not the cleavage of glycolglycine bonds within the peptide moiety by lysostaphin abolished the immunogenic properties of Sa PGN (Gust et al., 2007). Thus, it is likely that the chain length and possibly also the spatial structure of peptidoglycan are crucial parameters for the recognition. For instance, the Drosophila PGRP-LC cannot sense PGN monomers or dimers, indicating similar recognition preferences for longer glycan chains than the Arabidopsis PGN complex (Leulier et al., 2003). Moreover, the Drosophila PGRP-SA apparently perceives only PGN, which was preprocessed by the glucanase activity of GNBP1 (Filipe et al., 2005; Wang et al., 2006). Also plant LysM proteins seem to prefer carbohydrate ligands of specific chain length, because the rice LysM-domain receptor protein CEBiP was shown to require for binding chitin oligomers a degree of polymerisation (DP) > 6 (Ito et al., 1997; Okada et al., 2002).

Interestingly, the plant LysM proteins mediate via recognition of oligosaccharide structures as different processes as defense activation and symbiotic interactions. The lipochito-oligosaccharide-induced nodulation processes in leguminous plants require LysM-containing receptor kinases. For instance, the receptor pairs of NFR1/NFR5 (Figure 4-1) and LYK3/LYK4 are essential for the establishment of symbiosis with rhizobacteria in Lotus japonicus and Medicago truncatula respectively (Limpens et al., 2003; Radutoiu et al., 2003). Unfortunately, the formation of hetero-oligomeric complexes or direct binding to Nod factors has not been shown for neither in Lotus nor Medicago systems. The perception of chitin in rice is coordinated by the two receptor proteins CEBiP and OsCERK1 (Kaku et al., 2006; Miya et al., 2007; Wan et al., 2008) (Figure 4-1). In the presence of chitin oligosaccharides a portion of these proteins form a heteromeric complex in rice cells, however only CEBiP has been shown to bind to chitin (Kaku et al., 2006; Shimizu et al., 2010). In Arabidopsis, CERK1 is the only so far characterised LysM receptor protein essential for chitin-triggered activation defense responses and it also binds directly chitin in vitro (Iizasa et al., 2010; Miya et al., 2007) (Figure 4-1). The Arabidopsis PGN perception system mediated by CERK1 and LYM3 adds up to the versatility of LysM-protein functions in plants (Figure 4-1). As CERK1 displays no or only very weak PGN binding (Willmann, 2011), it seems to function exclusively as signal transducer upon PGN recognition by LYM3. There are in vitro indications for the
formation of an CERK1/LYM3 complex, as interaction was observed both in yeast two hybrid-assay and far western analysis (Willmann, 2011). Whether the functional PGN receptor consisting of CERK1 and LYM3 requires physical interaction between these two proteins \textit{in planta}, remains to be elucidated. Similarly, the possible participation of additional LysM proteins, like LYK3, within the recognition process is still unclear. Although all these plant carbohydrate ligand receptors share many structural characteristics with each other, there are yet differences in their binding and signaling properties rendering the needed specificity (Nakagawa et al., 2010; Wan et al., 2008). Intriguingly, the discovery of the \textit{Arabidopsis} CERK1/LYM3-based PGN perception system, that is analogous to the rice OsCERK1/CEBiP complex detecting chitin, indicates that plants employ two types of PRRs for the recognition of distinct types of PAMPs. The LRR-receptor kinases preferentially detect proteinous microbial signatures, whereas the LysM-domain containing receptor proteins contribute to plant innate immunity by perceiving carbohydrate-derived PAMPs.

\textbf{Figure 4-1: Plant perception and signaling of carbohydrate PAMPs/MAMPs}

Bacterial peptidoglycan is structurally similar to fungal chitin and rhizobacteria-derived lipochitooligosaccharides (Nod factors). The corresponding LysM receptor kinases or proteins from \textit{Arabidopsis thaliana} (At), \textit{Oryza sativa} (Os) and \textit{Lotus japonicus} (Lj) also share sequence similarities. OsCEBiP and OsCERK1 interact with each other \textit{in planta} (Shimizu et al., 2010). Based on Eckardt (2008).
4.2 CHIA chitinase is involved in plant innate immunity

