Characterisation of the *AT4G11100* gene, a negative regulator of disease resistance in *Arabidopsis thaliana*.

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List of Abbreviations

°C  degree Celsius
μ    micro
Avr  avirulence
bp   base pair(s)
cDNA complementary deoxyribonucleic acid
cfu  colony forming units
cir1 constitutively induced resistance 1 mutant
Col-0 Arabidopsis thaliana ecotype Columbia
DEPC diethylpyrocarbonate
DNA deoxyribonucleic acid
dNTP deoxynucleosidetriphosphate
DTT  dithiothreitol
EDTA ethylenediamine tetraacetic acid
ETI  effector-triggered immunity
g    gram(s)
g    gravity constant (9.81 ms⁻¹)
eGFP enhanced green fluorescent protein
h    hour(s)
HR   hypersensitive response
JA   jasmonic acid
kb   kilobase(s)
KB   King’s broth
RLU  relative light units
RNA  ribonucleic acid
SA   salicylic acid
SAR  systemic acquired resistance
SDS  sodium dodecyl sulphate
s    second(s)
U    unit(s)
UTR  Untranslated region
UV   ultraviolet
V    Volt
v/v  volume per volume
w/v  weight per volume
**Abstract**

Plants have evolved a complex system of defence to prevent pathogen establishment. The *Arabidopsis thaliana* cir1 (*constitutively induced resistance 1*) mutant displays enhanced resistance to infection by the virulent bacterial pathogen *Pseudomonas syringae* and constitutively expresses a number of defence genes. Evidence suggests that CIR1 is a negative regulator of plant immunity important in the absence of pathogen attack. Genetic mapping experiments indicate that *cir1* is located on the lower arm of chromosome 4 of *A. thaliana* and may be one of 8 known genes in the region. Analysis of T-DNA knockouts of these 8 genes suggests that *AT4G11100* is the mostly likely candidate for CIR1. This project established that the disease resistance phenotype of *cir1* is temperature dependent and linked to reduced plant growth. Genetic crosses between *cir1* and *at4g11100* T-DNA knockout mutants revealed that the mutants complement and therefore *AT4G11100* is not CIR1. However, like *cir1*, the *at4g11100* T-DNA knockout mutants display enhanced disease resistance. Over expression of *AT4G11100* leads to increased susceptibility to infection by *Pseudomonas syringae* (Pst) and reduced induction of the salicylic acid defence gene *PR2* following Pst infection, suggesting that *AT4G11100* may too be a negative regulator of immunity. Additionally, a plant line with exceptionally high *AT4G11100* expression levels displayed distinct leaf morphology, possibly implicating *AT4G11100* in leaf development. Unfortunately, efforts to determine the subcellular localisation of AT4G11100 were unsuccessful, potentially due to problems with the expression of an *AT4G11100::eGFP* fusion, or even the instability of the resulting fused protein. Thus AT4G11100 is a likely negative regulator of plant immunity, and the identity of CIR1 remains unknown. Parts of this research have been recently published (Carstens, M., McCrindle, T.K., Adams, N., Diener,
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Chapter 1: Introduction

Human population levels have dramatically increased over the last ten thousand years, and further growth is predicted to put the global population at between 9 and 11 billion by 2050 (Cohen, 2003). This growth will require that crop yields are improved in order to meet intensified food production targets. However, global food security is threatened by a number of factors, chiefly: high levels of poverty in the developing world, environmental degradation, climate change, and emergent plant pathogens (Godfray et al., 2010). Plant pathogens in particular are problematic, with more than 10% of the annual global crop harvest lost to pests and diseases (Strange & Scott, 2005) and microorganisms alone accounting for more than 200 billion $US worth of crop losses every year (Horbach et al., 2011). The lack of genetic diversity within crop species, and the continued practice of industrial scale monocultures, leaves many agriculturally important plants vulnerable to pathogen attack.

The development of transgenic plants able to resist infection by virulent pathogens has the potential to improve crop yields. It is therefore important to gain an understanding of the molecular mechanisms that underlie the interactions between plants and plant-pathogens. The convergence of studies of plant immunology and pathogen infectivity is central to elucidating how plant-pathogen interactions can be manipulated to enhance disease resistance (Dodds & Rathjen, 2010).
Plant immunity

To prevent pathogen establishment and disease, plants have evolved a complex system of defence. This defence system is analogous to the innate immune system present in vertebrates, but otherwise differs from animal immunity in a number of important ways. The most striking of these differences is that, unlike vertebrates, plants lack a circulatory system that would accommodate adaptive immunity, and so rely solely on an innate immune system (Boller & Felix, 2009).

The innate immune system involves the recognition of non-host (or non-self) entities. The ability of a plant species to resist infection by an entire pathogen species is called non-host resistance (Nuernberger & Lipka, 2005). Non-host resistance is the most common mechanism of plant defence, and enables protection against a wide range of pathogens by utilizing both preformed defences and inducible responses (Mysore & Ryu, 2004). The waxy cuticle and cell wall of epidermal cells, as well as actin microfilaments in the cytoskeleton, act as an effective barrier to pathogen entry into the intracellular space (Jones & Takemoto, 2004). Together with these physical barriers, the constitutive production of numerous antimicrobial enzymes and secondary metabolites make up the non-host preformed defence.

PAMP Triggered Immunity (PTI)

The inducible responses of non-host defence can be divided into two primary branches, each recognizing a different type of non-host molecule. The first response involves the recognition of highly conserved microbial proteins called pathogen associated molecular patterns (PAMPs) by transmembrane pattern recognition receptors (PRRs) (Zipfel & Felix,
This recognition induces a receptor-mediated defence response known as PAMP triggered immunity (PTI), which is characterised by the activation of basal defence mechanisms that include the activation of mitogen-activated protein kinases (MAPKs), oxidative bursts, plant hormone production, and the expression of defence related genes (Felix & Boller, 2003).

The activation of initial MAPKs stimulates several MAPK signalling cascades. These kinase cascades act as important signalling mediators between pathogen sensors and cellular defence responses (Meng & Zhang, 2013), and work as part of a complex signalling network that leads to the biosynthesis of plant hormones (including salicylic acid, ethylene, and jasmonic acid) and the induction of pathogenesis related (PR) defence gene expression.

A commonly cited example of PTI is the detection of the flg22 peptide of bacterial flagellin by the PRR FLAGELLIN SENSING 2 (FLS2). The flg22 peptide is a highly conserved 22 amino acid region of the flagellin protein, the primary constituent of bacterial flagella, which are used by bacteria for locomotion (Gómez-Gómez & Boller, 2000). Following detection of flg22, FLS2 associates with the receptor like kinase (RLK) BRI1-ASSOCIATED RECEPTOR KINASE 1 (BAK1), forming a heterodimer that is thought to initiate MAPK signalling cascades (Meng & Zhang, 2013); although the exact components involved in this initiation are still unknown, it has been established that FLS2/BAK1 complex is important for the establishment of innate immunity (Chinchilla et al., 2007).

Relatively little is known about the fundamental molecular mechanisms connecting PRR activation to initiation of MAPK signal transduction, however, the importance of MAPK
signalling cascades in activating cellular defence responses is well established (Colcombet & Hirt, 2008). For instance, following flg22 detection in A. thaliana, two MAPK signalling cascades are activated, whereupon the sequential transfer of phosphate groups is initiated; phosphate groups are transferred from MAPK kinase kinases (MAPKKKs) to MAPK Kinases (MAPKKs) and then onto MAPKs, which go on to regulate immunity (Nakagami, Pitzschke & Hirt, 2005). The first of these cascades consists of MEKK1 (a MAPKKK), MKK4/MKK5 (two redundant MAPKKs), and MPK3/MPK6 (two redundant MAPKs), with MPK3/MPK6 positively regulating the defence response (Asai et al., 2002). Similarly, the second cascade involves phosphate transfer beginning with MEKK1, but then branches to MKK1/MKK2 (two redundant MAPKKs), and MPK4, with MPK4 positively regulating basal defence and negatively regulating resistance protein mediated effector triggered immunity (Zhang et al., 2012).

**Effector Triggered Immunity (ETI)**

Because PAMPs are highly conserved molecules that are not easily lost or modified, it is difficult for pathogens to evolve PAMPs that do not trigger PTI. Instead, pathogens have evolved effector molecules able to subvert PTI, resulting in effector-triggered susceptibility (ETS) of the host plant (Jones & Dangl, 2006). Effector producing pathogens are able to directly introduce (via the type III secretion system) a wide variety of effector molecules into host cells, some of which are protein kinase inhibitors that prevent MAPK signalling and thus downstream defence activation (Xiang et al., 2008); in this way effector producing pathogens are able to greatly enhance their virulence (Dodds & Rathjen, 2010). Other effectors, such as AvrPto from *P. syringae*, interact directly with the intracellular kinase
domains of the PRRs themselves, preventing the autophosphorylation that occurs following binding of their PAMP ligands, and thus blocking downstream signalling (Xiang et al., 2008).

Enhanced pathogen virulence has exerted a selective pressure on host plants leading to the evolution of plant resistance (R) proteins, many of which are nucleotide-binding and leucine-rich repeat (NB-LRR) proteins (Eitas & Dangl, 2010). NB-LRR proteins are divided into two categories based on the structure of the N-terminus; TIR-NB-LRRs contain an N-terminus homologous to the animal immune system Toll and interleukin 1 receptor (TIR), whereas CC-NB-LRRs contain an N-terminus that possesses a coiled coil (CC) domain (Dangl & Jones, 2001; Gómez-Gómez, 2004). R proteins, encoded by R genes, recognise specific effector molecules in a gene-for-gene manner and are able to induce effector triggered immunity (ETI) by re-activating the MAPK signalling cascades (Abramovitch & Martin, 2004; Oh & Martin, 2011). ETI constitutes the second branch of the inducible non-host defence response, and differs from PTI in that it elicits a rapid, amplified and often prolonged defence reaction characterised by a hypersensitive response (HR) and programmed cell death.

Although there are some cases of direct recognition and interaction between R proteins and effector molecules (Dodds et al., 2006), most R proteins recognise effectors indirectly, usually by detecting effector induced modifications of certain host proteins, often components of PTI. This indirect recognition is known as the guard hypothesis, and a well-studied example of the guard hypothesis involves the “guardee” RPM1 INTERACTING PROTEIN 4 (RIN4). Phosphorylation of RIN4 is induced by P. syringae effectors, AvrB or AvrRpm1, and this modification of RIN4 is detected by the NB-LRR protein RPM1, which is
able to elicit ETI (Mackey et al., 2003; Liu et al., 2011). A third *P. syringae* effector, AvrRpt2, actually cleaves RIN4, and the resulting degradation of RIN4 is recognised by a second NB-LRR protein, RPM2, which is in turn able to activate ETI (Mackey et al., 2003). In this scenario, RPM1 and RPM2 act as “guards” of RIN4 and are able to detect modifications induced by AvrB, AvrRpm1, or AvrRpt2.

