Characterisation and expression of Receptor-like Cytoplasmic Kinases in *Medicago truncatula* during Rhizobial and Arbuscular Mycorrhizal symbioses

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Abstract

Legume plants have evolved the ability to form a mutualistic symbiosis with nitrogen-fixing bacteria known as rhizobia. Studies using mutant *Medicago truncatula* and *Lotus japonicus* have identified many genes that are necessary for this mutualism. Some of these genes have also been shown to be necessary for symbiosis with arbuscular mycorrhiza fungi (AMF). These common genes make up part of a signalling pathway known as the common symbiosis pathway (CSP) which serves to prepare the plant for the entry of the symbiont into the root. Although the core members of this signalling pathway are known there are still large gaps, and many of the genes involved in the infection process haven’t yet been identified. Using the *Medicago* Gene Expression Atlas (MtGEA) and in-house gene expression data three family Receptor-like Cytoplasmic Kinase (RLCK) genes were identified in *M. truncatula* which were differentially regulated during rhizobial and AMF colonisation. Analysis of *Tnt1* insertion mutants for the one of these genes, named *RLCK3* due to similarity to the rice gene family RLCK-OS3, revealed an apparently novel mycorrhizal phenotype with stunted arbuscule development. However, segregation analysis showed the phenotype was not linked to *RLCK3*; as such the locus was named *SCOOPY*. The other genes identified were similar to the rice gene family RLCK-XV and so named *RLCK1* and *RLCK2*. Both *RLCK1* and *RLCK2* encode soluble RD kinases with high homology to each other. Promoter-GUS analysis showed *RLCK1* and *RLCK2* expression associated with entry and accommodation of the symbionts. *rlck1* and *rlck2* mutants had a low level of arbusculation and misshaped arbuscules. *rlck1* was hyperinfected by rhizobia and had a significant reduction in nodule number. RNAi knockdown of *rlck1/rlck2* had a significantly reduced nodule number compared to the empty vector control. *RLCK1* and *RLCK2* possibly have partially redundant or synergistic roles during rhizobial and arbuscular mycorrhizal symbioses.
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<tr>
<td>AMF</td>
<td>Arbuscular Mycorrhiza Fungi</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine Tri-Phosphate</td>
</tr>
<tr>
<td>BR</td>
<td>Brassinosteroid</td>
</tr>
<tr>
<td>CSP</td>
<td>Common Symbiosis Pathway</td>
</tr>
<tr>
<td>dsRNA</td>
<td>Double Stranded RNA</td>
</tr>
<tr>
<td>DWA</td>
<td>Distilled Water Agar</td>
</tr>
<tr>
<td>EMS</td>
<td>Ethyl methanesulfonate</td>
</tr>
<tr>
<td>ENSA</td>
<td>Engineering Nitrogen Symbiosis for Africa</td>
</tr>
<tr>
<td>EV</td>
<td>Empty Vector</td>
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<tr>
<td>FST</td>
<td>Flanking Sequence Tag</td>
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<td>GFP</td>
<td>Green Fluorescent Protein</td>
</tr>
<tr>
<td>hpRNA</td>
<td>Hairpin RNA</td>
</tr>
<tr>
<td>LCO</td>
<td>Lipo-chitooligosaccharide</td>
</tr>
<tr>
<td>LRR</td>
<td>Leucine-Rich Repeat</td>
</tr>
<tr>
<td>LTR</td>
<td>Long Terminal Repeats</td>
</tr>
<tr>
<td>LysM</td>
<td>Lysin-motif</td>
</tr>
<tr>
<td>IT</td>
<td>Infection Thread</td>
</tr>
<tr>
<td>MAMP</td>
<td>Microbe-Associated Molecular Pattern</td>
</tr>
<tr>
<td>MAP</td>
<td>Mitogen-activated Protein</td>
</tr>
<tr>
<td>MtGEA</td>
<td>Medicago Gene Expression Atlas</td>
</tr>
<tr>
<td>mRNA</td>
<td>Messenger RNA</td>
</tr>
<tr>
<td>miRNA</td>
<td>MicroRNA</td>
</tr>
<tr>
<td>NBT</td>
<td>Nitro Blue Tetrazolium</td>
</tr>
<tr>
<td>NF</td>
<td>Nod Factor</td>
</tr>
<tr>
<td>PAM</td>
<td>Periarbuscular membrane</td>
</tr>
<tr>
<td>PAMP</td>
<td>Pathogen-Associated Molecular Pattern</td>
</tr>
<tr>
<td>PAS</td>
<td>Periarbuscular space</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase Chain Reaction</td>
</tr>
<tr>
<td>Acronym</td>
<td>Description</td>
</tr>
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<tr>
<td>PIT</td>
<td>Preinfection Thread</td>
</tr>
<tr>
<td>PPA</td>
<td>Prepeneratton Apparatus</td>
</tr>
<tr>
<td>PTI</td>
<td>PAMP-triggered Immunity</td>
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<tr>
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<td>Quantitative Reverse Transcription PCR</td>
</tr>
<tr>
<td>RD</td>
<td>Arginine-aspartate</td>
</tr>
<tr>
<td>RKN</td>
<td>Root Knot Nematode</td>
</tr>
<tr>
<td>RLK</td>
<td>Receptor-Like Kinase</td>
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<tr>
<td>RLCK</td>
<td>Receptor-Like Cytoplasmic Kinase</td>
</tr>
<tr>
<td>RNAi</td>
<td>RNA interference</td>
</tr>
<tr>
<td>RNS</td>
<td>Root Nodule Symbiosis</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive Oxygen Species</td>
</tr>
<tr>
<td>RTK</td>
<td>Receptor Tyrosine Kinases</td>
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<tr>
<td>sRNA</td>
<td>Silencing RNA</td>
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<tr>
<td>siRNA</td>
<td>Short Interfering RNA</td>
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<td>TGAC</td>
<td>The Genome Analysis Centre</td>
</tr>
<tr>
<td>TSL</td>
<td>The Sainsbury Laboratory</td>
</tr>
<tr>
<td>WGA</td>
<td>Wheat Germ Agglutinin</td>
</tr>
<tr>
<td>WGD</td>
<td>Whole Genome Duplication</td>
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<tr>
<td>WT</td>
<td>Wildtype</td>
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Chapter 1: Introduction
The sessile nature of plant life can make the necessary issue of finding nutrition and water a challenge. Plants can only make use of nutrients that are within the rhizosphere, which are sometimes obtained through the uptake of water such as mineral ions like potassium (K\(^+\)) and ammonia (NH\(_3\)) (Taiz and Zeiger, 1998). However, if the nutrients the plant requires are not within the rhizosphere, or not in such an accessible form, then the plant employs different strategies to obtain the nutrients it requires. One method which the plant uses is extended root growth, either through the lengthening of the tap root or increased branching, to increase the size of the rhizosphere. Plants also exude organic acids into the soil in order to free nutrients which are otherwise bound in salts; Phosphorous (P) is rarely found in a free (ionic) state in soil, tending to be in a salt on the surface of rocks and stones (Taiz and Zeiger, 1998). Despite all the strategies that plants have evolved to cope with these problems on their own, plants can still find themselves in environments where they are not able to find adequate nutrition or water. This is when beneficial symbiosis, or mutualism, becomes advantageous for the plant (Taiz and Zeiger, 1998).

The earliest evidence of a mutualistic symbiosis by a plant with a fungus is a fossil from the Rhynie Chert containing examples of the plant-mycorrhiza symbiosis from ~450 million years ago (mya) (Remy et al., 1994). It is thought that the symbiosis with mycorrhizal fungi was important in the colonisation of land by plants (Harrison, 2005). The first land plants did not have an extensive root system like higher plants. Ancient sea-dwelling plants did not need an extensive root system as the nutrients were dissolved in the water which was all around them. They did have a small root like structure (rhizoid) but this was more for anchorage than nutrient acquisition (Jones and Dolan, 2012). Early diverging lineages of plants, such as the hornworts and liverworts, are restricted to watery areas of land. The mycorrhizal fungi would have acted like a surrogate root system to the early land plants as they do to hornworts and liverworts today (Parniske, 2008). In exchange for providing water and nutrients for the plants, the mycorrhizal fungi receive a supply of carbon in the form of sugars made by the plant during photosynthesis. This symbiosis is present in ~70-90 % of modern day plants (Peterson et al., 2004; Parniske, 2008). It is believed that root nodule symbiosis (RNS), a symbiosis between plants in the Eurosid I clade, including Fabales, Fagales, Cucurbitales and Rosales, and Gram-negative soil bacteria collectively known as rhizobia, evolved from the earlier mycorrhizal fungi-plant symbiosis ~60 mya (Kistner and Parniske, 2002). It is believed that a Whole Genome Duplication (WGD) event at ~58 mya might have led to the recruitment of genes involved in the mycorrhizal symbiosis into the emerging symbiosis with nitrogen fixing bacteria (Parniske, 2000; Cannon et al., 2006; Parniske, 2008; Young et al., 2011).
1.1 Legume Endosymbiosis

Legumes (Fabaceae) are able to form symbioses with both arbuscular mycorrhizal fungi (AMF) and rhizobia within their root cortical cells. Model legumes *Medicago truncatula* (Medicago) and *Lotus japonicus* (Lotus), as well as other legumes such as *Glycine max* (soybean) and *Pisum sativum* (pea) and non-nodulating plants such as *Oryza sativa* (rice) and *Petunia hybrida* (petunia), have been used to investigate these root symbioses in more detail. Unless otherwise stated gene names used will be from *M. truncatula* with *L. japonicus* following in square brackets.