PGRPs and lysozymes carry out various tasks in metazoan innate immunity. The enzymatically active PGRPs and lysozymes can act directly bactericidal but also modulate the defense responses by PGN hydrolysis (Callewaert and Michiels, 2010; Dziarski and Gupta, 2010; Royet and Dziarski, 2007). Plants possess neither lysozyme nor PGRP encoding genes. However, among plant chitinases are members, which harbor lysozyme-like peptidoglycan hydrolysis activity (Heitz et al., 1994; Majeau et al., 1990; Park et al., 2002). One such well-characterised enzyme is the class III chitinase hevamine from *Hevea brasiliensis* (Bokma et al., 1997). Lysozyme and hevamine both hydrolyse $\beta(1\rightarrow4)$ glycosidic bonds in the glycan backbone, however they differ in the cleavage sites. Lysozyme cleaves between MurNAc and GlcNAc (muramidase activity) and hevamine between GlcNAc and MurNAc (glucosaminidase activity) (Bokma et al., 1997; Tipper et al., 1964). Only one *Arabidopsis* chitinase could be identified sharing high sequence homology to the rubber tree hevamine, and this acidic endochitinase (CHIA) is additionally the only member of the class III chitinases (Figure 3-13 and Figure 3-14).

Chitinases similar to other plant PR proteins are characterised by the inducibility of their expression upon pathogen infection (Kasprzewska, 2003). The analysis of the expression profile of *CHIA* revealed that its transcription in leaves was induced not only upon fungal infection but additionally also upon infection with phytopathogenic *Pseudomonas syringae* (Figure 3-15). The verification of the microarray data was achieved using transgenic *CHIA* promoter-GUS reporter plants. As GUS activity can be irreversibly stained in the analysed leaf tissue even weak activation of the promoter can be visualised. The fungal infections with *Alternaria brassicicola* and *Botrytis cinerea* resulted in relatively low but visible activation of the *CHIA* promoter (Figure 3-16B). Earlier data showed similar *CHIA* promoter activation upon treatment with *Rhizoctonia solani* (Samac and Shah, 1991). Bacterial infection turned on the *CHIA* promoter in a strong manner, however only using the less-virulent *Pto hrcC* and the non-compatible *Pph* strains (Figure 3-16C). The *CHIA* promoter was not activated upon treatment with the virulent *Pto* DC3000 strain. These results point to effector-mediated suppression indicating that *CHIA* might be essential combating bacterial pathogens. The activation of the *CHIA* promoter took exclusively place within the infected tissue region in the leaf, suggesting a local host response only. The slight discrepancy between the results with *Pto* DC3000 infection in the microarray and in the promoter-GUS analysis might result from differences in the condition of the used strains (Figure 3-15 and Figure 3-16). Depending on the age of the plated culture, the bacteria can display differences in their virulence. It is possible that the strain used for the microarray was less aggressive still allowing some stimulation of the *CHIA* promoter, whereas the strain used for the infection of *pCHIA::GUS*
plants was highly virulent suppressing the promoter completely. Interesting was also the observation that peptidoglycan and chitin treatment led either to no or only to moderate activation of the CHIA promoter, but the strong PAMP flagellin induced the promoter almost as good as Pto hrcC (Figure 3-16A and C). Obviously not only pathogenic fungi and bacteria but also conserved and highly immunogenic bacterial signatures are sufficient to trigger the expression of the CHIA gene.

CHIA is expressed constitutively at low levels in natural openings like hydathodes and stomatal guard cells (Samac and Shah, 1991). Although the accumulation of CHIA protein in the apoplastic space connected to these specific leaf regions has not been shown yet, it suggests that CHIA is indeed a part of the secreted defense armory of the plant. Possibly, some CHIA protein is always present but that a pathogenic attack leads to its enhanced production and accumulation in the apoplast. This postulation is supported by the finding that CHIA was among the secreted cell wall proteins in non-stressed cultured Arabidopsis cells (Kwon et al., 2005).