Another defence strategy evolved by plants involves the use of “decoys” to interfere with pathogen effector function. In the case of flg22 detection by the FLS2 PRR, the AvrPto effector molecule has been identified as a PTI suppressor upstream of MAPK cascade activation (He et al., 2006). AvrPto is a protein kinase inhibitor that binds FLS2, inhibiting autophosphorylation and thereby suppressing the MAPK cascade that would lead to defence activation (Xiang et al., 2008). Two plant resistance proteins are involved in the perception of AvrPto, namely PTO, a serine/threonine kinase, and PRF, a NB-LRR protein (Lin & Martin, 2007). PTO and PRF have been shown to constitutively interact in a manner that prevents any downstream signal initiation via PRF; however, in the presence of AvrPto, the effector molecule binds to PTO (potentially in competition with FLS2) which releases PRF from its interaction with PTO, and allows for activation of ETI responses (Xing et al., 2007). Here, PTO is thought to act as a “decoy” by having a kinase domain structurally similar to that of FLS2. Thus, AvrPto binds to both FLS2 and PTO, but whereas binding to FLS2 inhibits plant defence induction, binding to the PTO “decoy” has the opposite effect and results in the activation of ETI (van der Hoorn & Kamoun, 2008).
PTI and ETI, through various signalling pathways, elicit the expression of PR genes that enhance host resistance to infection. The inducible responses from both these defence stages lead to a third and final layer of innate immunity, namely, systemic acquired resistance (SAR). SAR occurs when the expression of PR genes in infected tissue causes the expression of the same PR genes in uninfect ed distal tissue, thereby conferring enhanced resistance to the rest of the plant and protecting from secondary infection (Fu & Dong, 2013). SAR is controlled via the production of the salicylic acid (SA) immune signal, which is able to trigger wide-spread transcriptional reprogramming and expression of PR genes (Durrant & Dong, 2004).

Figure 1 illustrates the example of PTI induction via the detection of the flg22 peptide, the suppression of PTI through the introduction of bacterial AvrPto effector molecules, and the induction of ETI through R protein mediated detection of AvrPto.
Figure 1) **An example of PAMP triggered immunity (PTI) and effector triggered immunity (ETI).** Detection of the flg22 pathogen associated molecular pattern (PAMP) by the FLAGELLIN SENSING 2 (FLS2) PAMP recognition receptor (PRR) triggers dimerization with BRI1-ASSOCIATED RECEPTOR KINASE 1 (BAK1) which initiates a cascade of mitogen-activated protein kinase (MAPK) signalling. MAPK signalling cascades lead to the induction of PTI through sequential transfer of phosphate groups. PTI is subverted by pathogen AvrPto effector molecules, introduced into the plant cell via the type three secretion system (TTSS). AvrPto prevents phosphorylation at the beginning of the MAPK cascade by binding to FLS2. ETI is induced when AvrPto effectors bind to the PTO serine/threonine kinase, which relaxes suppression of the PRF resistance (R) protein and leads to the activation of defence responses, including the hypersensitive response (HR).
Negative regulators of plant immunity

Plant defence responses are tightly controlled by a number of interconnected and complex networks of signalling pathways. These pathways interact in various ways to ensure appropriate defence responses are mounted to combat specific threats at specific times. Importantly, plants are able to suppress immune responses in the absence of pathogens. Activation of defence is energetically expensive and can have detrimental effects on overall plant health; for example, a number of mutants with constitutively active defence responses, including constitutive expressor of PR genes 1 (cpr1), suppressor of npr1-1, constitutive 1 (snc-1), and suppressor of rps4-RLD 1 (srfr1), display stunted growth and reduced reproductive capabilities (Zhang et al., 2003; Gou et al., 2009; Kim et al., 2010).

There are numerous proteins involved in the negative regulation of plant immunity; one such is the kinase-associated protein phosphatase (KAPP), which is able to bind to the kinase domains of a number of PRRs (including FLS2) and hinder autophosphorylation and downstream MAPK activation (Gómez-Gómez, Bauer & Boller, 2001). KAPP is an example of a moderator of the initial PTI, down-regulating this response once pathogen attack has abated.

Many of the key negative regulators of immunity work via controlling degradation of their target proteins. For example, NON-EXPRESSER OF PR GENES 1 (NPR1) is required for SA-induced PR gene expression during defence (Dong, 2004), and in the absence of SA, NPR1 is bound by its parologue NPR4 which marks it for degradation by the proteasome (Fu et al., 2012). NPR4 is an adapter protein of the Cullin 3 (CUL3) E3 ligase which is able to mediate the degradation of its substrate (NPR1) in a SA dependent manner. In the absence of SA,
NPR1 is bound by NPR4, but when SA is present it will bind NPR4, reducing its affinity for NPR1 and thereby enabling NPR1 to activate signalling. NPR4 therefore acts as a negative regulator of immunity in the absence of pathogen attack by degrading NPR1 and preventing PR gene expression.

A negative regulator of ETI has already been discussed; the PTO kinase supresses ETI in the absence of pathogen introduced effector molecules by constitutively interacting with and deactivating the PRF NB-LRR protein (Xing et al., 2007). A similar example involves the MPK4 MAPK, which (as mentioned above) positively regulates basal defence but negatively regulates R protein mediated ETI (Zhang et al., 2012). Here, the PTI MAPK cascade is targeted by the HopAI1 effector molecule, which binds and inactivates MPK4. However, the inactivation of MPK4 relaxes the suppression MPK4 exerts on the NB-LRR protein SUMM2, leading to activation of ETI. In this way it is possible for a plant to modulate its defence response in the presence of PAMPs but not effector molecules, ensuring that only PTI is induced and the HR is avoided.

Figure 2 illustrates the examples mentioned above; these negative regulators form part of greater regulatory networks that, together, enable plants to finely control defence activation. Negative regulators of immunity are vital for suppressing defence in the absence of pathogens, and counteracting the fitness costs of constantly mounting immune responses.
Negative regulation of PAMP triggered immunity (PTI) and effector triggered immunity (ETI) in the absence of pathogen attack. To prevent energetically expensive activation of defence responses in the absence of pathogen attack plants possess a number of negative regulators. The kinase-associated protein phosphatase (KAPP) directly inhibits autophosphorylation of the FLAGELLIN SENSING 2 (FLS2)/BRI1-ASSOCIATED RECEPTOR KINASE 1 (BAK1) heterodimer. This inhibits the sequential transfer of phosphate groups between mitogen-activated protein kinases (MAPKs) in the MAPK signalling cascade, and thus the induction of PTI is prevented. ETI activation by the resistance (R) protein PRF is prevented by constitutive interaction between PTO (serine/threonine kinase) and PRF. In the presence of pathogen associated molecular patterns (PAMPs), but not effectors, excessive activation of ETI is avoided though the suppression of the R protein SUMM2 by the MPK4 MAPK.
**Environmental temperature affects plant defence**

The effects of environmental temperature on plants are varied and depend, at least in part, on numerous other factors, including water availability, soil nutrients, and plant species (Garrett *et al.*, 2006). As with plant growth and development, there are numerous examples of how temperature affects plant defence, with abnormally high or low temperatures during growth inhibiting disease resistance (Zhu, Qian & Hua, 2010). Heat sensitivity of *R* gene-mediated disease resistance has been observed in plants exposed to bacterial, fungal, and viral infection (Whitham, McCormick & Baker, 1996; Xiao *et al.*, 2003; Wang *et al.*, 2009), with disease resistance abolished above temperatures between 28°C and 30°C.

There are a number of mutants that display temperature dependent enhanced disease resistance; one such is bonzai1 (*bon1*), a loss-of-function mutant that exhibits a constitutive defence response at 22°C, but not at 28°C (Yang & Hua, 2004). The *bon1* mutation leads to activation of *SUPPRESSOR OF NPR1-1, CONSTITUTIVE 1* (*SNC1*), an NB-LRR type *R* gene, resulting in downstream defence activation and SAR. The activation of *SNC1* and the consequent constitutive defence response, cause a dwarf phenotype in *bon1* at 22°C; however, at 28°C *SNC1* expression is suppressed, resulting in a loss of defence activation and no dwarf phenotype.

A *SNC1* gain-of-function mutant (*snc-1*) constitutively expresses *SNC1* and displays dwarfism, constitutive expression of defence genes and increased resistance to biotrophic pathogens. Like *bon1*, these phenotypes in *snc-1* are temperature dependant, with plants grown at 28°C not displaying dwarfism or enhanced defence (Zhang *et al.*, 2003).
Additionally, progeny of *bon1* and *snc1-11* (a null mutant) do not exhibit the *bon1* phenotypes, indicating that dwarfism and enhanced defence at 22°C are SNC1 dependent.

Dwarfism and temperature dependent disease resistance in other mutants (*cpr1* and *srfr1*) have also been shown to be SNC1 dependent (Gou *et al.*, 2009; Kim *et al.*, 2010), suggesting that SNC1 may be a temperature sensor that modulates immunity in response to environmental changes. The *snc1-3* mutant (a constitutive expresser of *SNC1*) provides further evidence for this theory as it exhibits both dwarfism and enhanced resistance to virulent biotrophic pathogen *P. syringae* (*Pst DC3000*) at both 22°C and 28°C (Zhu, Qian & Hua, 2010). Interestingly, the fact that SNC1 nuclear content decreases with increasing temperature in wild type plants, but not in the *snc1-3* mutant could suggest that a threshold SNC1 concentration is required in the nucleus in order to trigger immunity.

**The cir1 mutant**

Loss-of-function mutants that display constitutive expression of defence genes, like *bon1*, likely encode proteins that act as negative regulators of plant immunity. As discussed above, negative regulators occupy key positions in the plant defence signalling network, and enable tight regulation of defence gene expression (Trujillo *et al.*, 2008), importantly preventing immune activation in the absence of pathogens. Therefore the identification of the *constitutively induced resistance 1* (*cir1*) mutant as a potential negative regulator of SAR (Murray *et al.*, 2002) heralded a potentially interesting avenue for investigation.

The *cir1* mutant was identified following a mutant screen for increased luciferase activity in wild type *Arabidopsis thaliana* plants carrying a *PR-1:LUC* reporter construct. Increased
luciferase activity in the recessive cir1 mutant compared to both wild type and PR-1:LUC lines indicated elevated activity of the PR-1 promoter. Indeed, Northern blot analysis revealed the constitutive expression of a number of defence genes in cir1, including three PR genes, PLANT DEFENSIN 1.2 (PDF1.2), and GLUTATHIONE S-TRANSFERASE 1 (GST1) (Murray et al., 2002). Additionally, cir1 shows enhanced resistance to infection by the virulent biotrophic pathogens Pseudomonas syringae (Pst DC3000) and Hyaloperonospora arabidopsis.

The cir1 mutant shows enhanced resistance to infection by Pst DC3000 and to determine whether NPR1 is required for this defence phenotype, cir1 npr1 double mutants were created. It was found that npr1 reduced PR-1 expression, suggesting that NPR1 is required for cir1 mediated PR-1 expression. Additionally, the npr1 mutation was found to partially suppress disease resistance in the cir1 npr1 double mutants suggesting that both npr1-dependent and -independent SA-mediated resistance contribute to disease resistance in cir1 (Murray et al., 2002). The fact that the cir1 null mutant displays constitutive expression of defence genes, and enhanced resistance to Pst DC3000, suggests that CIR1 is a likely negative regulator of plant defence.