Before the symbionts come into physical contact they communicate via a chemical dialogue through the rhizosphere. Plants exude flavonoid or strigalactone signals, attracting symbionts towards the plant roots where the rhizobia attach to the growing root hair and the AMF form hyphopodia, a specialised, lobed hypha that attaches the fungus to the root surface (Firmin *et al.*, 1986; Redmond *et al.*, 1986; Besserer *et al.*, 2006; Peck *et al.*, 2006). The symbionts exude lipo-chitooligosaccharides, Nod Factors (NF) or Myc Factors (myc-LCOs), which are perceived by the plants and alerts them of a nearby symbiotic partner (Denarie *et al.*, 1996; Maillet *et al.*, 2011). AMF also exude tetra- and pentameric chitin oligomers (COs) that act as symbiosis signalling compounds (Genre *et al.*, 2013). NFs are perceived at the root epidermis by the LysM Receptor-like Kinases (LysM-RLKs) NFP [NFR5] and LYK3 [NFR1] (Limpens *et al.*, 2003; Madsen *et al.*, 2003; Radutoiu *et al.*, 2003; Radutoiu *et al.*, 2007). The Myc Factor receptor has not yet been identified. There is some evidence that a LysM receptor might be required for Myc Factor perception; Op den Camp *et al.* (2011) showed that a knockdown of the MtNFP orthologue in the non-legume *Parasponia abndersonii* (PaNFP) inhibited colonisation of roots by both AMF and rhizobia. Perception of these symbiont signals initiates a signalling cascade within the epidermal cell that results in the expression of symbiotic specific genes.

1.1.1 The Common Symbiosis Pathway and symbiotic gene expression

Genetic studies in *M. truncatula* and *L. japonicus* have identified a set of genes that are required for both RNS and arbuscular mycorrhization that are downstream of the NF receptors in the signalling cascade. A leucine-rich repeat-RLK *Does Not Make Infections (DMI) 2 [SYMRLK]* (Endre *et al.*, 2002; Stracke *et al.*, 2002), a nuclear envelope-localised cation channel *DMI1 [POLLUX]* (Ané *et al.*, 2004; Imaizumi-Anraku *et al.*, 2005; Charpentier *et al.*, 2008; Capoen *et al.*, 2011), a calcium and calmodulin dependent kinase *DMI3 [CCaMK]* and *Interacting protein of DMI3 (IPD3) [CYCLOPS]* (Catoira *et al.*, 2000; Lévy *et al.*, 2004; Mitra *et al.*, 2004; Messinese *et al.*, 2007; Horváth *et al.*, 2011)
are the core components of the common symbiosis pathway (CSP) with homologues being identified in both *M. truncatula* and *L. japonicus*. In *L. japonicus* nucleoporins NUP85, NUP133 and NENA (Kanamori et al., 2006; Saito et al., 2007; Groth et al., 2010), and cation channel CASTOR (Imaizumi-Anraku et al., 2005) have been shown to be part of the CSP, however their orthologues in *M. truncatula* have not yet been identified.

The CSP transfers the signal received at the cell surface to the nucleus to initiate calcium oscillations known as calcium spiking (Ehrhardt et al., 1996; Sieberer et al., 2009; Maillet et al., 2011). NF, Myc-LCOs and Myc-COs elicit distinct calcium spiking patterns (Denarie et al., 1996; Maillet et al., 2011; Genre et al., 2013). DMI2, DMI1 [POLLUX], CASTOR and the nucleoporins have all been shown to be required for spiking (Wais et al., 2000; Kanamori et al., 2006; Saito et al., 2007; Charpentier et al., 2008; Groth et al., 2010). It is widely believed that DMI3 [CCaMK] decodes the calcium spiking (Miller et al., 2013). dmi3 and ccamk mutants retain calcium spiking but are not able to form symbioses (Wais et al., 2000). Gain-of-function mutations in *DMI3* (*DMI3*/*ccamk*) are capable of producing nodule structures in the absence of rhizobia (Gleason et al., 2006; Tirichine et al., 2007) and results in the expression of mycorrhizal specific markers in the absence of AMF (Takeda et al., 2012). Recent evidence from Singh et al. (2014) show that CYCLOPS is able to activate nodule organogenesis independently from the CSP when serines 50 and 154 are replaced with aspartic acids (CYCLOPS-DD) adding to the evidence that IPD3/CYCLOPS may work with DMI3 in the decoding of the calcium spiking.

Several transcription factors act downstream of the CSP. *Nodule inception* (*NIN*), *ERF required for nodulation 1 and 2* (*ERN1/ERN2*), and *Nodulation signalling pathway 1 and 2* (*NSP1/NSP2*) have been shown to be required for nodulation (Oldroyd and Long, 2003; Kaló et al., 2005; Smit et al., 2005; Heckmann et al., 2006; Andriankaja et al., 2007; Middleton et al., 2007). NSP1 and NSP2 have been shown to form a DNA binding complex that is able to bind to the promoter of symbiosis reporter gene *Early Nodulin 11* (*ENOD11*) (Hirsch et al., 2009). *Required for Arbuscular Mycorrhization 1* (*RAM1*) has been shown to be required for mycorrhization and RAM1 has been shown to interact with NSP2 (Gobbato et al., 2012). There is also some evidence that NSP1 and NSP2 may have a role in strigolactone synthesis (Liu et al., 2011). This may explain the mycorrhizal phenotype seen in the *nsp2* mutant (Maillet et al., 2011).

There are a number of genes that have been shown to be required for both RNS and mycorrhization that have not been shown to be part of the CSP and as such are known as common symbiosis genes. It is possible that 3-hydroxy-3-methylglutaryl
coenzyme A reductase 1 (HMGR1) is a part of the CSP but as yet is not confirmed. There is some evidence that HMGR1 interacts with DMI2 and RNAi knockdown of HMGR1 expression causes inhibition of RNS and calcium spiking (Kevei et al., 2007) and unpublished evidence from the Ané Lab (presented on a poster at Plant Biology 2011, Minneapolis, Minnesota) suggests that HMGR1 is also required for colonisation by, and calcium spiking in response to, AMF (Jayaraman, 2011). However, no stable mutant data for HMGR1 have been published to date. Vapyrin, a gene encoding an N-terminal VAMP-associated protein (VAP)/major sperm protein (MSP) domain with a C-terminal ankyrin-repeat domain, appears to be involved in infection stages of both RNS and mycorrhizal symbiosis and acts downstream of the CSP (Feddermann et al., 2010; Pumplin et al., 2010; Murray et al., 2011). It is likely that other components of the CSP exist.

1.1.2 Early stages of Rhizobial and AMF colonisation of Legume roots

For RNS, the perception of NF initiates two spatially and temporally co-ordinated developmental processes, one which allows the entry of the rhizobia into the plant root (infection), and the other creation of the nodule that will house the endosymbiotic nitrogen-fixing bacteria (nodule organogenesis) (Murray, 2011; Oldroyd et al., 2011). For mycorrhization, colonisation of the root by AMF can be separated into three stages; precontact, intraradical development, and arbuscule development (Gutjahr and Parniske, 2013). Although the plant does not make specialist organs for AMF colonisation it has been shown that perception of diffusible signals from the germinating AMF spores by the plant stimulates lateral root formation, increasing the root area available for AMF colonisation (Oláh et al., 2005; Maillet et al., 2011; Gutjahr and Paszkowski, 2013). Although the infection of the root by rhizobia and AMF are morphologically distinct they bear striking similarities at the subcellular level (Kistner and Parniske, 2002).

1.1.2.1 Rhizobial infection and Infection thread formation

On the perception of NF legume root hairs switch from symmetrical polar growth to growth towards the source of the NF, sometimes including root hair swelling and branching (Esseling et al., 2003). The growing root hair cell curls around attached NF-producing rhizobium trapping it within an infection pocket (Geurts and Bisseling, 2002). The rhizobia within the infection pocket divide to form a microcolony or infection focus. The nucleus of the curling root hair cell moves up to the infection focus and the cytoskeleton aligns along the shaft of the hair forming a bridge through the cytoplasm (Fournier et al., 2008).