4.2.1 Chitinolytic activity of CHIA and its impact on fungal immunity

Many plant chitinases have been characterised regarding their ability to hydrolyse fungal cell walls and to contribute to innate immunity (Arlorio et al., 1992; Boller et al., 1983; Mauch et al., 1988; Onaga and Taira, 2008). However, so far only few Arabidopsis chitinases have been functionally analysed (Passarinho and De Vries, 2002). The enzymatic properties of the Arabidopsis class III chitinase, CHIA, were assayed using protein extracts from transgenic CHIA overexpression and knock-down lines. CHIA protein was detectable in the overexpression plants via the GFP-tag but also using an antibody against tobacco class III chitinases (Figure 3-22A-C). The class III chitinase antibody seemed also to recognise the Arabidopsis class III chitinase present in high amounts in the CHIA-oe leaves (Figure 3-22B). However, in the WT leaves the CHIA amounts were probably too low for detection.

The CHIA-oe leaf extracts displayed a very strong hydrolytic activity towards a chitin-derivate, 4-MUCT, in comparison to the WT extract, which only showed basal activity (Figure 3-27A). Colloidal chitin was more or less equally hydrolysed by the CHIA-oe and WT extracts (Figure 3-27B). The obtained experimental data strongly implicates that CHIA is a functional enzyme possessing chitinolytic activity. The results further suggest, that 4-MUCT could be a chitin substrate, which is preferentially cleaved by the class III chitinase, whereas colloidal chitin might be rather favoured by the other Arabidopsis chitinases, present in both CHIA-oe and WT leaf extracts. In fact, similar strong activity towards 4-MUCT was also obtained for tobacco class III chitinases, whereas chitinases of the classes I, II, IV and V showed only
Discussion

little or no enzymatic activity (Brunner et al., 1998). The analysis of the protoplast samples showed that the majority of 4-MUCT degrading activity was harbored within the CHIA-o£ secreted protein fraction (Figure 3-28A). Unexpectedly, no differences were seen between the activity in the chia-kd and WT or secGFP control samples. This raises the question whether the induction of CHIA upon treatment with pathogenic fungi or the fungal PAMP chitin could result in differences in the different genotypes (see also chapter 4.2).

The observed chitin-degrading activity of extracts from CHIA-o£ plants led to the assumption that CHIA could contribute to host immunity. Thus, the CHIA overexpression lines might be more resistant and the CHIA knock-down lines more susceptible to fungal pathogens. Fungal infection assays with the necrotrophs Botrytis cinerea and Alternaria brassicicola revealed only insignificant but still interesting tendencies (Figure 3-29 and Figure 3-30). Especially in the case of Alternaria infection, the degree of disease symptoms was somewhat lower in the CHIA-o£ and higher in chia-kd plants, but still the differences were not significant (Figure 3-30A and D). It is feasible, that the chitinolytic activity of CHIA supports the fungal resistance in Arabidopsis, but that due to redundancy among the chitinase members, the loss of one chitinase is not dramatically weakening the plant immunity towards fungal pathogens. Indeed, up to now only little in vivo data is presented for the role of single chitinases in fungal resistance (Benhamou et al., 1990; Broque et al., 1991). Moreover, even the depletion of the sole chitin receptor, CERK1, leads only to a very weak fungal growth phenotype in Arabidopsis (Miya et al., 2007; Wan et al., 2008). Further examinations are needed to strengthen the supposition of CHIA being a part of the fungal resistance machinery using in vitro antifungal assays and analysis of mutants carrying multiple chitinase gene deletions.