Genetic mapping experiments and segregation analysis indicate that cir1 is located on the lower arm of chromosome 4 of A. thaliana in a region containing 8 known genes (Carstens, 2008). Of these 8 genes, AT4G11100 has been identified as the most likely candidate for CIR1; an at4g11100 knockout line, containing a T-DNA insertion in the promoter region, displays elevated levels of PR-1 and EDS1 proteins, as well as enhanced resistance to infection by Pst DC3000 (Diener, 2012). AT4G11100 is a protein of unknown function that
has, interestingly, been shown to interact with two NB-LRR RLKs (AT2G36570 and AT3G50230) \textit{in vitro} during yeast two hybrid assays (Mukhtar \textit{et al}., 2011). Both RLKs appear to be involved in plant development, with evidence that AT2G36570 (\textit{PXC PXY/TDR-CORRELATED GENE 1}, or \textit{PXC1}) is important for secondary cell wall formation in xylem fibres (Wang \textit{et al}., 2013). Although \textit{in silico} analysis based on information from The \textit{Arabidopsis} Information Resource (TAIR) broadly implicates \textit{PXC} genes in defence and abiotic stress responses, Wang \textit{et al}.
(2013) found no clear link between \textit{PXC1} and plant defence.

\textbf{Project objectives}

Relatively early in the course of the project it was discovered the \textit{cir1} mutant displays a temperature sensitive growth phenotype. Since a number of other gain-of-resistance mutants display temperature dependant size and disease resistance, this novel finding prompted investigation into the effects of temperature on disease resistance in the \textit{cir1} mutant.

The initial aim of the project was to confirm the role of \textit{AT4G11100} as the gene responsible for the \textit{cir1} disease resistance phenotype, as suggested by Diener (2012). This was to be done by characterising a second independent \textit{AT4G11100} T-DNA insertion mutant, to ensure that the resistance phenotype displayed by the \textit{AT4G11100} T-DNA insertion mutant previously characterised (Diener, 2012) was due to the disruption of \textit{AT4G11100} and not due to multiple T-DNA insertions. While this second T-DNA insertion mutant did display the same phenotype as \textit{cir1}, complementation of \textit{cir1} and the \textit{at4g11100} mutants showed that \textit{AT4G11100} was not in fact \textit{CIR1}, but was nonetheless a likely negative regulator of plant immunity. This hypothesis was tested by generating \textit{AT4G11100} over-expressing transgenic
plants and analysing their resistance to *Pst* DC3000 and expression of several defence genes after infection. Finally, an attempt to determine the subcellular localisation of the AT4G11100 protein through the generation of eGFP-fusion proteins was undertaken.
**Chapter 2: Methods and Materials**

**Plant Material and Growth Conditions**

**Plant lines**

The wild type *Arabidopsis thaliana* seeds used in experiments were of the Columbia (Col-0) ecotype. The cir1 and PR1::LUC lines (in the Col-0 background) were obtained from Shane Murray (University of Cape Town). The T-DNA insertion lines SALK_062847C (promoter insertion) and SALK_096586 (exon 2 insertion) were obtained from the Nottingham Arabidopsis Stock Center (NASC) (Scholl, May & Ware, 2000).

**Soil-grown plants**

Seeds were hydrated in 0.1% (w/v) agar and left to stratify for 48 h at 4°C in the dark before being sown on a 1:1 mixture of peat (Jiffy Products, International AS, Norway) and vermiculite. Pots were covered with cling film to prevent soil desiccation and then placed at a constant temperature (22°C) under fluorescent light (100µM photons m⁻²s⁻¹) with a 16 h light/ 8 h dark cycle. The cling film was removed after seven days; any excess seedlings were removed at the same time so that each pot contained only an individual plant.

**Seed sterilization**

Seeds sown on agar were sterilized in a lamina flow cabinet before plating. Seeds were placed in 1.5mL Eppendorf tubes and washed with 70% (v/v) ethanol for 5 min (tubes were inverted periodically). After washing, the 70% ethanol was poured off and replaced with 100% ethanol which was immediately aspirated. Seeds were allowed to dry on sterilized filter paper before being sown onto agar.
**Agar-grown plants**

Seeds were plated onto half strength Murashige and Skoog (MS) agar plates (8% w/v). MS agar pH was adjusted to 5.7 using 0.1M potassium hydroxide (KOH). Seeds were allowed to stratify on the plates for 48 h at 4°C in the dark, after which plates were placed at a constant temperature (22°C) under fluorescent light (100μM photons m$^{-2}$s$^{-1}$) with a 16h light/ 8h dark cycle.

**Kanamycin selection**

Plants expressing the kanamycin resistance gene (*NptII*) are resistant to the kanamycin antibiotic, and *NptII* homozygous lines (which display 100% resistance) can be selected for by growth on MS agar (0.8% w/v) containing 50mg/mL kanamycin. Between 1 and 2 weeks of growth, kanamycin sensitive plants can be distinguished from resistant individuals by their stunted leaf and root growth. Kanamycin resistant individuals were transferred to pots of soil and allowed to grow under normal soil growth conditions.

**Microbial Strains and Plant Infection**

**Escherichia coli**

*E. coli* strains DH5α and One Shot® (Life Technologies) were cultured in Luria-Bertani (LB) media (Sambrook *et al*, 1989) on either plates (1.5 % w/v agar) or in liquid culture, containing plasmid-dependent selective antibiotic(s). Bacteria were cultured at 37°C for 16 h overnight, with constant shaking (250rpm) for liquid cultures.
**Agrobacterium tumefaciens**

The *A. tumefaciens* strain GV3101 (Holsters *et al*., 1980) was cultured on LB agar plates (1.5% w/v agar) containing rifampicin (150 μg/mL) and gentamycin (15 μg/mL) at 28°C in the dark.

**Pseudomonas syringae**

The virulent strain *P. syringae* pv. *tomato* DC3000 (*Pst* DC3000) (Whalen *et al*., 1991) was cultured in King’s Broth (KB) medium (King *et al*., 1954) on either plates (1.5% w/v agar) or in liquid culture, containing 50 μg/mL rifampicin. Bacteria were cultured at 30°C for 16 h overnight, with constant shaking for liquid cultures.

**A. thaliana infection assays**

Four week old soil grown *A. thaliana* plants were infected with a virulent *P. syringae* strain, *Pst* DC3000, according to the following protocol. A 5 mL overnight *P. syringae* culture was centrifuged and cells were then washed and resuspended in 10mM MgCl₂. Cells were diluted to give a final OD₆₀₀ of 0.002, measured using the Beckman DU 650 Spectrophotometer (Beckman Coulter, Inc., CA, USA), in a total volume of 25mL. An OD₆₀₀ of 0.002 corresponds to 1 x 10⁶ colony forming units (c.f.u)/mL (Katagiri *et al*., 2002). Three leaves per plant were pressure infiltrated with the bacterial suspension using a needleless syringe. Five plants per plant line per time point (for harvesting at 4 or 48 h post infection) were infiltrated and an additional plant per line was infiltrated with 10mM MgCl₂ as a mock infection negative control. Infiltrated plants were covered with cling film and placed under normal growth conditions for 48 h. Infiltrated leaves were harvested at 4 h and 48 h post infection for bacterial growth analysis. Bacterial concentrations were determined by grinding three 0.5cm² leaf discs per plant (one for each infiltrated leaf) in 1mL 10mM MgCl₂.
and making a serial dilution of the resulting bacterial suspension. Each dilution was then plated onto KB agar plates (50 μg/mL rifampicin) and incubated for 48 h at 30°C. Colony forming units were recorded for each sample and bacterial titre per cm² leaf area calculated.

**DNA Manipulation**

**Isolation of genomic DNA from *A. thaliana***

*A. thaliana* genomic DNA was isolated from 100mg of leaf tissue per sample using a modified version of the Dellaporta (1983) DNA mini-preparation technique. The following modifications were made to the original protocol: Leaf tissue was not frozen with liquid nitrogen before being ground, but was homogenised directly in the extraction buffer.

**Isolation of bacterial plasmid DNA**

For moderate yields, bacterial plasmid DNA was isolated using the Wizard Plus DNA Purification System (Promega Corporation, Madison, US) according to the manufacturer’s instructions. For larger yields, bacterial plasmid DNA was isolated using a modification of the DNA miniprep method used by Serghini *et al.* (1989). A single *E.coli* colony was inoculated into 5mL LB and incubated for 16 h overnight, after which 1.5mL samples of culture were centrifuged at 13 000 rpm in 1.5mL Eppendorf tubes for 45 s. After centrifugation the supernatant was discarded and 50μl of a TEN buffer (10mM Tris-HCl, pH 8.0, 1mM EDTA, 100mM NaCl) was added to each sample. Cells were resuspended by vortexing for 2 min and 50μl of a phenol/chloroform/isoamyl alcohol mixture (25:24:1 v/v) was added. Following a brief mix by vortexing, samples were centrifuged at 13 000 rpm for 5 min, and the supernatant transferred to a new 1.5mL Eppendorf tube containing 100μl isopropanol.
After the transfer, 17µl of 7.5M ammonium acetate (NH₄OAc) was added to samples, which were briefly mixed before being centrifuged at 13 000 rpm for 1 minute. After centrifugation the supernatant was discarded and the pellet rinsed with 70% (v/v) EtOH. Samples were aspirated and the DNA resuspended in 40µl H₂O.

**DNA amplification**

DNA was amplified using Supertherm DNA Polymerase (Separations Scientific SA Pty Ltd., Honeydew, South Africa) in a polymerase chain reaction (PCR). Table 1 describes the PCR primers used in experiments, their sequence information, and annealing temperatures (Tₘ). Each reaction had a total volume of 20µl and included 1.5mM MgCl₂, 1 x PCR Buffer, 0.4µM of each primer, 0.2mM dNTPs (Fermentas, Ontario, Canada), 0.5U Supertherm DNA Taq polymerase. Typically between 100 – 200ng template DNA was used in each reaction. The following cycling conditions were used for all reactions: Initial denaturation at 94°C for 5 min followed by 35 cycles (94°C for 15 s, Tₘ for 30 s, and 72°C for 15 s/kilobase of product) and a final elongation at 72°C for 10 min. All PCR reactions were carried out in a Gene Amp PCR System 2700 (Applied Biosystems, Foster City, USA).

**High fidelity PCR**

When cloning, a high fidelity Velocity DNA polymerase (Bioline Ltd., London, UK) was used in PCR reactions to reduce the polymerase error rate. Reactions were prepared according to the manufacturer’s protocol using the following cycling conditions: Initial denaturation at 98°C for 2 min followed by 35 cycles (98°C for 30 s, Tₘ for 30 s, and 72°C for 30 s/kilobase of product) and a final elongation at 72°C for 10 min.
Table 1 | PCR primers used during the course of this project.

<table>
<thead>
<tr>
<th>Primer name</th>
<th>Sequence</th>
<th>Annealing Temp (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cloning</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AT4G11100 – F</td>
<td>AGGATCCATGATGTTTGGTTAGAAACG</td>
<td>55</td>
</tr>
<tr>
<td>AT4G11100 – R</td>
<td>AGAATTCAGAGGCTTTTGGGAATCTTCC</td>
<td>55</td>
</tr>
<tr>
<td>AT4G11100stop – R</td>
<td>AGAATTCCTAAGGCTTTGGGAATCC</td>
<td>55</td>
</tr>
<tr>
<td><strong>Genotyping</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>At-LP</td>
<td>AAACGATGACTTTGGAGCATG</td>
<td>55</td>
</tr>
<tr>
<td>At-RP</td>
<td>AACTCCTGACAAAAACAGAAAGC</td>
<td>55</td>
</tr>
<tr>
<td>LBb1.3</td>
<td>ATTTTGCCGATTTTCGGAAC</td>
<td>55</td>
</tr>
<tr>
<td><strong>Sequencing</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GFP fusion</td>
<td>AGATGAACTTCAGGGTCAG</td>
<td>50</td>
</tr>
</tbody>
</table>

**DNA electrophoresis**

Electrophoresis of DNA was performed using a 1% (w/v) agarose gel containing 0.016µg/mL ethidium bromide (EtBr). Gels were prepared using 1 x TAE buffer (40mM Tris, 1mM EDTA, 0.11% (v/v) glacial acetic acid). DNA was separated alongside a 1 Kb size marker (O’ Gene Ruler ™ DNA ladder, Fermentas, Ontario, Canada). All gels were electrophoresed in 1 x TAE buffer for approximately 1 h, after which DNA was visualised with a long wavelength (365 nm) UV transilluminator.
**Gel extraction of DNA**

DNA was purified from the gel slice using the Wizard SV Gel and PCR Clean up system (Promega Corporation, Madison, USA) according to the manufacturer's protocol.