An invagination of the plant cell wall and plasma membrane occurs at the infection focus growing inwardly through the root hair cell, following the migrating nucleus along the cytoplasmic bridge down towards the base of the cell creating the infection thread (IT). An
influx of calcium ions seen at the root hair tip which is accompanied by the generation of reactive oxygen species (ROS) is associated with the change of polar growth at the infection foci from an outward to an inward growth (Miwa et al., 2006; Murray, 2011; Oldroyd et al., 2011). The IT is a tube filled with a glycoprotein matrix, which is still technically external to the root hair cell, through the root hair along which the rhizobia grow and divide (Brewin, 1998). As the IT reaches the base of the root hair cell, the nucleus in the outer cortical cell directly below the progressing thread moves to the upper plasma membrane and guides the formation of the preinfection thread (PIT) (Van Brussel et al., 1992; Timmers et al., 1999; Niwa et al., 2001) which is very similar to what occurs in the root hair during infection. The formation of PITs and ITs continue in this way, ramifying and guiding the bacteria through the cell layers of the root, until it reaches the inner cortical cells comprising the nodule where the bacteria are released and begin to fix nitrogen (Timmers et al., 1999). Several mutants have been identified that are impaired in IT formation, creating arrested, lumpy or misshaped ITs, including lyk3, vapyrin, ern1, ern2, a remorin symrem1, lumpy infections (lin), and the knockdown mutants of flotillins flot2 and flot4 (Kuppusamy et al., 2004; Andriankaja et al., 2007; Middleton et al., 2007; Smit et al., 2007; Haney and Long, 2010; Lefebvre et al., 2010; Murray et al., 2011; Tóth et al., 2012).

1.1.2.2 AMF infection and PPA formation
AMF enter the plant root via atrichoblasts rather than root hair cells; despite this difference the changes within the epidermal cells that allow symbiont entry are very similar. Upon perception of mycorrhizal specific diffusible signals, myc-LCOs and hyphopodia formation on the root surface, the epidermal cell undergoes differential gene expression and cytoskeletal changes (Genre et al., 2005). The nucleus moves first to the top of the cell directly beneath the fungal hyphopodia, then moves down through the plant cell forming the pre-penetration apparatus (PPA) a tunnel of cytoskeleton and endoplasmic reticulum (similar to the PIT) through which the AMF hyphae can grow (Genre et al., 2005). The PPA guides the invagination of the plant cell wall creating a tube, like the rhizobial IT but larger, through which the AMF penetrate (Genre et al., 2005). As in rhizobial infection, this process is controlled by the plant and the AMF remains technically external to the plant cell. Once through the epidermal cell the hyphae either travels through the cell directly below the epidermal cell, which has already formed a PPA, or can travel intercellularly until it reaches the root inner cortex, but the AMF must pass intracellularly through at least one plant cell before it is able to form arbuscules (Demchenko et al., 2004).
1.1.3 Nodule development in Legumes

Whilst the root hairs at the epidermis are forming ITs to allow the entry of the rhizobial symbiont into the root, the nodule has already started to develop in the root cortex (Oldroyd and Downie, 2008). There are two types of legume root nodule: determinate, as seen on *L. japonicus* root, and indeterminate, as seen on *M. truncatula* roots. Determinate root nodules develop from the outer cortical cells, grow to a predetermined size and are normally round in shape. Indeterminate root nodules develop from the inner cortical cells and maintain an apical meristem causing them to be more elongated in shape (Sprent, 2001). I shall be referring to indeterminate root nodules unless otherwise stated.

To develop the nodule organ the cortical cells directly below the developing infection thread divide (Oldroyd and Downie, 2008). As the proliferation of cortical cells continues they eventually break through the epidermis causing the characteristic outgrowth, or bump, of the nodule primordia. The growing infection thread ramifies through the nodule primordia distributing the rhizobia across the nodule. When the rhizobia reach the cortical cells they are taken up via a process very similar to endocytosis, such that each bacterial cell is surrounded by a plant derived membrane known as the peribacteroid membrane (Udvardi and Day, 1997). At this stage the rhizobia irreversibly differentiate into bacteroids that are completely dependent on the plant (Mergaert *et al.*, 2006). Plant derived leghaemoglobin binds to oxygen to support respiration of the bacteroids and protect the nitrogenase allowing the bacteroids to fix nitrogen and giving the nodule its pink colour (Appleby, 1984; Starker *et al.*, 2006).

Mature indeterminate nodules contain five zones that progress from the nodule tip to the base (fig. 1.1.3.1) (Vasse *et al.*, 1990; Foucher and Kondorosi, 2000; Timmers *et al.*, 2000). Zone I is the meristem of the nodule. These cells continue to divide and elongate the nodule. Cells that remain in Zone I are never colonised by rhizobia. Zone II is the infection zone. The rhizobia continue to colonise this zone throughout the life of the nodule through ramifying infection threads which can be clearly seen on root nodule sections. Zone III is the nitrogen fixation zone where the cells are packed full of bacteroids. Zone IV is the senescent zone where the bacteria and leghaemoglobin are degraded, often a green-ish yellow in colour. In some nodules a zone between II and III, called the interzone (IZ) or Zone II-III, can be seen where cells undergo many symbiotic specific cell changes (Vasse *et al.*, 1990; Foucher and Kondorosi, 2000). Timmers *et al.* (2000) showed that Zone V comprises a saprophytic zone at the very base of the nodule. Cells start their life in Zone I and progress into Zone V as the nodule grows and matures, Zones III to V may not be seen in young developing nodules.
**Figure 1.1.3.1.** The Zones of an indeterminate nodule

A diagram to demonstrate the zones observed during indeterminate nodule development. Zone I is the nodule meristem. Zone II is the infection zone. Zone III is the fixation zone. Zone IV is the senescence zone. Zone V is the saprophytic zone.

1.1.4 Arbuscule development and genetic regulation in root cortical cells

AMF form arbuscules in the inner cortical cells of the plant root. Arbuscules are highly branched, tree-like structures that are the site of nutrient exchange between the AMF and the plant. The arbuscules are the defining structures of, and give their name to, the AMF. The arbuscule is surrounded by a plant derived membrane called the periarbuscular membrane (PAM), which is continuous with the plasma membrane, and an interfacial glycoprotein-rich matrix known as the periarbuscular space (PAS) made up of AMF and plant cell wall material (Parniske, 2008; Balestrini and Bonfante, 2014). The PAM contains nutrient transporters such as the phosphate transporter MtPT4 (Javot et al., 2007; Yang et al., 2012).

The formation of an arbuscule in a plant inner cortical cell can be genetically divided into 5 stages: 1) PPA formation or pre-arbuscule stage, 2) fungal entry/arbuscule trunk formation, 3) birdsfoot stage, 4) hyphal branching/mature arbuscule and 5) arbuscule collapse/senescence (fig.1.1.4.1) (Gutjahr and Parniske, 2013). The first stage of arbusculation is very similar to the PPA formation in epidermal cells prior to hyphal entry into the plant root. Laser capture microdissection shows that cortical cells ahead of the advancing intraradical mycorrhizal hyphae have differential gene expression compared to cortical cells on non-mycorrhizized plants, showing these cells are preparing to receive the AMF (Gomez et al., 2009; Hogekamp et al., 2011; Gaude et al., 2012).
CSP mutants do not complete stage one of arbuscule formation, although a gain-of-function DMI3 (CCaMK) mutant can induce the gene expression patterns seen in stage one (Takeda et al., 2012). In stage 2, the fungal hypha enters the cell forming the unbranched arbuscule trunk. Mutants such as vapyrin in *Medicago or penetration and arbuscule morphogenesis*1 (pam1) in petunia occasionally show the formation of arbuscule trunks but do not progress to the birdsfoot stage of arbuscule development (Reddy D. M. R et al., 2007; Feddermann et al., 2010; Murray et al., 2011).

The third stage of arbuscule development initiates the hyphal branching. This stage is referred to as the birdsfoot stage due to the resemblance of the hyphae to the toes on a bird’s foot (Gutjahr and Parniske, 2013). Mutants in two half-ABC transporter genes in *Medicago, Stunted arbuscule* (str and str2), progress only to the birds foot stage of arbusculation (Zhang et al., 2010b). The *Medicago* mutant of *Required for Arbuscular Mycorrhization* (RAM) 2 occasionally forms birdsfoot arbuscules when the AMF does get through to the cortex (Wang et al., 2012). In addition, overexpression of KP1106 (protease inhibitor), knockdown of SCP1 (serine carboxypeptidase 1) or ERF1 (Ethylene response factor 1) all result in the birdsfoot type arbuscules in *Medicago* inner cortical cells (Devers et al., 2013; Rech et al., 2013). Stage four of arbuscule development is the ramification of the AMF hyphae to form the highly branched, tree-like mature arbuscules. Live cell imaging in rice (*Oryza sativa*) determined the average life span of an arbuscule was on average 2 to 3 days but could be as little as one day (Kobae and Hata, 2010). MtPT4 and its rice homologue OsPT11 have been shown to be required for the maintenance of the mature arbuscules in mutant studies when nutrients other than phosphate were readily available (Javot et al., 2007; Yang et al., 2012). However, the pt4 phenotype was supressed when *Medicago* plants were nitrogen starved, yet not when starved of sulphate (Javot et al., 2011), suggesting a complex system of regulation for arbuscule maintenance that is dependent on nitrogen and phosphate availability.
Figure 1.1.4.1. Stages of arbuscule development.

A diagram to demonstrate the stages of arbuscule development. Stage I is PPA development. Stage II is the entry of the hyphae into the cell. Stage III is the birdsfoot stage showing a few larger hyphal ramifications. Stage IV is the hyphal branching/mature arbuscule. Stage V is the collapsed/senescing arbuscule.