4.2.2 PGN-lytic activity of CHIA and its importance to bacterial immunity

The chitinolytic activity observed in the transgenic plants overexpressing CHIA, demonstrated that this enzyme can cleave complex carbohydrate structures. Whether this Arabidopsis chitinase also degrades the heteromeric glycan backbone of peptidoglycan and displays a lysozyme-like activity characteristic for some plant chitinases of the classes III and V (Passarinho and De Vries, 2002), was determined in turbidity assays. The reduction of turbidity in a suspension of insoluble substrate, like purified peptidoglycan or bacterial cells containing intact cell walls, is a good indicator for enzymatic degradation of such complex structures into soluble and less turbid fragments. The PGN-hydrolysing activity of lysozyme, the standard PGN hydrolase used as reference in turbidity assays (Brunner et al., 1998; Park et al., 2002), leads to complete lysis of bacteria and clarification of the suspension within
short incubation times. The leaf protein extract of CHIA-oe displayed towards the Micrococcus luteus cell walls very high hydrolysis activity, whereas the activity of both WT and chia-kd extracts was low (Figure 3-27C). In the beginning, the activity of CHIA-oe extract was lower than that of lysozyme, reaching however similar reduction in turbidity after 4 hours. Since the approaches to purify CHIA from the leaf extracts by immunoprecipitation (Figure 3-22) or to heterologously express it in a highly active form (Figure 3-24 and Figure 3-25) failed so far, the determination of specific enzymatic properties of CHIA are still to be clarified. Despite the indications for a formation of CHIA dimers in vivo (Figure 3-22 and Figure 3-24), the homooligomerisation could not be observed in vitro (Figure 3-26). It is also possible that other yet unknown plant proteins present in the crude extract are interacting with and affecting the enzymatic properties of CHIA. Further experiments are needed to show in which functional form CHIA is operating and whether CHIA is interacting with components of the PGN receptor complex. Interestingly, in addition to the Gram-positive M.luteus cell walls containing Lys-type PGN, CHIA-oe extracts also hydrolysed purified Dap-type PGN from Bacillus subtilis (Figure 3-27D). This indicates that CHIA is able to cleave the glycosidic bonds in both PGN subtypes. Also the Arabidopsis PGN receptor proteins, LYM3 and CERK1, display similar broad PGN specificity. Not all peptidoglycans are suitable substrates for CHIA or other PGN hydrolases though. The CHIA-oe leaf extract was unable to degrade Staphylococcus aureus PGN (data not shown). Similar results have also been reported for lysozyme (Bera et al., 2007). The resistance against PGN degradation displayed by S.aureus enhances virulence and is due to peptidoglycan modifications including O-acetylation and heavy cross-linking and associated wall teichoic acids. The majority of the PGN-hydrolysing activity in the CHIA-oe leaves was harbored in the secreted protein fraction as shown by the protoplast assays (Figure 3-28B). Furthermore, the CHIA protein was detectable in the secretion sample (Figure 3-28C). These observations once more support the earlier indications of CHIA being targeted into the apoplast (see chapter 4.2).

Intriguingly, the PGN-degrading properties of CHIA seem to be linked to bacterial immunity. The infection of CHIA knock-down lines with the hypervirulent Pto DC3000 resulted in increased bacterial propagation in comparison to WT plants (Figure 3-31A). Even the hypovirulent P.syringae mutant strain lacking the effectors avrPto and avrPtoB (Pto DC3000 ΔavrPto/PtoB) was able to grow better in the chia-kd plants (Figure 3-31B). Thus, CHIA obviously contributes to bacterial resistance in Arabidopsis and can be regarded as an innate immunity protein. Similarly, the depletion of lysozyme M and the resulting decrease of PGN hydrolysis and antimicrobial activity led to enhanced susceptibility towards Streptococcus pneumonia infection in mice (Shimada et al., 2008). However, the analysis of the CHIA overexpression lines in bacterial infection assays delivered unexpected results. Instead of
being more resistant against *P. syringae*, the CHIA-oe lines showed similar susceptibility as the chia-kd lines upon infection with either hypervirulent or less-virulent strains (Figure 3-31A-C). Despite the dramatic differences in the PGN hydrolysis properties of CHIA overexpression and knock-down lines, both genotypes display a similar output phenotype being less resistant towards the Gram-negative phytopathogen *P. syringae*. One possible role of CHIA in defending *Arabidopsis* plants during bacterial attack is the release of immunogenic PGN fragments with a defined chain length, which upon recognition by a PGN receptor could enhance the activation of downstream responses. In such case, both the complete loss of PGN hydrolase activity but also too strong activity cleaving PGN into fragment lengths, which no longer stimulate the innate immune system, would have devastating effects. This correlates with the precedence of the rice LysM protein, OsCEBiP, to bind chitin chains only with a specific degree of polymerization (Ito et al., 1997; Okada et al., 2002) and might also be true for AtLYM3 regarding PGN binding. Hence, the detailed analysis of the PGN fragments produced by CHIA is crucial for understanding the molecular basis for its action. Another possibility is that CHIA is, like lysozyme, able to directly inhibit bacterial growth by massive lysis. However, this hypothesis would not explain why the CHIA overexpression lines are also more susceptible. Although the CHIA-oe plants do not show any dramatic phenotypic alterations, the presence of overexpression-derived artefacts cannot be totally excluded. The *in vitro* analysis of antibacterial activity with either highly purified *Arabidopsis* CHIA or recombinantly expressed CHIA could enlighten its mode of action. CHIA gets posttranslationally glycosylated (Figure 3-23), hence expression systems relying on eukaryotic hosts, such as yeast or insect cells, are needed for the production of active CHIA. In addition, it would also be important to test a Gram-positive phytobacterium in the bacterial infection assays to see whether the impact of CHIA, but also CERK1 and LYM3, is similar to that observed upon *Pseudomonas* infection. For example, the Gram-positive phytopathogen *Rhodococcus fascians* (Hogenhout and Loria, 2008) could be tested to see whether the broad PGN specificity observed for all three innate immunity proteins, CHIA, CERK1 and LYM3, also contributes to the formation and maintenance of resistance against both Gram-negative and Gram-positive pathogenic bacteria in *Arabidopsis thaliana*.