**Restriction endonuclease digestion of DNA**

Fermentas International Inc. (Ontario, Canada) restriction enzymes (REs) were used in the digestion of plasmid or purified PCR DNA. Reactions typically required between 5 - 10 Units of RE, and were carried out according to the manufacturers recommended conditions.

**Ligation of DNA**

Ligation reactions consisted of purified insert and vector DNA, 3 – 5 Units of T4 DNA Ligase (Fermentas International Inc., Ontario, Canada), and 1 x T4 DNA Ligase buffer. A vector:insert ratio of 1:3 was used in all ligation reactions, with the amount of insert required (ng) calculated using the following equation:

\[
\text{amount of insert (ng)} = \frac{\text{quantity of vector DNA (ng)} \times \text{size of insert DNA (Kb)}}{\text{size of vector DNA (Kb)}} \times \text{insert:vector molar ratio}
\]

Reactions were performed according to the manufacturer’s protocol and incubated overnight at 4 °C.

**LR Gateway cloning**

Gateway cloning utilizes `attL`- and `attR`- sites to facilitate the transfer of a desired fragment of DNA from an entry vector to a destination vector. The transfer of a DNA fragment from
between the attL- sites of the entry vector to between the attR- sites of the destination vector is performed through LR recombination. LR recombination was performed according to the manufacturer’s protocol.

**Entry Vector**

*pENTR™4 Dual selection vector*

This vector contains a *ccdB/chloramphenicol* fusion gene between the attL- and attR- sites which allows for negative selection of DH5α *E.coli* cells that are sensitive to the ccdB protein. The empty vector was propagated in One Shot® *ccdB Survival™2 T1R* *E.coli* cells that are ccdB insensitive, in the presence of 50 µg/mL kanamycin, which was also used to select for positive DH5α transformants.

**Destination Vectors**

*pFAST-G02*

This vector contains a *ccdB* gene between the attL- and attR- sites which allows for negative selection of DH5α *E.coli* cells that are sensitive to the ccdB protein. The empty vector was propagated in One Shot® *ccdB Survival™2 T1R* *E.coli* cells that are ccdB insensitive, using either 50 µg/mL spectinomycin or 50 µg/mL streptomycin to select for positive transformants. pFAST-G02 contains the herbicide resistance gene *Bar*, which allows for the identification of transformed plants via phosphinothricin selection. Alternatively, transformed plant seeds can be identified under a fluorescent microscope as they express enhanced green fluorescent protein (EGFP).
pK7FWG2.0

This vector contains a *ccdB/chloramphenicol* fusion gene between the *attL*- and *attR*- sites which allows for negative selection of DH5α *E.coli* cells that are sensitive to the *ccdB* protein. The empty vector was propagated in One Shot® *ccdB* Survival™2 T1R *E.coli* cells that are *ccdB* insensitive, using 50 µg/mL spectinomycin, 50 µg/mL streptomycin, or 50 µg/mL kanamycin to select for positive transformants. pK7FWG2.0 contains an *EGFP* gene immediately following the *attR2* site, which is designed to translationally fuse with genes transferred through LR recombination.

**E. coli competent cell preparation**

A single *E.coli* colony was inoculated into 5mL LB media and grown for 16 h overnight. Two mL of the overnight culture were added to 250mL LB media containing 20mM MgSO₄, which was left to grow at 37°C while shaking until the OD₆₀₀ was between 0.4 and 0.6. The culture was then centrifuged in a JA-21 Beckman centrifuge at 5000 x g for 5 min at 4°C and the supernatant discarded. Cells were resuspended in 100mL ice cold TFB1 (a filter sterilized buffer containing 30mM KAc, 100mM RbCl, 10mM CaCl₂, 50mM MnCl₂, 15% glycerol (v/v), adjusted to pH 5.8 using glacial acetic acid) and kept on ice for 5 min. The suspension again centrifuged at 5000 x g for 5 min at 4°C and the supernatant discarded. The pellet was resuspended in 10mL ice cold TFB2 (a filter sterilized buffer containing 10mM MOPS, 75mM CaCl₂, 10mM RbCl and 15% glycerol (v/v)) and kept on ice for between 15 and 60 min. Competent cells were divided into pre-cooled 1.5mL Eppendorf tubes in 100µL aliquots, and frozen immediately in liquid nitrogen. Cells were stored at -80°C.
Transformation of competent *E.coli* cells

Between 50 and 100ng of plasmid DNA were added to 100 µl ice-thawed competent *E. coli* cells. After the addition of the DNA, cells were left on ice for 30 min, followed by a heat shock at 42°C for 45 s and then snap cooled on ice for 2 min. Nine hundred µL of LB media was added and cells were incubated at 37°C for 60 min with shaking. Following incubation, 100µl of each culture was plated on LB plates (1.5 % w/v agar) with the appropriate antibiotics before being incubated at 37°C for 16 h overnight.

Glycerol stocks

Glycerol stocks of positive transformed cells lines containing a vector of interest were created for storage. Glycerol stocks were created by adding 680µL of freshly grown overnight culture to 320µL of pre-sterilized 50 % glycerol in a 1.5 mL Eppendorf. Stocks were flash frozen in liquid nitrogen and stored at -80°C.

DNA sequencing and analysis

All DNA sequencing was performed at the Central Analytical Facility (Stellenbosch, South Africa) on an ABI3730xl DNA analyser (Applied Biosystems, Foster City, USA). Sequence data was analysed using Chromas software (Version 2.01, Technelysium Pty Ltd, Queensland, Australia) and BLAST (http://blast.ncbi.nlm.nih.gov/Blast.cgi).
RNA Manipulation

Isolation of RNA from *A. thaliana*

Total RNA was extracted using a TRizol reagent (100mM NaAc pH 5.2, 800mM guanidine thiocyanate, 400mM ammonium thiocyanate, 5% glycerol (v/v) and 38% phenol (v/v) pH 4, made up in DEPC-treated dH2O) based on the Chomczynski & Mackey (1995) RNA extraction protocol. RNA was extracted according to the TRizol reagent protocol (Invitrogen, Carlsbad, USA) with the following modification: Plant tissue was homogenised in 1.5mL Eppendorf tubes each containing 3 stainless steel ball bearings and 1mL TRizol reagent. Tubes were subjected to 4 min of mechanical shaking in a paint shaker.

Electrophoresis of RNA

Electrophoresis of RNA was performed using a RNA formaldehyde-agarose denaturing gel, containing 1 x MOPS pH 7 (0.4M MOPS, 0.1M NaAc, 10mM EDTA), 1.2% (w/v) agarose, and 2.25% (v/v) formaldehyde. Prior to electrophoresis, RNA samples were mixed with 0.2 volumes of RNA sample application buffer (4 x MOPS, 2.7% (v/v) formaldehyde, 30.8% (v/v) formamide and 0.01 mg/mL EtBr), heated at 65°C for 5 min and snap cooled on ice. RNA was visualised on a Gel Doc™ XR UV transilluminator (Bio-Rad Laboratories, UK).

DNase treatment of RNA

RNA to be used in cDNA synthesis was first treated with DNase from the Turbo DNA-free™ kit (Life Technologies, California, USA) as instructed by the manufacturer.
cDNA synthesis

cDNA synthesis was performed using Superscript III Reverse Transcriptase (Life Technologies, California, USA) according to the manufacturer’s protocol with the following modifications: 2.5µg total RNA and half volume of the Superscript™ III enzyme were used; cDNA synthesis was performed at 42°C for 2 h, followed by a heat inactivation step at 72°C for 15 min.

Quantitative real-time PCR (qPCR)

All qPCR reactions were performed using the KAPA SYBR® FAST qPCR Kit, according to the manufacturer’s protocol. A Rotor-Gene® 6000 Real-Time PCR machine (QIAGEN, Limburg, Netherlands) was used to amplify cDNA under the following cycling conditions: 95°C for 3 min, followed by 40 cycles (95°C for 3 s, 60°C for 20 s, and 72°C for 1 s), and a final step at 72-95°C for 5 min. Rotor-Gene® Series Software (version 1.7) was used for melt curve analysis and quality control (with a minimum $R^2$ value of 0.99, slope values between 3.3 and 3.5 required for a run to be considered “good”; slope values near 3.3 reflect a reaction efficiency of near 100%). Table 2 details all primers used for qPCR experiments in this project, their sequences, and the concentrations at which they were effective.
Table 2 | qPCR primers used during the course of this project.

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Locus</th>
<th>Primer pair sequences</th>
<th>Final concentration (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACT2</td>
<td>AT3G18780</td>
<td>AGTGGTCGTACAACCGGTATTGT CATGAGGTAATCAGTAAGGTCACGT</td>
<td>900 300</td>
</tr>
<tr>
<td>SNC1</td>
<td>AT4G16890</td>
<td>GCTCGCCGACTTTACAGACT GGAAGATGATAACAATTTATCCAGA</td>
<td>200 200</td>
</tr>
<tr>
<td>AT4G11100</td>
<td>AT4G11100</td>
<td>GTAGTAGCTTCGTTACTCAAAGT GCATCTGTCACAGCATGTTTC</td>
<td>200 200</td>
</tr>
<tr>
<td>PR2</td>
<td>AT3G57260</td>
<td>TCTTCAACCACACAGCTGGA TCTGAACCTGGGAACGTCGAG</td>
<td>200 200</td>
</tr>
<tr>
<td>ICS1</td>
<td>AT1G74710</td>
<td>GCTAGCAGTGTTACACGCGTG AAGCTTCACTGCAGACACCT</td>
<td>200 200</td>
</tr>
</tbody>
</table>

Western Blots

Crude protein extraction from A. thaliana

A. thaliana leaf tissue was homogenised in an extraction buffer (10mM Potassium Phosphate pH7.2, 5mM DTT) and centrifuged for 5 min at 10 000 rpm. Protein concentrations of extracts were determined though a Bradford assay.

SDS Polyacrylamide Gel preparation

A 15% polyacrylamide gel was prepared and electrophoresed using a Mini-PROREAN® 3 system (BioRad Laboratories, Inc. Hercules, USA). In all gels a 40% (w/v) Acrylogel acrylamide/bisacrylamide (29:1 v/v) stock solution (Sigma-Aldrich Inc., St Louis, USA) was utilized. A resolving gel (20% (v/v) Acrylogel, 375mM Tris-HCl ph8.8, 0.2% (w/v) SDS, 0.1%
(w/v) APS, 0.1% (v/v) TEMED) was overlaid by a stacking gel (4% (v/v) Acrylogel, 125mM Tris-HCl pH6.8, 0.1% (w/v) SDS, 0.09% (w/v) APS, 0.1% (v/v) TEMED).