Figure adapted from Gutjahr and Parniske (2013). *Annual Review of Cell and Developmental Biology*. 29: 593-617.
1.2 Similarities between symbiosis and pathogenesis of plants

There have been many reviews highlighting the similarities between the mutualisms of rhizobia and AMF, and the pathogenesis by fungi, oomycetes and nematodes with plants (Ausubel and Bisseling, 1999; Parniske, 2000; Kogel et al., 2006; Paszkowski, 2006; Rey and Schornack, 2013). Mutualism and pathogenesis are very different in terms of cost and gain for the plant, but they can also be seen as a sliding scale of biotrophism, with mutualism at one end and plant pathogens at the other with lots of examples in between.

For example, Kiers et al. (2011) demonstrated that two Glomus species and Rhizophagus irregularis (nee. G. intraradices) acquire differential amounts of carbon from the plant as well as differing in the amount of phosphorus they give in exchange. In fact G. aggregatum had almost double the carbon cost per phosphorus molecule than R. irregularis (Kiers et al., 2011), it could then be argued that G. aggregatum is less mutualistic and more parasitic than R. irregularis to the host plant M. truncatula. Another example of the blurred lines between friend and foe is the fungal endophyte Epichloë festucae and its host ryegrass Lolium perenne. A mutation in the NoxA gene in E. festucae is enough to switch it to a pathogen that causes great damage to L. perenne (Tanaka et al., 2006).

Many aspects of plant colonisation by beneficial symbionts are similar to those seen in plant-pathogen interactions. For example, the Root Knot Nematode (RKN) Meloidogyne incognita causes root hair deformation and nuclear movement similar to that seen with NF application and successful cortical colonisation by RKN is dependent on the CSP member DMI2 (Weerasinghe et al., 2005). Damiani et al. (2012) showed an overlap in gene expression in response to rhizobia in Zone II and RKN Giant Cells both of which are responsible for the accommodation of the microbe. NFP, a gene regarded as specific to Nod Factor perception, has been shown to play a role in resistance of M. truncatula to Aphanomyces euteiches, a pathogenic root oomycete (Rey et al., 2013). Chitin oligomers, which are thought to play a positive role in AMF colonisation of the root (Genre et al., 2013) also activate plant defence through the chitin receptor CERK1 in A. thaliana and rice (Petutschnig et al., 2010; Shimizu et al., 2010). However, as A. thaliana does not form a mutualistic relationship with AMF, plants that have co-evolved with AMF and/or rhizobia presumably have evolved a way of distinguishing the chitin signals through the mixture of chitin oligomers or a suite of receptors that modulate the response, for instance M. truncatula has a greatly expanded set of LysM RLKs (Arrighi et al., 2006; Petutschnig et al., 2010; Genre et al., 2013; Rey and Schornack, 2013). It has also been shown through M. truncatula mutant ram2 that cutin C16 monomers are required for the formation of both AMF hyphopodia and oomycete appresoria on the root surface (Wang et al., 2012).
The nuclear movement and cytoskeletal rearrangements seen during AMF infection can also be seen during oomycete and fungal infection, with the nucleus moving directly beneath the contact site. If the plant is resistant to the oomycete, the plant locally thickens the cell wall before the nucleus moves away. If the oomycete continues to invade the cell the association of the nucleus with the haustorium persists until hypersensitive cell death occurs. In susceptible cells, the nucleus does not move toward the oomycete (Guest, 1984; Guest, 1986; Freytag et al., 1994; Daniel and Guest, 2005). During fungal pathogenic infection the nucleus moves to the site of infection in both resistant and susceptible cells. If the fungus successfully enters the cell the nucleus moves away again, and cell death occurs if the plant is resistant. If the fungus is not able to penetrate the nucleus remains at the contact site (Heath et al., 1997). The receptor-like MLO has been shown to interfere with the polarisation of the actin cytoskeleton during infection (Opalski et al., 2005). Interestingly, the barley mlo mutant is more resistant to the biotrophic fungal pathogen *Blumeria graminis* in leaves but is also less able to form arbuscules in the roots (Ruiz-Lozano et al., 1999; Bhat et al., 2005; Opalski et al., 2005).

1.3 Plant Receptor-like Kinases
The ability to perceive and respond to information at the cell surface via receptors at the cell surface is a basic property for all living organisms (Stone and Walker, 1995). The ability to transfer that signal to the nucleus requires a signalling cascade through the cytoplasm. The phosphorylation and de-phosphorylation of proteins have been shown to play a central role in signal transduction from the cell surface (Stone and Walker, 1995). Protein kinases catalyse the transfer of the terminal phosphate group from ATP to the side chains of serine, threonine, tyrosine or histidine in a protein (Grebe and Stock, 1999; Alberts, 2002). De-phosphorylation is achieved through the action of protein phosphatases (Alberts, 2002). Approximately 1-3% of a eukaryotic genome is predicted to encode protein kinases, suggesting that they are important in many cellular processes (Stone and Walker, 1995).

In plants, the largest kinase superfamily is the receptor-like kinases (RLKs) (Shiu and Bleecker, 2001a), the first being discovered in *Zea mays* (Maize) by Walker and Zhang (1990). Generally plant RLKs are made up of a signal sequence, an external amino-terminal domain with a transmembrane region and an internal carboxy-terminal kinase domain, similar to the structure of animal receptor tyrosine kinases (RTKs), although the majority of plant RLKs are serine/threonine kinases (Walker, 1994; Shiu and Bleecker, 2001a; Shiu and Bleecker, 2001b; Shiu and Bleecker, 2003). In *A. thaliana* a predicted 610 genes belong to the RLK super family, making up approximately 2.5% of...
the protein-coding genes within the genome (Shiu and Bleecker, 2001a). Nearly twice the number of RLKs of *A. thaliana* has been predicted in the rice (*O. indica*) genome (Shiu and Bleecker, 2003; Shiu et al., 2004). Based on their extracellular domains there are 16 families of RLK: C-type Lectin, Crinkly4-like, CrRLK1-like, DUF26, Extensin, Legume Lectin, LRK10-like, Leucine-rich repeat (LRR), Lysin-motif (LysM), URK I, PERK-like, RKF3-like, S-Domain, Thaumatin, WAK-like and receptor-like cytoplasmic kinases (RLCKs) (Shiu and Bleecker, 2001a).

RLCKs are different from most of the RLKs as they lack the external receptor domain, and are similar in structure to the animal non-receptor tyrosine kinases (Shiu and Bleecker, 2001b; Shiu and Bleecker, 2003). Approximately 25% of plant RLKs are RLCKs and can be divided into 19 to 23 subfamilies, with rice having 4 more subfamilies than *A. thaliana* (Shiu et al., 2004). RLCKs can also be organised into different types of kinase; more than 70% of rice RLCKs having only a kinase domain, but the other 30% contain domains that are normally found in the extracellular domains of RLKs such as LRRs, Lectin, DUF, U box and LysM as well as PPR, ECH, USP, UBQ, SPERM and JACALIN (Vij et al., 2008).

RD kinases have an arginine adjacent to an aspartate in the active site of the kinase which can interact with a tyrosine, serine or threonine further down the active site when the aspartate is in a non-phosphorylated. In this inactive state the protein is folded by the amino acid interaction with the active site hidden inside the fold. When the aspartate is phosphorylated the interaction with the tyrosine, serine or threonine is broken and the protein is able to unfold opening up the active site of the kinase. In this activated state RD kinases are then able to phosphorylate other proteins. Therefore RD need to be activated by another kinases before they can function. RLKs can be broadly classified into two groups: The first controlling plant growth and development, such as CLAVATA1 (meristem control) (Clark et al., 1997) and the second involved in plant-microbe interactions, both mutualistic and pathogenic, such as FLS2 (flagellin perception) (Gómez-Gómez and Boller, 2000) and SYMRK (symbiont perception) (Stracke et al., 2002). Typically the RLKs that recognise pathogenic signals are non-RD kinases, but those involved in the recognition of a broad range of pathogens are often RD kinases. In contrast, most RLKs involved in developmental processes are often of the RD class of kinases. However, many non-RD RLKs associate with cytoplasmic RD kinases for signal transduction (Schwessinger and Ronald, 2012). The signalling pathways of plant growth via brassinosteroid signalling and plant defence through the FLS2 have been well studied in *A. thaliana*. Both of these signalling pathways (fig. 1.3.1) contain RLKs at the cell
surface and RLCKs, both RD and non-RD, and share similarities with the CSP, providing useful examples to compare with plant symbiosis signalling.

1.3.1 The brassinosteroid signalling pathway
Brassinosteroid (BR) is a plant hormone that regulates plant growth. Plants that cannot sense BR are dwarfed in height. The BR signalling pathway, characterised through mutant studies in A. thaliana and rice, is a signalling cascade from perception of BR at the cell surface through to regulation of gene expression (Kim and Wang, 2010). An RD, LRR-RLK, BRI1 (BR insensitive 1) is able to bind BR extracellularly which causes autoactivation of the internal kinase domain (Russinova et al., 2004; Wang et al., 2005b). BRI1 is found in the plasma membrane in homodimeric receptor complexes or in a heterodimer with another LRR-RLK BAK1 (BRI1-Associated receptor kinase 1) (Li et al., 2002; Nam and Li, 2002; Russinova et al., 2004; Wang et al., 2005b). The addition of BR was found to increase the levels of homo- and heterodimerisation of BRI1 (Wang et al., 2005a; Wang et al., 2008).