4.3 Peptidoglycan perception systems have arisen through convergent evolution in metazoans and plants

The experimental data obtained during this work and in Willmann (2011) revealed novel insights into peptidoglycan perception in plants allowing for the first time a kingdom-wide comparison. Similar to invertebrates and higher animals, also plants possess a peptidoglycan pattern recognition machinery operating in the innate immunity and especially
in the resistance against bacterial pathogens. Whereas the recognition of peptidoglycan structures in metazoans is mainly mediated by proteins containing LRRs or PGRP domains (Royet and Dziarski, 2007), the newly characterised plant PGN receptors are designated by the presence of LysM domains. Also PGN hydrolysing activity that contributes to host immunity is found across the phyla. So far, all identified plant PGN hydrolases belong to glucosaminidases, thus displaying different cleavage specificity than lysozymes (muramidase) or enzymatically active PGRPs (amidase) in animals (Bokma et al., 1997; Dziarski and Gupta, 2010; Park et al., 2002; Royet and Dziarski, 2007). Thus, the proteins involved in the PGN degradation and recognition processes vary not only in their domain features but also in their specificities pointing strongly to convergent evolution of these systems in the different eukaryotic lineages.
5 Summary

Plants as sessile organisms cannot escape, when they are confronted with harmful pathogens. Instead, they are weaponed with sophisticated and highly complex molecular responses that allow them to defend themselves. The innate immunity forms the basis for the self-defense of higher organisms, including mammals, invertebrates and plants. During the basal immune response, conserved microbial signatures are perceived by pattern recognition receptors and trigger a variety of defense reactions leading to protection of the plant tissue and resistance. The bacterial cell wall component peptidoglycan (PGN) is one of such conserved signatures activating the plant defense responses, however the molecular mechanisms of its recognition was until now not understood.

The importance of LysM-domain containing plant proteins in the recognition of carbohydrate ligands, such as chitin and lipochitooligosaccharides, has been elucidated in the last years (Kaku et al., 2006; Limpens et al., 2003; Madsen et al., 2003; Miya et al., 2007). Due to the structural similarities between chitin, lipochitooligosaccharide and peptidoglycan, members of this protein family provide interesting candidates for a putative PGN receptor. The reverse genetics approach performed in this work revealed a LysM receptor kinase, CERK1, to be involved in PGN perception. CERK1 is not only essential for the PGN-mediated activation of defense responses in Arabidopsis thaliana, it also contributes to bacterial resistance. Consequently, CERK1 is a dual player within plant immunity, as it is important for both the recognition of carbohydrate signatures derived of both fungi and bacteria. The in parallel identified LysM protein, LYM3, binds PGN and is also essential for PGN-responsiveness and bacterial resistance (Willmann, 2011). As LYM3 lacks a signaling domain the obtained data suggest that CERK1 and LYM3 form a functional receptor complex and are both needed for PGN-triggered immunity towards bacterial infection.