### SDS Polyacrylamide Gel Electrophoresis (SDS-PAGE)

Heat denatured crude protein samples, together with 1x loading buffer, and a molecular weight marker (PageRuler™ Prestained Protein Ladder; Fermentas International Inc., Ontario, Canada) were loaded onto the polyacrylamide gel. Samples were electrophoresed in a running buffer (25mM Tris-HCl pH8.8, 192mM glycine and 0.1% (w/v) SDS) for 30 min at 60 volts, as they moved through the stacking gel. The voltage was then increased to 200 volts until the dye front had run off the bottom of the resolving gel.

### Protein transfer

Protein samples were transferred from the polyacrylamide gel onto a nitrocellulose membrane (Schleicher and Schuell BioScience, Dassal, Germany) for 1 hour at 15 volts, using the Biorad semi-dry transblotter apparatus (BioRad Laboratories, Inc. Hercules, USA) as per manufacturer’s instructions. The membrane and blotting paper were pre-soaked in transfer buffer (25mM Tris, 192mM glycine and 20% (v/v) methanol).

### Blocking and eGFP incubation conditions

Nitrocellulose membranes with transferred protein were blocked with blocking buffer (1 x PBS(58mM Na₂HPO₄.2H₂O, 18mM KH₂PO₄, 1.37M NaCl, 26mM KCl), 5% milk powder (w/v), 0.1% (v/v) Tween) for 30 min at 22°C with shaking. Roche mouse anti-GFP antibodies were used to detect the presence of eGFP in protein samples.
**A. thaliana transformation**

**Plant preparation**

Soil grown plants were allowed to grow for 4 weeks until the first inflorescence shoot began to appear. These inflorescence shoots were removed as this encourages growth of multiple secondary inflorescences. The next inflorescence shoots were again removed, after which the plants were allowed to grow until a large number of unopened flower heads appeared.

**A. tumefaciens competent cell preparation**

A 10 mL LB medium, supplemented with the appropriate antibiotics, was inoculated with a single colony of *A. tumefaciens* and incubated at 28°C for 16 h overnight with shaking. 2 mL of this culture was transferred to 50 mL fresh LB media (with appropriate antibiotics) and incubated with shaking at 28°C until the OD600 reached 0.5–1.0. The culture was chilled on ice before the cells were harvested by centrifugation at 3000 x g for 5 min at 4°C in a J2-21 Beckman centrifuge (Beckman Coulter, Inc., CA, USA). Pelleted cells were resuspended in 1 mL ice cold 20 mM CaCl2. Aliquots of 100μl were then immediately frozen in liquid nitrogen and stored at -80°C.

**Transformation of competent A. tumefaciens**

The 100μl competent *A. tumefaciens* cell aliquots were defrosted on ice for approximately 5 min. After the cells had defrosted, between 50 and 100ng of plasmid DNA was added to the Eppendorf tube. Following the addition of the DNA, cells were heat shocked at 37°C for 5 min. Nine hundred μL of LB media were added and cells were incubated at 30°C for 6 h while shaking. Following incubation, 100μl of each culture was plated on LB plates (1.5 %
w/v agar) with the appropriate antibiotics before being incubated at 30°C for between 2 and 3 days.

**Floral dip transformation of *A. thaliana***

*A. thaliana* plants were transformed following a protocol based on the floral dip method used by Clough and Bent (1998). Any open flower heads were removed before submerging the aerial parts of plants in an *A. tumefaciens* suspension for 5 s. Dipped plants were left on their sides in a paper towel-lined tray which was then covered with clingfilm and left for 16 h overnight at 22°C. Plants were then placed upright in a tray and allowed to grow normally until siliques were fully formed, after which the plants were allowed to dry for seed collection.

**Herbicide selection of plants containing pFAST-G02**

Transgenic plants expressing the *Bar* gene are resistant to herbicides containing glufosinate ammonium (phosphinothricin), which can be used to select for *Bar* containing plants (homozygous plants will display 100% resistance). After 5 days of growth, soil grown transgenic plants were sprayed with a 0.015% (w/v) Basta solution (Basta is a herbicide containing glufosinate ammonium) so that all leaf surfaces were damp. Two more rounds of herbicide spraying took place at 8 and 11 days of growth, after which resistant plants could be distinguished by healthy green leaves, and sensitive plants by pale yellow leaves.
**A. thaliana protoplast isolation and transfection**

*A. thaliana* protoplasts were isolated and transfected with plasmid DNA according to the method used by Yoo *et al.* (2007). A Zeiss LSM 510 Meta confocal microscope was used for imaging transfected protoplasts, utilizing a LD C-Apochromat 403/1.1 W M27 objective. Excitation of GFP was performed at 488 nm and emission detected in the 500 to 520 nm range.

**Luciferase assays**

Luciferase activity was detected using the Luciferase Assay System (Promega Corporation, Madison, US) according to the manufacturer’s instructions, utilizing a Modulus microplate luminometer.

**Data analysis**

All data analysis was performed using StatSoft Statistica software. Fisher LSD post hoc tests were used in the Analysis of Variance (ANOVA) tests of differences between groups.
Chapter 3: Results and Discussion

The cir1 growth and disease resistance phenotypes are temperature dependent

Environmental temperature modulates both growth and disease resistance in wild type plants (Garrett et al., 2006); with higher growth temperatures often associated with compromised immunity and greater plant growth. Similarly, in a number of gain-of-resistance mutants, including constitutive expressor of PR genes 1 (cpr1), suppressor of rps4-RLD 1 (srfr1), bonzai1 (bon1), and suppressor of npr1-1, constitutive 1 (snc-1), enhanced resistance to pathogen attack and constitutive expression of defence genes are abolished when these plants are grown at higher temperatures (Zhang et al., 2003; Gou et al., 2009; Kim et al., 2010). The constitutively induced resistance 1 (cir1) mutant is also a constitutive expresser of a number of plant defence genes, including PATHOGENESIS RELATED-1 (PR-1), and displays enhanced resistance to infection by virulent bacterial pathogens including Pseudomonas syringae (Murray et al., 2002), but whether these phenotypes are similarly modulated by environmental temperature is unknown.

The effects of temperature on the growth of the cir1 mutant were investigated by growing cir1 and PR-1:LUC (cir1 genetic background) plants at 18°C, 22°C (standard growth temperature), and 25°C for 4 weeks (Figure 3). Temperature clearly influenced plant growth in both lines, with much smaller plants observed at 18°C and progressively larger individuals at 22°C and 25°C respectively. A marked difference in the size of cir1 and PR-1:LUC plants was observed at 18°C, with cir1 individuals growing considerably smaller than PR-1:LUC individuals. This size difference was less obvious at 22°C, and was completely absent at 25°C.
Figure 3 | Temperature sensitive growth phenotype of \textit{cir1}. The \textit{cir1} mutants displays reduced growth at 18°C as compared to \textit{PR-1:LUC} control plants. Plants were grown for four weeks under a 16 hour light/8 hour dark cycle at 18°C, 22°C or 25°C. Scale bar represents 10mm.

Thus, like other gain-of-resistance mutants, \textit{cir1} displays a temperature dependent growth phenotype. However, \textit{cpr1}, \textit{srfr1}, \textit{bon1}, and \textit{snc-1} all display a temperature modulated growth phenotype where plants are dwarfed at temperatures between 22°C and 28°C (Zhang \textit{et al}., 2003; Gou \textit{et al}., 2009; Kim \textit{et al}., 2010). The \textit{cir1} mutant clearly differs from these mutants as it only exhibits reduced growth at 18°C, which is completely abolished at 22°C.

To determine whether the enhanced resistance to infection by \textit{P. syringae} displayed by \textit{cir1} under normal growth conditions (22°C), is also modulated by temperature, 4 week old \textit{cir1} and \textit{PR-1:LUC} plants grown at 18°C, 22°C, and 25°C were infected with a virulent \textit{P. syringae} pv. \textit{tomato} DC3000 (\textit{Pst DC3000}) strain, and bacterial growth quantified 48 h post infection
The cir1 mutant shows significantly lower bacterial titres when compared to PR-1:LUC at both 18°C (p < 0.05) and 22°C (p < 0.05), but not at 25°C. Interestingly, the fold difference in bacterial titre 48 h post infection between cir1 and PR-1:LUC is greater at 18°C (29 fold) than at 22°C (7 fold) and bacterial titres in the cir1 mutant are significantly lower at 18°C versus 22°C (Figure 4). This demonstrates that there is an increased defence response in cir1 at 18°C that gradually lessens as growth temperature increases, and is abolished at 25°C. Taken together, the reduced growth and increased resistance to Pst DC3000 of cir1 at 18°C suggests that these are temperature dependent characteristics (i.e. enhanced resistance is lost at higher temperatures) and that there is likely an energetic cost associated with the constitutive expression of defence genes in cir1 which results in reduced biomass production at 18°C.
Figure 4 | Temperature sensitive pathogen resistance phenotype of cir1. Mean P. syringae titre (+SE, n = 5) 48 h post infection in cir1 and PR-1:LUC control lines. Plants were grown for four weeks at 18°C, 22°C and 25°C prior to infection with Pst DC3000. Letters indicate significant differences (p < 0.05) determined by ANOVA with Fisher LSD. This experiment is representative of three independent experiments.

The temperature conditional phenotype of cir1 could explain the “variable nature of the cir1 mutant” described in previous work (Carstens, 2008; Diener, 2012). This variability refers to several instances where the bacterial titres in cir1 plants did not significantly differ from those of control lines during pathogen assays performed on plants grown in the departmental plant growth room, which is subject to temperature fluctuations several degrees above 22. In contrast, the experiments described above were carried out in Perceval growth chambers where the temperature is more reliably kept constant through a computer controlled temperature regulator. Given what is now known about the
temperature sensitivity of the cir1 mutant, the instances where cir1 displayed a “variable nature” may well have been the result of fluctuating temperatures during growth leading to the repression of the cir1 phenotype.

The nucleotide-binding and leucine-rich repeat (NB-LRR) R protein, SNC1, is essential for the dwarfed growth and constitutive defence gene expression in the gain-of-resistance mutants bon1, cpr1, and srfr1, which display elevated SNC1 RNA and/or protein levels (Yang & Hua, 2004; Gou et al., 2009; Kim et al., 2010). It is possible SNC1 acts as a temperature sensor that modulates plant immunity in response to changes in environmental temperature, and may do this once a threshold amount of SNC1 has localised in the nucleus (Zhu, Qian & Hua, 2010). To investigate whether SNC1 might be similarly important to the growth and resistance phenotypes in cir1, SNC1 expression was analysed in cir1 and PR-1:LUC plants grown at 18°C and 22°C (Figure 5). However, quantitative real-time PCR (qPCR) analysis of SNC1 mRNA from 4 week old plants revealed no significant differences in SNC1 transcript levels between cir1 and PR-1:LUC, at either 18°C or 22°C, suggesting that the cir1 phenotypes may be independent of SNC1 mRNA levels. In order to definitively determine whether or not SNC1 is required for the defence phenotype of cir1, a cross between the cir1 mutant and a snc1 knockout mutant would need to be performed. If the offspring of such a cross continued to display enhanced disease resistance, this would indicate that the cir1 defence phenotype is indeed SNC1 independent.
Identification and characterisation of a second *at4g11100* T-DNA insertion mutant

The *cir1* mutant was originally identified and characterised by Murray and colleagues (2002); subsequent work has involved attempts to identify the gene responsible for the *cir1* phenotype. Through a series of genetic mapping experiments, the location of the *CIR1* gene was narrowed down to a region of chromosome IV, where 8 possible gene candidates are present (Carstens, 2008). Homozygous T-DNA insertion mutants for each of the 8 candidate genes were obtained and analysed in order to determine if any phenocopied *cir1*. Unlike the other seven mutants, the *at4g11100-p* mutant (SALK_062847 - containing a T-DNA insertion in the promoter region of the *AT4G11100* gene) displayed elevated levels of PR-1 protein.
and was found to be more resistant to infection by Pst DC3000 as compared to wild type A. thaliana; for this reason AT4G11100 was considered to be the most likely candidate for the CIR1 gene (Diener, 2012).