BRI1 kinase inhibitor protein (BKI1) is able to inhibit BRI1 by associating with the kinase domain of BRI1 in the absence of BR (Wang and Chory, 2006). When BR binds to BRI1, BKI1 is phosphorylated and dissociates from BRI1 and the plasma membrane enabling access to BRI1 for BAK1 (Wang and Chory, 2006). Modified BKI with an N-terminal myristoylation site stays associated with BRI1 even with BR application and causes the same dwarfed phenotype as the bri1 mutants (Wang and Chory, 2006). BR signalling kinases (BSKs), RLCKs of the RLCK-XII subfamily, also interact with BRI1 at the plasma membrane. BSKs are phosphorylated upon BR treatment and then dissociate from the membrane (Tang et al., 2008). Overexpression of the BSKs was able to rescue the strong bri1 phenotype (Tang et al., 2008), suggesting that these RLCKs are the next step in the signal cascade after the membrane bound RLKs.

BIN2 (Brassinosteroid insensitive 2) is a Gibberellin Signalling Kinase (GSK)-like protein that was identified as part of the BR signalling pathway though a semi-dominant mutation bin2 that displayed a dwarf phenotype like bri1 (Li et al., 2001). Therefore BIN2 plays a negative role on BR signalling. BIN2 phosphorylates the transcription factors BZR1 (Brassinole resistant 1) and BZR2/BES1 (bri1-EMS suppressor 1) (He et al., 2002; He et al., 2005) leading to their interaction with 14-3-3 proteins and ultimately proteasomal degradation (Yin et al., 2002; Yin et al., 2005; Vert and Chory, 2006; Gampala et al., 2007). The phosphorylation and association with 14-3-3 proteins stop BZR1 and BZR2 from entering the nucleus where they act (Vert and Chory, 2006; Gampala et al., 2007). BZR1 and BZR2 are dephosphorylated under treatment with BR (He et al., 2002).
et al. (2011) showed that phosphorylated BKI1 is able to bind 14-3-3 proteins and competes with BZR2 for the interaction. It is possible that this acts as another layer of BZR activation upon BR perception by BRI1.

BIN2 is dephosphorylated upon BR treatment by phosphatase BSU1. The dephosphorylated BIN2 is then subject to proteasomal degradation (Mora-García et al., 2004; Kim et al., 2009). Over expression of BSU1 is able to rescue the bri1 phenotype whilst knockdown of BSU1 causes a severe dwarf phenotype (Mora-García et al., 2004). BSU1 is located mostly in the nucleus but also at the plasma membrane in the absence of BR (Mora-García et al., 2004; Kim et al., 2009). It has been shown that BSU1 interacts with the BSKs at the plasma membrane (Kim et al., 2009). Recently constitutive differential growth 1 (CDG1), an RLCK VII-subfamily member, has been shown to interact with BRI1 and BSU1 suggesting multiple levels of BSU1 regulation (Muto et al., 2004; Kim et al., 2011). The BR signalling pathway is believed to be one of the more completely understood plant signalling pathways and has been used as the basis of understanding other plant signalling pathways (fig. 1.3.1).
Figure 1.3.1. BR and flagellin signalling pathways

A schematic showing the BR (left), flagellin (middle) and nodulation (right) signalling pathways upon activation by BR, flg22 or NF. Complexes of BAK1 with either BRI1 or FLS2 at the plasma membrane (PM) activate RLCKs BSKs/CDG1 or BIK1. In the absence of BR, BIN2 phosphorylates BZR1/2 which is targeted for degradation by interacting with protein 14-3-3. In flagellin signalling, BIK1 interacts with RbohD producing a ROS burst. MAP kinases carry the flagellin triggered signal through the cytoplasm. Perception of NF is at the cell surface by a receptor complex possibly consisting of NFP, LYK3 and DMI2. Activation of DMI1 at the nuclear membrane lead to calcium spiking in the nucleus which is decoded by the DMI3/IPD3 complex and activates symbiosis specific transcription factors. All 3 pathways lead to regulation of gene expression.
1.3.2 The flagellin defence signalling pathway

Plant pathogen recognition is controlled on several levels by RLKs both at the plasma membrane and in the cytoplasm. Pathogens are recognised by molecules that are essential to the life of the pathogen known as microbe-associated molecular patterns (MAMPs) or pathogen-associated molecular patterns (PAMPs) such as bacterial flagellin (Felix et al., 1999), as well as the broader danger-associated molecular patterns (DAMPs) that can be found on more than one type of pathogen such as chitin or peptidoglycan. These patterns trigger defence signalling cascades and gene expression regulation involved in PAMP-triggered immunity (PTI) (Jones and Dangl, 2006; Schwessinger and Ronald, 2012).

The flagellin signalling pathway has been well studied however it is currently not as complete as the BR signalling pathway. The LRR-RLK FLS2 is able to recognise flagellin peptides including flg22, flg15 (in tomato) and a 28-amino acid region in the C-terminal end of the protein flgII-28, as well as full length flagellin (Gómez-Gómez and Boller, 2000; Bauer et al., 2001; Zipfel et al., 2004; Robatzek et al., 2007; Takai et al., 2008; Cai et al., 2011). FLS2 forms a receptor complex at the plasma membrane with BAK1 and the RLCK Botrytis-induced kinase 1 (BIK1) (Veronese et al., 2006; Chinchilla et al., 2007; Lu et al., 2010; Zhang et al., 2010a; Roux et al., 2011; Schwessinger et al., 2011). BIK1 is a PBS-like, RD kinase of the RLCK-VII subfamily (Zhang et al., 2010a). BIK1 and PBL1 (also an RLCK-VII member) have both been shown to be required for full immune responses to Pseudomonas syringae and could act partially redundantly in the signalling pathway (Zhang et al., 2010a). flg22 application decreases the association between BIK1 and FLS2, suggesting that BIK1 dissociates from the complex upon activation (Zhang et al., 2010a). Evidence from Lu et al. (2010) and Schulze et al. (2010) suggests that rapid transphosphorylation occurs between FLS2, BAK1 and BIK1 upon flg22 perception. Mitogen-activated protein (MAP) kinases have been shown to act downstream of the FLS2 receptor complex (Nühse et al., 2000; Asai et al., 2002; Suarez-Rodriguez et al., 2007). However, it is not yet known how they are linked to the receptors (fig.1.3.1).

Recent research by Li et al. (2014) has shown that an NAPDH oxidase RbohD interacts with BIK1 and FLS2 and is activated upon flg22 perception. PTI has been shown to initiate a transient influx of calcium ions into the cytosol followed by a transient burst of ROS leading to the closure of stomata and callose deposition to strengthen the cell wall (Blume et al., 2000; Lecourieux et al., 2002; Nühse et al., 2007; Zhang et al., 2007). This research has given some insight into the mechanism controlling pathogen-activated ROS generation. There has been increasing evidence of cross-talk between the BR and flg22
signalling pathways. For example, BIK1 has been shown to interact with BRI1 and the BSKs with FLS2 (Lin et al., 2013; Shi et al., 2013b). Brassinosteroids are known to inhibit the FLS2 signalling pathway and flg22 inhibits plant growth (Bethke et al., 2009; Albrecht et al., 2012). It is possible that the activity of RLCKs could holistically modulate the activity of signalling pathways within the cell (Laluk et al., 2011).

1.3.3 Comparison of the CSP with BR and flg22 signalling
Parallels can be drawn between these two signalling pathways and the signalling cascade downstream of the symbiosis RLKs. In the BR and flagellin signalling pathways RLCKs transfer the signal from receptor complexes at the cell surface onto downstream signalling components that ultimately leads to transcriptional changes. It is likely that the receptors NFP, LYK3, and DMI2 may sit in complexes at the plasma membrane along with cytoplasmic components such as a remorin (SYMREM1) and an E3 ubiquitin ligase (PUB1) (Lefebvre et al., 2010; Mbengue et al., 2010; Tóth et al., 2012). It is possible that MAP kinases perform signal transduction from the receptors to the nucleus (Chen et al., 2012). It is also conceivable that RLCKs may act as the next step in the signal relay between the RLKs and the MAP kinases. In symbiosis there are known transient influxes of calcium and ROS in response to NF and during nodule development (Miwa et al., 2006; Pauly et al., 2006). It is possible that currently unknown RLCKs may interact with NFP, LYK3 or DMI2 to regulate the ROS machinery as BIK1 does in flg22 signalling (Peleg-Grossman et al., 2012; Li et al., 2014). RLCKs may also play a role in parallel to the CSP, co-ordinating developmental and defence signalling pathways during symbiosis (Peleg-Grossman et al., 2012).