The second focus of this work was to analyse the possible PGN processing properties of Arabidopsis. Studies on analogous, complex polymeric ligands, like fungal chitin and oomycete ß-glucan and the corresponding perception systems in plants (Fliegmann et al., 2004; Ito et al., 1997), but also peptidoglycan perception in animals (Filipe et al., 2005; Leulier et al., 2003) gave indications for such processing events. In addition, both complex and fragmented peptidoglycans act immunostimulatory in Arabidopsis, and are recognised by the same CERK1/LYM3-based receptor system. Therefore, the class III chitinase CHIA, a putative PGN hydrolase, was analysed regarding its role in peptidoglycan degradation and bacterial immunity. CHIA possesses both chitinolytic and peptidoglycan-hydrolysing activity, as observed with CHIA-oe leaf protein extracts but also with protein derived from the secreted fraction of CHIA-oe protoplasts. Both Gram-positive and Gram-negative
peptidoglycan subtypes can be cleaved by CHIA. The posttranslational glycosylation and apoplastic localisation of CHIA depend on the N-terminal secretion signal. Absence or excess of CHIA-dependent PGN hydrolysis affects the plant defense response towards bacterial pathogens in a negative manner, suggesting that CHIA is a plant innate immunity protein contributing to bacterial resistance.
6 Zusammenfassung


Im zweiten Teil dieser Arbeit wurde Fokus auf die PGN Prozessierung in Arabidopsis gelegt. Studien an analogen und ähnlich komplexen polymeren Liganden, wie zum Beispiel Chitin aus Pilzen und ß-Glucan aus Oomyceten, und den dazugehörigen pflanzlichen Erkennungssystemen (Fliegmann et al., 2004; Ito et al., 1997), aber auch der PGN-Perzeption der Tiere (Filipe et al., 2005; Leulier et al., 2003) lieferten Indizien für solche Prozessierungsvorgänge. Darüber hinaus stimulieren sowohl komplexes als auch fragmentiertes Peptidoglycan die Immunantwort in Arabidopsis und beides wird über das
References


Bera, A., Herbert, S., Jakob, A., Vollmer, W., and Götz, F. (2005). Why are pathogenic staphylococci so lysozyme resistant? The peptidoglycan O-acetyltransferase OatA is the


# Appendix

Table 8-1 summarizes the primers used in the frame of this work.

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Table 8-1: Used oligonucleotides
Figure 8-1: ClustalW2 protein sequence alignment of CHIA (At5g24090) and selected plant class chitinases

A multiple sequence alignment of the full-length protein sequences of Arabidopsis thaliana CHIA, Hevea brasiliensis HevamineA, Capsicum annuum chitinase I, Medicago truncatula chitinase, Nicotiana tabacum chitinase, Oryza sativum chitinase III and Vitis vinifera chitinase II using the ClustalW2 algorithm. Black boxes indicate differences in the amino acid residues.
Danksagung

Diese Arbeit wurde vom November 2007 bis August 2011 am Institut für Pflanzenbiochemie des Zentrums der Molekularbiologie der Pflanzen der Eberhard-Karls-Universität in Tübingen angefertigt.

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Lebenslauf

Heini Marjatta Grabherr

Geburtsdatum und -ort  15.12.1981 in Helsinki, Finnland
Staatsangehörigkeit  finnisch
Familienstand  verheiratet

Schulbildung

1997 – 2000    Gymnasium in Savonlinna, Finnland
Abschluss der allgemeinen Hochschulreife

Hochschulbildung

2002 – 2007     Studium der Biologie an der Johann Wolfgang Goethe-Universität Frankfurt am Main mit den Schwerpunkten Genetik, Mikrobiologie und Zell- und Entwicklungsbiologie
Mündliche Diplomprüfungen in allen Schwerpunktfächern
Titel der Arbeit: „Functional characterization of HsfC1 in Arabidopsis thaliana“
Seit 11/2007 Promotion bei Prof. Dr. T. Nürnberger am Institut für Pflanzenbiochemie des Zentrums für Molekulare Biologie der Pflanzen an der Eberhard Karls-Universität Tübingen
Titel der Arbeit: „Characterisation of the role of LysM-receptor-like kinases and the CHIA chitinase in the perception of peptidoglycan and in the innate immunity of Arabidopsis thaliana“