In the present study, to ensure that the phenotype of the at4g11100-p mutant was indeed due to a disruption of that gene and not an artefact of multiple T-DNA insertions in this plant line, a second at4g11100 T-DNA insertion mutant was obtained - at4g11100-e (SALK_096586 - containing a T-DNA insertion in exon 2). However, a homozygous at4g11100-e line was unavailable and so it was necessary to screen a segregating line to identify homozygous individuals. Screening was undertaken using a PCR based method that utilized either AT4G11100 gene-specific primers, or T-DNA insert-specific primers (Figure 6A). The AT4G11100 gene-specific primers (At-LP and At-RP) were designed in positions flanking the predicted site of T-DNA insertion, and because the insert is almost 5kb in size, amplification using the gene-specific primers is impossible in the presence of the insert; similarly, the T-DNA insert-specific primers (At-RP and LBb 1.3) generate a PCR product only in the presence of the insert. Therefore, using these two primer pairs it is possible to determine if an individual plant is homozygous for either the wild type or T-DNA insertion at4g11100 alleles, or if an individual is a heterozygote containing both alleles (Figure 6B). Once homozygous at4g11100-e plants were identified, they were separated from the others and allowed to set seed; homozygous at4g11100-e seed stocks were generated in this way.
Figure 6 | \textit{at4g11100-e} T-DNA insert position and identification of a homozygous T-DNA insertion line. A) The At-LP and At-RP gene-specific primers are designed to amplify wild type \textit{AT4G11100} and flank the predicted site of T-DNA insertion. The LBb1.3 insert-specific primer is located on the border of the T-DNA insert and generates a product with At-RP only in the presence of a T-DNA insertion. Exons are indicated by boxes, while primers and the direction of their amplification are indicated by arrows. B) PCR amplification of genomic DNA extracted from a wild type Columbia (Col-0) plant and three plants (a, b, and c) from the \textit{at4g11100-e} T-DNA insertion line. DNA was amplified using either T-DNA insertion specific primers (LBb1.3 + At-RP) or wild type \textit{AT4G11100} primers (At-LP + At-RP). The red box highlights the lack of amplification of Col-0 DNA when using T-DNA insertion specific primers. The yellow boxes highlight the lack of amplification of DNA from plants a and b using wild type \textit{AT4G11100} primers. Plants a and b are homozygous for the T-DNA insertion. Plant c is heterozygous for the insertion.
Additionally, the PCR product obtained with the LBb1.3 and At-RP primers was sequenced to determine the exact site of T-DNA insertion. The predicted site of T-DNA insertion in \textit{at4g11100-e} according to \textit{The Arabidopsis} Information Resource (TAIR) is 814bp downstream of the ATG start codon, in exon 2; however, sequencing data revealed that the actual site of insertion is approximately 250bp further upstream, at 563bp downstream of the ATG start codon, in the middle of an intron (Figure 7).

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{predicted_actual_tdna_insertion.png}
\caption{\textbf{Predicted and actual position of T-DNA insertion in \textit{at4g11100-e}.} The predicted site of T-DNA insertion in \textit{at4g11100-e}, according to \textit{The Arabidopsis} Information Resource (TAIR – arabidopsis.org), is 814bp downstream of the ATG start codon. Sequencing data shows the actual site of T-DNA insertion in \textit{at4g11100-e} is 563bp downstream of the ATG start codon.}
\end{figure}
It is also necessary to determine that T-DNA insertion lines are true knock-outs for the gene of interest, especially given the fact that the T-DNA insert in this line is in an intron. To this end, reverse transcriptase PCR (RT-PCR) was used to ensure that no full-length \textit{AT4G11100} mRNA was present in the \textit{at4g11100-e} plants. cDNA generated from \textit{at4g11100-e} and Col-0 total RNA was subjected to PCR using \textit{AT4G11100} primers specific to the 5' and 3' ends of the cDNA sequence (Figure 8). Amplification was only observed from the Col-0 cDNA, indicating that the T-DNA insert was sufficient to prevent the transcription of wild-type \textit{AT4G11100} mRNA.

![Figure 8](image)

Figure 8 | Reverse transcriptase PCR (RT-PCR) of SALK_096586 plants homozygous for T-DNA insert. PCR amplification of cDNA obtained from wild type Columbia (lane 1) and two plants from the homozygous T-DNA insertion line (lanes 2 and 3) using \textit{AT4G11100} specific primers (At-LP & At-RP). The yellow boxes highlight the lack of amplification of cDNA from the two plants homozygous for the T-DNA insertion.
The identification of the \textit{at4g11100-e} mutant was followed by a characterisation of its growth and resistance phenotypes. Considering the temperature dependent nature of \textit{cir1} growth, the effects of temperature on the growth of the \textit{at4g11100-p} and \textit{at4g11100-e} mutants were investigated by growing both \textit{at4g11100} mutants and Col-0 (\textit{at4g11100-e} genetic background) at 18\textdegree{}C, 22\textdegree{}C, and 25\textdegree{}C for 4 weeks. However, unlike \textit{cir1}, neither \textit{at4g11100} mutant displayed any reduction in growth at 18\textdegree{}C or 22\textdegree{}C when compared to Col-0 (data not shown). Next, 4 week old \textit{at4g11100-p}, \textit{at4g11100-e}, and Col-0 plants were infected with \textit{Pst} DC3000, and bacterial growth was quantified 4 h and 48 h post infection (Figure 9). Both \textit{at4g11100} mutants had significantly lower bacterial titres (p < 0.05) 48 h post infection than infected Col-0, with an approximate 10 fold reduction in bacterial titre; there was no significant difference in bacterial titre between the \textit{at4g11100} mutants. Therefore it appears that although neither \textit{at4g11100} mutant exhibits the temperature sensitive growth phenotype of \textit{cir1}, both do show enhanced resistance to \textit{Pst} DC3000 infection; and importantly, the \textit{at4g11100-e} mutant displays the same phenotypes as \textit{at4g11100-p}, indicating that this is likely due to disruption of \textit{At4g11100}.
**Heterologous expression of AT4G11100 does not complement the cir1 phenotype**

If the cir1 mutation is in AT4G11100 then the introduction of a functional AT4G11100 gene to the mutant should abolish the cir1 phenotypes, including constitutive PR-1 expression. To test this, the wild type AT4G11100 open reading frame (ORF) was amplified from Col-0 A. thaliana cDNA and cloned into the pENTR™4 entry vector. The primers used to amplify and clone AT4G11100 (Table 1) contained EcoRI and BamHI restriction enzyme sites, and so to confirm the presence of AT4G11100 in pENTR™4, a double digest was performed (Figure 9).

**Figure 9** | *at4g11100* T-DNA insertion mutants display increased resistance to *P. syringae* infection. Mean *P. syringae* titre (+SE, n = 5) in wild type Columbia (Col-0) and two AT4G11100 knockout lines 4 and 48 h post infection. Plants were grown for four weeks at 22°C prior to infection. Letters indicate significant differences (p < 0.05) determined by ANOVA with Fisher LSD. This experiment is representative of three independent experiments.
10A). Once the presence of AT4G11100 in pENTR™4 had been confirmed, the AT4G11100 coding region was sequenced to ensure that there were no mutations present in the gene, and the AT4G11100 insert was transferred to the pFAST-G02 vector (Shimada, Shimada & Hara-Nishimura, 2010) through Gateway LR recombination. E. coli DH5α cells were then transformed with the resulting pFAST-G02::AT4G11100 construct, and the presence of AT4G11100 was again confirmed through a PCR (Figure 10B).
Figure 10 | Confirmation of AT4G11100 in pENTR™4 and pFAST-G02.
A) Double digest of the pENTR4::AT4G11100 vector with EcoRI and BamHI restriction enzymes. Bands correspond with the 864bp AT4G11100 ORF and the 2400bp pENTR™4 vector backbone. B) Lanes 6 and 7 show PCR amplification of AT4G11100 from the pFAST-G02::AT4G11100 construct (using At-LP & At-RP primers). Bands correspond to the 864bp AT4G11100 sequence. Lanes 1 – 5 show amplifications from non-related cloning experiments.
A GV3101 *Agrobacterium tumefaciens* strain was transformed with the *pFAST-G02::AT4G11100* construct, and the *pFAST-G02* empty vector. These *A. tumefaciens* strains were in turn used to transform 6 week old *cir1* and *PR-1:LUC* *A. thaliana* plants. The presence of enhanced green fluorescent protein (eGFP) in the *pFAST-G02* vector meant that transgenic seed from the transformed plants could be identified using a fluorescent microscope, where they fluoresced brightly after excitation at 395nm.

The *cir1* mutant was generated in a *PR-1:LUC* genetic background which contains a *PR-1* promoter transcriptionally fused to a luciferase reporter gene; this means that *PR-1* promoter activity can be inferred by the measurement of luciferase activity. The *cir1* mutant displays elevated levels of *PR-1* transcription, and therefore elevated luciferase activity when compared to its *PR-1:LUC* genetic background. With this in mind, the transgenic *cir1* and *PR-1:LUC* seeds were grown for two weeks at 22°C before being assayed for luciferase activity, to determine if the presence of a functional *AT4G11100* gene would have any effect on *PR-1* promoter activity (Figure 11).
Luciferase activity in cir1 and PR1:LUC plants transformed with AT4G11100. Mean luciferase activity from tissue of 2 week old plants was assayed in four plant lines: the cir1 mutant (+SE, n = 3), cir1 transformed with the pFAST-G02::AT4G11100 construct (+SE, n = 2), PR-1:LUC transformed with the pFAST-G02 empty vector (+SE, n = 4), and PR-1:LUC transformed with the pFAST-G02::AT4G11100 construct (+SE, n = 4). In vector-containing plant lines, biological replicates represent independent transformation events. Luciferase activity was normalised for total protein content using a Bradford assay. All plants were grown at 22°C.

Figure 11 clearly shows the elevated luciferase activity in cir1 versus PR-1:LUC plants, as previously reported (Diener, 2012). However, the presence of AT4G11100 has no effect on the luciferase activity of the cir1 mutant, with no significant difference between luciferase activity in cir1 and cir1+AT4G11100 plants, suggesting that AT4G11100 may not be the gene responsible for the cir1 phenotype.
CIR1 and AT4G11100 complementation - AT4G11100 is not CIR1

A second experiment to determine whether the AT4G11100 gene is in fact CIR1 was performed; here, complementation was tested by cross pollinating the at4g11100-e and cir1 mutants. Because the cir1 mutation is recessive, two copies of the mutant allele are necessary to elicit the cir1 phenotypes (i.e. constitutive expression of PR-1, and enhanced disease resistance); thus, in a genetic cross between at4g11100-e and cir1, if the mutation responsible for the cir1 phenotype is present in the AT4G11100 gene, then there will be constitutive expression of PR-1 and high levels of luciferase activity. Conversely, if the mutation is in a different gene, no elevated expression of PR-1 should occur and luciferase activity will be low (i.e. complementation has occurred).