In this study, three RLCK genes were selected using the Medicago Gene Atlas (MtGEA) and in-house microarray data. These genes were investigated for their roles in both RNS and mycorrhizal symbiosis through mutant studies and RNAi. Resources for RLCK protein interaction were also made.
Chapter 2: Materials and methods

2.1. Plant material

*Medicago truncatula* lines used are listed in table 2.1.1. Homozygous *Tnt1* mutants were identified in the lab by PCR. A homozygous EMS mutant was identified by sequencing of PCR products. See appendix I for primers. *Nicotiana benthamiana* plants were cultivated by Horticultural services and used for infiltration from 3 weeks post sewing.

<table>
<thead>
<tr>
<th>Line</th>
<th>Background</th>
<th>Description</th>
<th>Mutation</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>R108</td>
<td>R108</td>
<td>Wild type</td>
<td>WT</td>
<td>Hoffmann <em>et al.</em> (1997)</td>
</tr>
<tr>
<td>A17</td>
<td>A17</td>
<td>Wild type</td>
<td>WT</td>
<td>Van den Bosch and Stacy (2003)</td>
</tr>
<tr>
<td>NF12390 (rck1-1)</td>
<td>R108</td>
<td><em>Tnt1</em> insertion at position 447 (genomic) in <em>RLCK1/SPK1</em></td>
<td><em>Tnt1</em></td>
<td>Samuel Roberts Noble Foundation, USA</td>
</tr>
<tr>
<td>NF91796 (rck1-2)</td>
<td>R108</td>
<td><em>Tnt1</em> insertion at position 621 (genomic) in <em>RLCK1/SPK1</em></td>
<td><em>Tnt1</em></td>
<td>Samuel Roberts Noble Foundation, USA</td>
</tr>
<tr>
<td>NF11296 (rck1-3)</td>
<td>R108</td>
<td><em>Tnt1</em> insertion at position 878 (genomic) in <em>RLCK1/SPK1</em></td>
<td><em>Tnt1</em></td>
<td>Samuel Roberts Noble Foundation, USA</td>
</tr>
<tr>
<td>NF9810 (rck2-1)</td>
<td>R108</td>
<td><em>Tnt1</em> insertion at position 1342 (genomic) in <em>RLCK2</em></td>
<td><em>Tnt1</em></td>
<td>Samuel Roberts Noble Foundation, USA</td>
</tr>
<tr>
<td>NF7569 (rck2-2)</td>
<td>A17</td>
<td>Single base pair mutation C to T at position 1523 (genomic) of <em>RLCK2</em></td>
<td>EMS mutagenesis</td>
<td>RevGenUK, Norwich, UK</td>
</tr>
<tr>
<td>NF5270 (rck3-1/Scooby)</td>
<td>R108</td>
<td><em>Tnt1</em> insertion line. <em>Scooby</em> phenotype.</td>
<td><em>Tnt1</em></td>
<td>Samuel Roberts Noble Foundation, USA</td>
</tr>
</tbody>
</table>

Table 2.1.1. Plant material.

2.1.1 Seed preparation

Mature seed pods were collected by hand from dry *M. truncatula* plants. They were then placed at 37°C for 4-7 days to dry further. Pods were then opened using wooden blocks covered in corrugated rubber to access the seeds. For germination, seeds were scarified using sand paper. The seeds were then treated with 12.5% bleach for 2 minutes and then a series of five 2 minute washes with dH2O, until all the bleach was no longer present – tested using indicator blue tissue. Seeds were then immersed in water to imbibe for at least 1 hour before plating out onto Distilled Water Agar (DWA) plates and inverting. Seeds were placed into the dark at 4°C for 3 days for stratification or 7 to 14 days for vernalisation.
### 2.2 Bacterial and Yeast strains

All bacterial strains used are in table 2.2.1.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Resistance</th>
<th>Species</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>DH5α</td>
<td>-</td>
<td><em>Escherichia coli</em></td>
<td><em>E. coli</em> strain for plasmid amplification</td>
</tr>
<tr>
<td>DB3.1</td>
<td>-</td>
<td><em>E. coli</em></td>
<td><em>E.coli</em> strain resistant to the ccdB cassette</td>
</tr>
<tr>
<td>AR1193</td>
<td>Rif &amp; Carb</td>
<td><em>Agrobacterium rhizogenes</em></td>
<td>For transformation of <em>M. truncatula</em> roots</td>
</tr>
<tr>
<td>GV3101</td>
<td>Rif &amp; Gen</td>
<td><em>Agrobacterium tumefaciens</em></td>
<td>For transformation of <em>N. benthamiana</em></td>
</tr>
<tr>
<td>AH109</td>
<td>-</td>
<td><em>Saccharomyces cerevisiae</em></td>
<td>For yeast 2-hybrid analysis</td>
</tr>
<tr>
<td>Sm1021</td>
<td>Tet</td>
<td><em>Sinorhizobium meliloti</em></td>
<td>Rhizobial symbiont of <em>M. truncatula</em></td>
</tr>
</tbody>
</table>

**Table 2.2.1.** Bacterial and yeast strains.
2.3 Growth Media

The composition of the media used for plant, bacterial and yeast growth are given in table 2.3.1.

<table>
<thead>
<tr>
<th>Media</th>
<th>Composition for 1L</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rhizobium complete medium (TY)</td>
<td>5 g Difco tryptone, 3 g Difco yeast extract, 1.325 g CaCl₂ (containing 15 g agar for solid growth medium).</td>
</tr>
<tr>
<td>Lennox (L)</td>
<td>10 g tryptone, 5 g yeast extract, 5 g NaCl, 1 g D-Glucose (containing 10 g agar for solid growth medium).</td>
</tr>
<tr>
<td>SOC</td>
<td>20 g tryptone, 5 g yeast extract, 0.58 g NaCl, 0.19 g KCl, 2.03 g MgCl₂, 2.46 g MgSO₄·7H₂O, 3.6 g D-Glucose.</td>
</tr>
<tr>
<td>Distilled Water Agar (DWA)</td>
<td>1.5 % w/v Bacto agar, pH 5.7 (adjusted with KOH)</td>
</tr>
<tr>
<td>Fahraeus plant medium (FP)</td>
<td>0.1 g CaCl₂·2H₂O, 0.12 g MgSO₄, 0.01g KHPO₄, 0.150 g NaHPO₄·12H₂O, 5 mg ferric citrate, 2.86 g H₃BO₃, 2.03 g MnSO₄·0.22 g ZnSO₄·7H₂O, 0.08 g CuSO₄·5H₂O, 0.08 g H₂MoO₄·4H₂O, pH 6.3-6.7. For solid medium 0.5% (w/v) LabM No. 1 agar was added.</td>
</tr>
<tr>
<td>Modified FP</td>
<td>FP medium containing 0.5 mM NH₄NO₃</td>
</tr>
<tr>
<td>SD’-LW</td>
<td>1.9g Yeast Nitrogen Base without Ammonium Sulphate and Amino Acids, 5g (NH₄)SO₄, 20g D-Glucose, 20g Formedium agar, 380mg Leucine, 76mg Tryptophan. pH 5.8</td>
</tr>
<tr>
<td>SD’-AHLW</td>
<td>1.9g Yeast Nitrogen Base without Ammonium Sulphate and Amino Acids, 5g (NH₄)SO₄, 20g D-Glucose, 20g Formedium agar, 380mg Leucine, 76mg Tryptophan, 76mg Alanine, 76mg Histidine. pH 5.8</td>
</tr>
<tr>
<td>Medicago mix (compost)</td>
<td>6:6:1 Mix of Levington F2 compost, John Innes No. 2 compost and 4 mm grit.</td>
</tr>
<tr>
<td>50:50 mix Terragreen:Sand</td>
<td>1:1 mix of terragreen (Oil-dry UK ltd, UK) and sharp sand (BB Minerals, UK)</td>
</tr>
</tbody>
</table>

Table 2.3.1. Media used for plant, bacterial and yeast growth.

2.4 Agrobacterium transformation, growth and plant transformation

2.4.1 Agrobacterium rhizogenes ‘Hairy root’ transformation of M. truncatula

Agrobacterium rhizogenes strain AR1193 was transformed with constructs via electroporation at 200Ω and 2.5V. The A. rhizogenes was then shaken at 28°C for 1 hour with SOC medium. The Agrobacterium was plated onto selection plates and left for 3 days at 28°C to grow. Colonies were then grown overnight in liquid culture at 28°C with antibiotic selection and shaking. 600µl were plated out onto selection and left to grown for 3 days on antibiotic selection LB plates to form a thick layer of A. rhizogenes. The
transformed rhizogenes was then used to inoculate 1 day old A17 seedlings. 3mm of root tip were removed from the seedlings using a scalpel, then the seedling dipped into the A. rhizogenes and placed onto square Modified FP (ModFP) plates, poured at a slant and covered in a sterile filter paper. The bottom half of the plates were then covered in black plastic to block out the light and placed in growth chambers at 20°C/15°C day/night temperature with 16 hour days. After 1 week the seedlings were transferred to fresh ModFP plates and the non-transformed roots, identified by fluorescence microscopy, were removed.

2.4.2 Agrobacterium tumefaciens transformation of Nicotiana benthamiana
Agrobacterium tumefaciens strain GV3101 was transformed with constructs via electroporation at 200Ω and 2.5V. The A. tumefaciens was then shaken at 28°C for 1 hour with SOC medium and plated onto antibiotic selection plates and grown for 3 days at 28°C. A colony was grown overnight in liquid culture at 28°C with shaking and antibiotic selection. The liquid culture was spun down and the supernatant removed. The bacteria were resuspended in dH$_2$O. The A. tumefaciens was infiltrated into the leaf by pressure infiltration using a 1ml syringe. dH$_2$O was used as a mock treatment.

2.5 Image capture and microscopy
Light microscopy was performed on either a Nikon Eclipse E800 with a Pixera Pro 600ES camera or a Zeiss Axiophot with a Retiga-2000R Fast 1394 Color camera, QImaging. Fluorescence microscopy was performed on a Zeiss Axiophot with a Retiga-2000R Fast 1394 Color camera, QImaging, or a Leica MZFLIII Fluorescence stereoscope. Confocal microscopy was performed on a Zeiss 780 LSM. Scale bars were added to the images either using the QCapture Pro software (QImaging) or the Image J software.

2.6 DNA amplification, purification and sequencing
All DNA extraction was carried out using a Qiagen DNeasy Plant Kit by Richard Goram, Norwich. All plasmid extraction was carried out using Qiagen Mini-prep spin columns as per manufacturer’s instructions.

2.6.1 Standard Polymerase Chain Reaction (PCR)
GoTaq® Green Master Mix (Promega) was used as the standard Taq polymerase for genotyping and colony PCR following the manufacturer’s instructions for a 25µl reaction volume using cycling parameters 95°C 30s (10 minutes if for colony PCR), [94°C 30s, T$_m$-3°C 30s, 72°C 30s/(kb)x28 - 35, 72°C 2-10 minutes. Phusion High-Fidelity Taq polymerase (New England Biolabs Ltd) was used for cloning PCRs following manufacturer’s instructions for a 20µl or 50µl reaction volume using cycling parameters.
98°C 30s, [98°C 10s, Tm°C 20s, 72°C 1min/kb]x35, 72°C 2-10 minutes. All PCRs were cooled to 10°C after cycling and stored at 4°C or chilled on ice if to be used directly. All PCR amplifications were performed using a G-Storm GS1 thermal cycler (Gene Technologies Ltd). See appendix I for primers.

2.6.2 Restriction Digest
Purified plasmids were digested using restriction enzymes. Digests were performed as per the manufacturer’s instructions (Life Technologies or New England Biolabs), using 1U of enzyme to a 10µl or 20µl reaction with an incubation time of 1 hour at 37°C.

2.6.3 Sequencing
DNA sequencing was carried out by performing the BigDye® reaction with either purified PCR product or plasmid according to manufacturer’s protocol and then completed by The Genome Analysis Centre (TGAC), Norwich or Eurofins (MWG Operon), UK. Purified plasmid of Golden Gate constructs was sent for Next Generation Sequencing by IMGM Laboratory, Germany.

2.7 qRT-PCR
Root material was ground on liquid nitrogen using a pestle and mortar and RNA was extracted using a QIAGEN® RNeasy mini kit. The RLT buffer was pre-warmed to 50°C and the elution volume was reduced to 30µl. RNA was then treated with Turbo DNasease (Life Technologies™) to remove the DNA. The quality of the DNA was tested on a 1% Agarose gel containing ethidium bromide and the quantity was tested on a NanoDrop (Thermo Fisher Scientific). cDNA was synthesized using SuperScript®II Reverse Transcriptase (Life Technologies™). The input RNA was normalised to the lowest sample with a lower threshold of 400ng/total RNA.

qRT-PCR was carried out on a CFX96 Touch™ c1000 thermal cycler (Bio-Rad) using SYBR™ Green JumpStart Taq ReadyMix (without MgCl2) in a 96 well plate. Each well contained the following: 2.6µl MgCl2; 5µl SYBR Green, 2µl diluted cDNA, 1.6µl each 20µM primers to give a total of 10µl. The primer efficiencies were calculated using a serial dilution of control cDNA. The following parameters: 95°C 30s, [94°C 30s, 60°C 30s, 72°C 30s]x49. Melt curve 65°C to 95°C in 0.5°C increments in 5s.

The efficiency of all primer pairs was calculated using a dilution series and linear regression of the resulting Ct data points. Ubiquitin, EF1α and TIP41-like protein were determined to be the most stable references. Normalized relative quantities were calculated using the qBase model (Hellemans et al., 2007), which allows for multiple housekeeping genes and primer specific efficiencies. Values based on 3 technical reps
per sample. The expression was then calculated relative to the control. Standard error was calculated as relative standard error to the control. See Appendix I for primers.

2.8 Cloning

2.8.1 Gateway® Cloning, Invitrogen
The desired gene or promoter region was amplified via PCR using Phusion High Fidelity Polymerase, New England Biolabs. The primers were designed to contain the attB site required for the Gateway® BP, for recombination of the PCR fragment into pDONR207 (Karimi et al., 2002). The donor plasmid was then propagated in *E. coli* strain DH5α and correct colonies were identified by PCR. A Gateway® LR reaction facilitates recombination from the donor vector to the destination vector. The destination plasmid was then propagated in *E. coli* strain DH5α and potentially correct colonies were identified by PCR and verified by sequencing.

2.8.2 TOPO® Cloning, Life technologies
The desired gene or promoter region was amplified via PCR using Phusion High-fidelity Polymerase, New England Biolabs. The primers were designed to contain a CACC 5’ extension compatible with TOPO® cloning, Life Technologies. The PCR fragments were cloned into pENTR-D entry vector following the TOPO manual; the only difference was the incubation at room temperature of the vector, the insert and the clonase was overnight rather than the recommended 5 to 30 minutes. The inserts in the pENTR-D entry vector were then transferred to the compatible Gateway® vector using the LR reaction as described above.

2.8.3 Golden Gate Cloning
Golden Gate cloning was carried out as described in Engler et al. (2008) and Weber et al. (2011), using compatible sites with the Engineering Nitrogen Symbiosis for Africa (ENSA) project, see below (Oldroyd, communications). Level 0 modules were synthesised by Life Technologies GeneArt® Gene Synthesis. Level 1 modules and level 2 constructs were assembled in the lab. Level 0 modules and Level 1 End linkers are Spec resistant. All Level 1 modules, except the End linkers, are Amp resistant. Level 2 constructs are Kanamycin resistant. After assembly each level was propagated in *E. coli* strain DH5α.
Level 1 modules were checked by colony PCR before being sequenced. Level 2 constructs were checked by restriction digest before being sent for next generation sequencing.
2.9 Nodulation Phenotyping

*Medicago truncatula* seeds were germinated on DWA and then planted in a 50:50 mix terragreen and sand. Plants were left to grow in the mix for one week before inoculation. *Sinorhizobia meliloti* 1021 LacZ was grown overnight in 10ml TY plus Tetracycline to an OD600 absorbance between 0.2 and 0.5. The rhizobia were diluted to an OD600 absorbance between 0.05 and 0.001 with water. 2ml of this diluted rhizobia was pipetted at the base of each plant. After 21 days the plants were removed from the soil, washed and the numbers of white and pink nodules per plant were counted.

2.9.1 Histochemical staining procedure

Root material was submerged in 2.5% glutaraldehyde and placed under a vacuum for 15 minutes. The glutaraldehyde was removed and fresh glutaraldehyde was added before leaving at room temperature for a minimum of 1 hour. The fixed material was then washed with Z buffer with one 5 minute wash followed by a 1 hour wash. Fresh LacZ staining solution was made according to the following: 1mM 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-gal) or 5-bromo-6-chloro-3-indolyl-β-D-galactopyranoside (Magenta-gal); 0.1M sodium phosphate; 10mM Potassium chloride; 1mM Magnesium Sulphate; 5mM Potassium Ferricyanide; 5mM Potassium Ferrocyanide.30 to 40μm longitudinal sections of the nodule were made on a Vibratome.

2.10 Mycorrhizal Phenotyping

2.10.1 Growth media and inoculation

Several sources of AMF inoculum were used in the course of this study. Seedlings were planted directly into one of the following inoculums.

**Chive inoculum:** *Rhizophagus irregularis* was grown in a 50:50 mix with chive plants. Inoculum was made by chopping up the root mass and soil containing *R. irregularis*. A 50:50 mix of terragreen and sand was made and then mixed with the chive root inoculum at a ratio of 1:20 inoculum : sand mix.

**PlantWorks inoculum:** A commercial inoculum was obtained from PlantWorks, UK, and mixed with the 50:50 sand mix at a ratio of 1:10 inoculum : sand mix.

**Symplanta inoculum:** Another commercial inoculum was obtained from Symplanta, Munich, 10,000 spores/ml of *Rhizophagus irregularis* (syn: *Glomus irregulare*) SYMPLANTA-001 research grade. For a more concentrated inoculum, 750 spores/plant was use. For a less concentrated inoculum, 500 spores/plant was used.

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For mutant colonisation assays extra R108 WT plants were included in each experiment and level of WT colonisation was checked weekly from 5 weeks post-inoculation until a minimum level of colonisation of 50% was achieved. Once sufficient WT colonisation was achieved all plants were then ready for staining and scoring.