Seeds from two at4g11100-e x cir1 crosses were soil grown for two weeks at 22°C before being assayed for luciferase activity. As before, the elevated PR-1 transcription in cir1 plants, and the low basal levels of PR-1 transcription in PR-1:LUC plants, is reflected by the high and low levels of luciferase activity in cir1 and PR-1:LUC respectively (Figure 12). Offspring from both at4g11100-e x cir1 crosses displayed significantly lower luciferase activity when compared to those of cir1 individuals; instead, offspring from both crosses were not significantly different from PR-1:LUC plants in luciferase activity, indicating that complementation had occurred. Thus, it seems likely that CIR1 is not AT4G11100. Finally, DNA sequencing of the AT4G11100 locus (including 1.5 kb of upstream sequence) in cir1 showed no nucleotide differences compared to the wild type Col-0 sequence.
Figure 12 | Luciferase activity in the offspring of a cross between the cir1 and at4g11100-e mutants. Mean luciferase activity (+SE, n = 3) of tissue from 2 week old plants was assayed for luciferase activity in the cir1 x at4g11100 offspring, PR-1:LUC and cir1. Luciferase activity was normalised for total protein content using a Bradford assay. All plants were grown at 22°C. Letters indicate significant differences (p < 0.05) determined by ANOVA with Fisher LSD. This experiment is representative of two independent experiments.

Generation and characterisation of AT4G11100 over expressers

Although AT4G11100 is not CIR1, the at4g11100 insertion mutants nonetheless exhibit increased resistance to Pst DC3000, suggesting that this gene may be involved in the negative regulation of innate immunity. Therefore, characterising the AT4G11100 gene could yield valuable insight into the mechanisms underlying immunity in A. thaliana.

Because, like CIR1, AT4G11100 is most likely a negative regulator of the immune response, the effect of over-expressing AT4G11100 is an interesting avenue for investigation. The rationale behind creating AT4G11100 over-expressing mutants was based on the hypothesis that over-expressing a negative regulator of plant immunity would inhibit a plant’s ability to
resist disease. Therefore once such over-expresser plants were created, the effect of elevated AT4G11100 expression on plant immunity and defence gene expression could be examined.

To generate transgenic plants over-expressing AT4G11100, the pFAST-G02::AT4G11100 containing A. tumefaciens strain described above was used to transform 4 week old Col-0 A. thaliana plants. The transgene inserted into the pFAST-G02 is under the control of a CaMV 35S viral promoter and is therefore expressed constitutively. The phosphinothricin-containing herbicide BASTA was sprayed on T1 offspring of the transformed plants to select for transgenic individuals. T2 seed was harvested from the T1 survivors of herbicide selection, and again subjected to herbicide selection. T2 lines that displayed a 3:1 survival ratio were allowed to set seed as this indicated that only a single insertion event had occurred. T3 seedlings were treated with herbicide, and lines that displayed 100% survival were taken to be homozygous for the pFAST-G02::AT4G11100 transgene. In this way 7 plant lines homozygous for the transgene were identified.

Transformation of plants using A. tumefaciens will result in random insertion of the transgene into the plant genome, and depending on where in the genome the transgene inserts, there will be different levels of expression of the insert. For this reason, it was important to isolate pFAST-G02::AT4G11100 transgenic plant lines that originated from different transformation events and then to analyse their expression of AT4G11100. Expression of AT4G11100 in seven homozygous T3 transgenic plant lines arising from independent transformation events was assessed using qPCR (Figure 13). Relative to the ACTIN2 reference gene, three pFAST-G02::AT4G11100 transgenic plant lines (O-E 1, O-E 2,
and O-E 3) were identified as good candidates for further analysis based on their low, mid, and high \( AT4G11100 \) expression levels respectively.

Figure 13 | Relative \( AT4G11100 \) mRNA levels in \( AT4G11100 \) over-expressing (O-E) plant lines. O-E lines were made by transforming wild type Columbia (Col-0) plants with a \( pFAST-G02::AT4G11100 \) construct; different O-E lines represent different transformation events. mRNA levels were normalised against \( ACTIN-2 \). Plants were grown for four weeks under a 16 hour light/8 hour dark cycle at 22°C.

O-E 1 had \( AT4G11100 \) mRNA levels that were not significantly different to those in wild type Col-0 plants (and so was used as a vector control), but both O-E 2 and O-E 3 had considerably higher \( AT4G11100 \) mRNA levels than Col-0, with 8 and 30 times more respectively. It was fortuitous to have found transgenic lines with this range of \( AT4G11100 \) expression as it could make any \( AT4G11100 \) dose dependant effects easier to find.
AT4G11100 over-expressers display increased susceptibility to Pseudomonas syringae

If AT4G11100 is a negative regulator of plant immunity, then it is reasonable to expect that over-expression of AT4G11100 will result in increased susceptibility to infection; the over-expression of a negative regulator might dampen the defence response mounted against pathogen attack. The effect of over-expression of AT4G11100 on plant defence was examined by infecting 4 week old Col-0 and O-E plants with Pst DC3000 and quantifying bacterial growth 4 h and 48 h post infection (Figure 14).

Figure 14 | AT4G11100 over expressers (O-E) are more susceptible to P. syringae infection than wild type plants. Mean P. syringae titre (+SE, n = 5) in wild type Columbia (Col-0) and three AT4G11100 over expresser lines 4 and 48 h post infection. Plants were grown for four weeks at 22°C. Letters indicate significant differences (p < 0.05) determined by ANOVA with Fisher LSD. This experiment is representative of three independent experiments.
The AT4G11100 over expressers displayed enhanced susceptibility to Pst DC3000 infection when compared to wild type Col-0, and interestingly, they did so in a manner mirroring the expression level of AT4G11100. At 48 h post infection, O-E 1, with AT4G11100 expression no higher than Col-0, did not have a significantly higher bacterial titre than Col-0; whereas both O-E 2 and O-E 3, with significantly higher levels of AT4G11100 expression, had significantly higher bacterial titres than Col-0 (three and five fold increases respectively). This would appear to confirm the role of AT4G11100 as a negative regulator of the defence response, with increased expression of AT4G11100 leading to suppression of immunity and therefore increased pathogen susceptibility.

Additional experiments were performed to determine if there is any association between expression of AT4G11100 and defence gene expression, with the hypothesis that over-expression of At4g11000 would lead to reduced induction of defence genes after infection with Pst DC3000. Here, expression of the defence genes PATHOGENESIS RELATED 2 (PR2) and ISOCHORISMATE SYNTHASE 1 (ICS1) was examined in 4 week old O-E 1 and O-E 3 plant tissue infected with Pst DC3000, at 4h and 48h post infection. O-E 1 and O-E 3 were chosen for initial experiments as they display wild-type and highest expression of AT4G11100 respectively, and behave differently in response to Pst DC3000 infection. Figures 15 shows preliminary data from the single experiment carried out.
Figure 15 | Relative PR2 and ICS1 mRNA levels in two AT4G11100 over-expressing (O-E) plant lines. Relative expression of PR2 (+SE, n = 3) and ICS1 (+SE, n = 3) in two AT4G11100 O-E lines, 4 and 48 h post infection with P. syringae. mRNA levels were normalised against ACTIN-2. Plants were grown for four weeks under a 16 hour light/8 hour dark cycle at 22°C. Letters indicate significant differences (p < 0.05) determined by ANOVA with Fisher LSD.
Initial data shows that PR2 expression following infection with Pst DC3000 is significantly lower in O-E 3, where AT4G11100 expression is higher, compared to O-E 1, where AT4G11100 expression is at wild-type levels. This suggests that the elevated expression of AT4G11100 in O-E 3 might be inhibiting the induction of PR2 expression. The opposite trend is observed when ICS1 expression is examined; ICS1 is expressed at significantly lower levels after infection with Pst DC3000 when AT4G11100 expression is low (O-E 1), compared to when AT4G11100 expression is high (O-E 3).

Although these opposite patterns in defence gene expression might appear contradictory, the relative positions of ICS1 and PR2 in the defence signalling pathway might provide an explanation. ICS1 is responsible for the synthesis of salicylic acid (SA), an important defence molecule that acts at the beginning of a signalling pathway that results in the expression of a large number of plant defence genes, including the PR defence genes (Yan & Dong, 2014). Additionally, in the absence of pathogen attack, ICS1 expression is down regulated by the transcription factor CALMODULIN BINDING PROTEIN 60a (CBP60a) (Seyfferth & Tsuda, 2014). Figure 15 shows that over expression of AT4G11100 lowers induction of PR2, but not that of ICS1. It is conceivable that AT4G11100 may prevent the negative regulation of ICS1 by CBP60a and in this way block SA signalling downstream of SA production, but above PR2 production. This would explain the difference in expression observed between the two genes in AT4G11100 over expressers, and potentially gives insight into where AT4G11100 might act in its regulation of plant immunity. However, these results are preliminary and the experiments need to be repeated, specifically with the inclusion of expression data from wild type plants.
**AT4G11100 over-expresser has distinct leaf morphology**

While the AT4G11100 over expressing O-E 3 plant line displayed increased susceptibility to *P. syringae* infection, it is also worth noting that O-E 3 plants exhibited a leaf morphology distinct from both the other over expressing lines (O-E 1 & O-E 2; both of which express AT4G11100 at considerably lower levels than O-E 3), and wild type plants. At 4 weeks, compared to both wild type Col-0 and other over expresser lines, O-E 3 plant leaves have more serrated edges, particularly in younger leaves (Figure 16). The shapes of the O-E 3 leaves are noticeably different from wild type, and also appear to have a rougher surface.

![Figure 16](image_url)

**Figure 16**  | **Distinct leaf morphology of plants highly over expressing AT4G11100 (O-E 3).** Leaves are arrayed from oldest to youngest (left to right). Arrays from wild type Columbia (Col-0) and a moderate AT4G11100 over expresser (O-E 2) are shown. Plants were grown for four weeks under a 16 hour light/8 hour dark cycle at 22°C.
The surface of O-E 3 plant leaves was examined using a dissecting microscope, and it became apparent that the rough texture of O-E 3 leaves visible in the leaf arrays is due to an increased number of trichomes (Figure 17A). The number of trichomes per leaf area was determined, and it was clear that O-E 3 had a significantly higher number of trichomes per leaf area (approximately double) than both wild type and other AT4G11100 over expressing plants (Figure 17B).
Figure 17 | The highest AT4G11100 over expresser (O-E 3) has more trichomes than wild type plants. Leaf 10 from O-E 3 and Columbia (Col-0) was examined under a dissecting microscope (A) and the number of trichomes per cm$^2$ (+SE, n = 5) was calculated for leaf 6 from Col-0, O-E 2 and O-E 3 (B). Letters indicate significant differences (p < 0.05) determined by ANOVA with Fisher LSD. Plants were grown for four weeks under a 16 hour light/8 hour dark cycle at 22°C.
It is important to note that while the distinct morphological phenotype of O-E 3 is interesting, the role of AT4G11100 in generating the phenotype should not be over stated. The data show that O-E 3 has an increased number of trichomes compared to wild type, not necessarily that increased expression of AT4G11100 leads to higher numbers of trichomes. The high number of trichomes present in O-E 3 leaves might be due to the position in the genome that the pFAST-G02::AT4G11100 construct inserted. If the construct happened to disrupt a locus involved in leaf development, this might explain the trichome phenotype. However, if the increased expression of AT4G11100 does lead to higher numbers of trichomes, this could provide additional insight into the function and role of AT4G11100. To more definitively determine the effect of AT4G11100 expression on leaf morphology it would be useful to analyse additional independent AT4G11100 over expressers with high transgene expression from a separate transformation event.

However, it is interesting to note that in a series of yeast two hybrid assays AT4G11100 was found to interact with two NB-LRR RLKs, AT2G36570 and AT3G50230 (Mukhtar et al., 2011), both of which are implicated in plant development. And it has been suggested that AT2G36570 (PXC PXY/TDR-CORRELATED GENE 1, or PXC1) in particular is involved in plant vascular development and secondary cell wall formation in xylem fibres (Wang et al., 2013). It is possible that altered levels of AT4G11100 may disrupt function of these NB-LRR RLKs, the first step towards testing this would be to determine whether AT4G11100 actually interacts with AT2G36570 and AT3G50230 in planta. One means of testing such an interaction would be through the use of a bimolecular fluorescence complementation assay in which two halves of a fluorescent protein (yellow fluorescent protein, or YFP, as
AT4G11100:GFP may be unstable – see below) are fused to AT4G11100 and an RLK, and if they interact in planta, the two halves of YFP will come together and fluoresce.

Subcellular localization of AT4G11100

It seems likely that AT4G11100 plays a role in regulating plant immunity; manipulating the expression of AT4G11100 has a clear impact of the ability of A. thaliana to resist bacterial infection. Understanding the mechanisms governing how AT4G11100 behaves on a subcellular level is therefore of great interest, particularly with regard to how AT4G11100 may regulate the immune response. With this in mind, finding where AT4G11100 localises in the cell could help to elucidate how AT4G11100 functions.

To determine where AT4G11100 localises, a pk7FGW2::AT4G11100 vector was created through Gateway LR recombination of the AT4G11100 containing pENTR™4 vector (see page 8 above) with the pk7FGW2 destination vector. The resulting pk7FGW2::AT4G11100 vector allows for the translational fusing of the C-terminus of AT4G11100 with eGFP, resulting in an AT4G11100::eGFP fusion protein when expressed. The pk7FGW2::AT4G11100 vector was sequenced to confirm that the GFP tag was in frame with AT4G11100 and transfected into two week old maize protoplasts which were then examined under a confocal microscope (Figure 18).
No fluorescence in \textit{pk7FGW2::AT4G11100} transfection of maize protoplasts. Two week old maize protoplasts transfected with either \textit{pk7FGW2::IMPL2-1-60} as a positive control (+), the \textit{pk7FGW2::AT4G11100} fusion vector (\textit{AT4G11100}), or transfection solution without DNA as a negative control (-). The \textit{IMPL2-1-60} gene has been truncated so that its protein will not contain plastid transit peptides and thus localises in the cytoplasm. Green fluorescence was detected using a confocal microscope after laser excitation at 395nm. Plants were grown under a 16 hour light/8 hour dark cycle at 22°C. Red bars indicate a distance of 10µm.

The transfection was successful with the \textit{pk7FGW2::MYOINOSITOL MONOPHOSPHATASE-LIKE2 (IMPL2)-1-60} construct acting as a positive control. The \textit{IMPL2-1-60} gene has been truncated to remove the first 60 amino acids coded; this region contains the transit peptides necessary for IMPL2 plastid localisation, and thus the truncated IMPL2-1-60:eGFP fusion protein will localise in the cytoplasm (Petersen \textit{et al.}, 2010). The \textit{pk7FGW2::IMPL2-1-60} transfection showed clear green fluorescence in the protoplast cytoplasm. Unfortunately,
none of the protoplasts transfected with the *pk7FGW2::AT4G11100* fusion vector showed any green fluorescence above background levels. The transfection experiments were repeated a number of times, using both *A. thaliana* and maize protoplasts; however green fluorescence was never observed in protoplasts transfected with *pk7FGW2::AT4G11100*.

Concurrently with the transfection experiments, the creation of stable plant lines containing the *pk7FGW2::AT4G11100* fusion vector was carried out. The rationale behind the creation of these transgenic plants was that *in vivo* observations of AT4G11100::eGFP might be possible. Once the transfections failed to produce any information on the subcellular localisation of AT4G11100, it was hoped that the *pk7FGW2::AT4G11100* containing plants might yield better results.

Therefore, a GV3101 *A. tumefaciens* strain was transformed with *pk7FGW2::AT4G11100*, which was in turn used to transform 4 week old wild type *A. thaliana* plants. The pk7FGW2 vector contains a kanamycin resistance gene that could be used to select for positive transformants, and so multiple homozygous *pk7FGW2::AT4G11100* containing lines, from various separate transformation events, were then isolated through several generations of kanamycin selection.

Once plants homozygous for the *pk7FGW2::AT4G11100* vector had been identified, whole leaves from various lines at various stages of development were examined under a confocal microscope. Figure 19 shows images taken of a two week old transgenic plant; no GFP fluorescence above background levels was detected at any of the developmental stages examined. The same result was found in all lines studied.
The lack of fluorescence in a single pk7FGW2::AT4G11100 transgenic plant line might be due to the insertion of the vector in a transcriptionally silent region of the genome; however, because multiple transgenic plant lines had been created from multiple insertion events, it seemed unlikely that the pk7FGW2::AT4G11100 construct was being silenced in all lines. A western blot was performed using total protein extracted from pk7FGW2::AT4G11100 transgenic plant lines and eGFP antibodies in an effort to determine whether the eGFP fusion protein was actually expressed in the transgenic plant lines (Figure 20). However, none of the several independently transformed lines tested exhibited any binding of eGFP antibody, suggesting that no eGFP was being produced in any of the transgenic plant lines.

Figure 19 | No fluorescence in transgenic pk7FGW2::AT4G11100 containing plant leaves. Leaves of two week old transgenic A. thaliana plants containing the pk7FGW2::AT4G11100 fusion vector, viewed with a confocal microscope. The left hand panel shows the leaf under laser excitation at 395nm. The right hand panel is the same section with no light excitation. Plants were grown under a 16 hour light/8 hour dark cycle at 22°C. Red bars indicate a distance of 20µm.
Figure 20 | eGFP protein was not detected in pk7FGW2::AT4G11100 containing transgenic plant lines. 40mg total protein was extracted from 4 week old A. thaliana plants containing the pk7FGW2::AT4G11100 fusion vector (4 independent lines), and probed with an eGFP antibody in a Western blot (lanes 5 – 8). Bacterially expressed eGFP was used as a positive control (lanes 1 and 2), and total protein from 4 week old Columbia plants used as a negative control (lanes 3 and 4). A 30kDa band corresponds with the size of the eGFP protein.

The absence of eGFP in any of the pk7FGW2::AT4G11100 transgenic plant lines explains why no fluorescence was detected in these plants; however, the cause of the absence of eGFP in these plants is not clear. A total lack of expression of the pk7FGW2::AT4G11100 construct in the transgenic plants seems an unlikely cause as all the stable lines exhibit kanamycin resistance, which is conferred by the vector. If the expression of the AT4G11100::eGFP fusion is being somehow prevented, this would offer a reason for the failure of eGFP fluorescence. Equally, if the AT4G11100::eGFP protein fusion is unstable, this would explain
why both the protoplast transfections and the transgenic plant lines do not fluoresce. In either case, further experiments would need to be performed to determine why the eGFP is not being produced. RT-PCR analysis of the transgenic plant lines would reveal if the \textit{AT4G11100::eGFP} is actually being transcribed. Another course of action may be to create another fusion of \textit{AT4G11100} and eGFP, but as an N-terminal fusion as opposed to the C-terminal fusion performed here; this might circumvent any stability issue that may be associated with the current fusion protein.
Chapter 4: Conclusion

Plants possess a complex system of defence to prevent pathogen establishment, including two branches of innate immunity. Tight control of these defence systems is necessary to ensure appropriate timing and severity of immune responses in reaction to specific pathogens, including suppression of the immune system in the absence of pathogen attack. The *A. thaliana* cir1 null mutant is characterised by constitutive expression of defence genes, and enhanced resistance to *Pst* DC3000, suggesting that *CIR1* is a likely negative regulator of defence (Murray *et al.*, 2002).

Relatively early in the course of this project it was discovered the cir1 mutant displays a temperature sensitive growth phenotype (Figure 3). This led to an investigation into the effects of temperature on the cir1 defence phenotype, which revealed that the cir1 defence phenotype is actually temperature dependent (Figure 4). Efforts to identify the gene responsible for *CIR1* have previously led to the identification of *AT4G11100* as a likely *CIR1* candidate (Carstens, 2008; Diener, 2012), and this continued to appear likely as *at4g11100* T-DNA knockouts displayed a similar defence phenotype to cir1 (Figure 9). However, through complementation tests and genetic crosses (Figure 11 & Figure 12), this project has revealed that *AT4G11100* is not *CIR1*.

The creation and analysis of *AT4G11100* over-expressers shows that increased expression of *AT4G11100* causes increased susceptibility to infection by *Pst* DC3000 (Figure 14), suggesting that, like cir1, *AT4G11100* is a likely negative regulator of defence. Additionally, a transgenic plant line with high levels of *AT4G11100* expression displays distinct leaf
morphology, with serrated edges (Figure 16) and increased trichome density (Figure 17). This may implicate \textit{AT4G11100} in plant development, though independent \textit{AT4G11100} over-expressing lines would need to be created and analysed to confirm this.

Unfortunately, subcellular localisation of the \textit{AT4G11100} protein could not be determined despite efforts to create an \textit{AT4G11100}:eGFP fusion protein (Figure 18 & Figure 19), potentially as a result of problematic expression or stability of the fusion.

The creation and analysis of independent \textit{AT4G11100} over-expressing plant lines could help determine if the distinct leaf morphology seen in the over-expressor line described in this project is actually due to high \textit{AT4G11100} mRNA/protein levels. Additionally, further examination of the two NB-LRR RLKs (AT2G36570 and AT3G50230) that are thought to interact with \textit{AT4G11100} (Mukhtar \textit{et al.}, 2011) could prove useful, specifically in determining if they do in fact interact with \textit{AT4G11100} \textit{in planta} via a bimolecular fluorescence complementation assay, as the yeast two hybrid method used by Mukhtar \textit{et al.} (2011) often gives false positives. The qPCR experiments examining relative expression of \textit{ICS1} and \textit{PR2} in an \textit{AT4G11100} over-expressor (Figure 15) need to be repeated to confirm that the expression data found is significant. Furthermore, it may be interesting to investigate whether \textit{AT4G11100} interacts with the negative regulator of \textit{ICS1}, CBP60a, and possibly explain why \textit{ICS1} and \textit{PR2} expression differs in \textit{AT4G11100} over-expressing plant lines. To help dissect where in the SA defence pathway \textit{AT4G11100} is exerting a negative effect, expression levels of more defence genes in \textit{AT4G11100} over-expressers could be examined.
Overall, this study has identified novel, temperature dependant phenotypes of the cir1 mutant; ruled out AT4G11100 as the gene responsible for the cir1 phenotypes; and implicated AT4G11100 in the negative regulation of the plant immune system. Although further studies will need to be carried out to identify CIR1, the findings presented here contribute to the understanding of the complex defence systems utilised by A. thaliana in defence against pathogen attack.
Chapter 5: References