2.10.2 Mycorrhizal staining and scoring

Root material was first washed in fresh 10% KOH at 95°C for 15 minutes, then rinsed in water before being transferred to a 95°C ink and acetic acid solution (5% ink, 5% acetic acid) for 6 minutes. The roots were then left in water over night to de-stain. Colonisation of the root by AMF was scored using the grid method as described by Giovannetti and Mosse (1980).

Alternatively, the AMF was stained using WGA-AlexaFluor549. Roots were placed in 50% ethanol overnight before being transferred to 20%KOH for 2-3 days at room temperature. Roots were rinsed with dH₂O and submerged in 0.1M HCl for 1-2 hours. After another rinse with dH₂O, and again with phosphate buffered saline (PBS) solution, the roots were submerged in PBS/WGA-AF549 (final concentration of 0.2µg/ml) and kept in the dark at room temperature overnight.

2.11 Gene silencing – RNA interference

2.11.1 Constructs

The double knockdown construct was obtained from Myriam Charpentier. It was designed against the RLCK1 CDS and subsequently shown to knockdown both RLCK1 and RLCK2. The vector used was pK7GW|WG2D(II)R modified from pK7GW|WG2D(II) (Karimi et al., 2002; Capoen et al., 2011).

The RLCK2 knockdown construct was designed against the RLCK2 5’ UTR region. It covers 161bp upstream of the start codon. It was cloned into pENTR-D TOPO entry vector using the CACC overhang on the forward primer. It was then cloned into the destination vector pK7GW|WG2D(II)R using GATEWAY® LR reaction, Invitrogen. See Appendix I for sequencing primers.

2.11.2 Transformation of RNAi constructs

A. rhizogenes transformation was carried out as described in 2.4.1. Two weeks after the plate transfer, the plants were then transferred to sterilised 50:50 mix in small greenhouse boxes (Sigma) to keep the humidity high. The lid was slowly removed over several days to reduce the humidity without shocking the plants.
2.11.3 Rhizobial Inoculation and scoring

After one week in the pots the plants were inoculated with Sm1021 as described above for nodulation phenotyping. Plants were checked for nodules at 21dpi. Only transformed roots displaying the dsRED transformation marker were included in the count. The transformed roots were then removed from the plants, frozen using liquid nitrogen and stored at -80°C for qRT-PCR.

2.12 Gene expression analysis

2.12.1 In silico gene expression analysis (MtGEA)

In silico gene expression analysis was carried out using the Medicago Gene Expression Atlas (MtGEA) Database (Benedito et al., 2008; He et al., 2009), Samuel Roberts Nobel Foundation. Probesets were identified using the BLAST function using the CDS sequence.

2.12.2 Promoter-GUS analysis

RLCK1 promoter:GUS construct (pRLCK1:GUS) was obtained from Nicolas Pauly, INRA France (Andrio et al., 2013). The RLCK2 promoter:GUS (pRLCK2:GUS) construct was made by cloning the 2328bp region upstream of the RLCK2 start codon into pDONR207. The promoter region was amplified by PCR and confirmed by sequencing. The primers contained overhangs compatible for the Gateway® BP reaction. An LR reaction was then performed to clone the promoter region into the pKGWFS2 promoter:GUS vector. The Lotus japonicus Ubiquitin1 (LjUB1) promoter:GUS construct was used as a positive control for the staining. See Appendix I for primers.

Plants were transformed with the constructs using A. rhizogenes transformation. Plants were inoculated with either LacZ containing Sm1021 and checked at 4, 7, 14d and 21dpi or R. irregularis (chive inoculum) and checked at 5, 10 and 20dpi.

GUS staining solution was freshly made according to the following: 2mM 5-bromo-4-chloro-3-indolyl-β-glucuronic acid (X-GlcA); 10mM Na-EDTA; 1mM potassium ferricyanide; 0.1 M sodium phosphate; 0.1% V/V Triton X-100.

The plant roots were immersed in the GUS staining solution and incubated at 37°C and removed when the level of staining seemed appropriate – approximately 5 hours. The GUS stained roots were then either counterstained with a Wheat Germ Agglutinin (WGA) Alexa Fluor® 594 dye (Life Technologies) for AMF colonisation or fixed and counterstained with magenta-gal for rhizobial colonisation.
2.13 Phylogenetic Analysis
The protein sequences of MtRLCK1 and MtRLCK2 were used as queries for BLASTP searches against the predicted protein sequence databases of the genomes of *Amborella trichopoda, Arabidopsis lyrata, A. thaliana, Brachypodium distachyon, Brassica rapa, Chlamydomonas reinhardtii, Glycine max, L. japonicus, M. truncatula, N. benthamiana, Oryza brachyantha, Oryza glaberrima, Oryza indica, O. sativa, Panicum virgatum, Phaseolus vulgaris, Physcomitrella patens, Populus trichocarpa, Selaginella moellendorffii, Setaria italica, Solanum lycopersicum, Solanum tuberosum, Sorghum bicolor, Triticum aestivum, Vitis vinifera and Zea mays* using Geneious. The top 50 blast hits for each of these genomes were taken forward and aligned with the queries using ClustalW with the default parameters. The resultant alignment was used to construct trees using the Neighbour-joining tree algorithm implemented in Geneious with 1000 bootstrap replicates and default values for the other parameters. From the resulting tree, the RLCK1 and RLCK2 branches were extracted and the proteins were realigned as described above to give the final tree.

2.14 Protein Analysis

2.14.1 *In silico* protein analysis
Proteins were analysed *in silico* using InterProScan4 (Hunter *et al.*, 2012) and the ExPASy Bioinformatics Portal (Artimo *et al.*, 2012) in 2011. In 2014 proteins were analysed *in silico* using InterProScan5 (Jones *et al.*, 2014) and the Calmodulin target database ([http://calcium.uhnres.utoronto.ca/ctdb/ctdb/home.html](http://calcium.uhnres.utoronto.ca/ctdb/ctdb/home.html)).

2.14.2 Protein localisation constructs
Protein localisation constructs were created by PCR fusion of RLCK1 and RLCK2 to mCherry using successive PCR to splice PCR products together. The first PCR (PCR1) amplified *RLCK1* and *RLCK2* CDS with overhanging attB site at the 5’ end and complementary sequence to mCherry at the 3’ end. mCherry was also amplified with a 5’ overhang that was complementary to *RLCK1* or *RLCK2* CDS and 3’ attB. The second PCR (PCR2) takes place in two steps. Step a combines the products from PCR1 as the complementary overhangs act as primers for the polymerase to create the fused product. Step b has the addition of 5’ and 3’ primers for the full length fused product *RLCK1:mCherry or RLCK2:mCherry*. See appendix I for primers.

PCR1: Composition: 0.05U Phusion Taq, 1x GC buffer, 1µM dNTP, 0.25µM Forward Primer, 0.25µM Reverse Primer, 50-100ng Template DNA. Cycling parameters: 98°C 30s, [98°C 10s, 55°C 20s, 72°C 2m]x35, 72°C 4m.
**PCR2a:** Composition: 0.05U Phusion Taq, 1x GC buffer, 0.8µM dNTP, 50-100ng Gene CDS, 50-1—ng mCherry CDS. Cycling parameters: 98°C 30s, [98°C 10s, 50°C 20s, 72°C 2m30s]x10, 72°C 5m.

**PCR2b:** Add to PCR2a reaction tubes: 0.1µM Forward Primer and 0.1µM Reverse Primer. Cycling parameters: 98°C 30s, [98°C 10s, 60°C 20s, 72°C 5m]x28, 72°C 10m.

The fused fragments were cloned into pDONR207 by a BP reaction. They were then put into the pUB-GW-GFP via an LR reaction (Maekawa et al., 2008). Constructs were checked by sequencing, see appendix I for primers. Constructs were transformed into *N. benthamiana* leaves by *A. tumefaciens* infiltration and localisation was checked in leaf discs using a confocal microscope at 1, 2 and 3dpi.

### 2.14.3 Yeast 2-Hybrid

*RLCK1* and *RLCK2* CDS were amplified using Phusion High-Fidelity Taq polymerase and cloned into pENTR-D using TOPO® cloning. The CDS was then transferred by LR reaction into pGADT7 and pGBK7 Gateway® vectors for expression of RLCK1 and RLCK2 proteins as bait and prey respectively. AH109 yeast strain was transformed using the lithium acetate method (Daniel Gietz and Woods, 2002). The yeast was grown at 28°C for 3 days on SD⁺ agar minus leucine and tryptophan (SD⁺-LW) or SD⁺ agar minus alanine, histidine, leucine, and tryptophan (SD⁺-AHLW) for interaction selection.

### 2.15 Co-Immunoprecipitation

Cloning was performed using the Golden Gate system to make Level 2 constructs encoding myc-tagged RLCK1 or RLCK2 and GFP-tagged MtNFP, MtLYK3, MtHMGR1 and MtDMI2. See Appendix I for construct details and plasmid maps. Constructs were transformed into *N. benthamiana* leaves by *A. tumefaciens* infiltration.

#### 2.15.1 Protein Extraction

Leaf discs were taken at 1, 2 and 3dpi, frozen in liquid nitrogen and stored in a -80°C freezer. Samples were ground either using a pestle and mortar chilled using liquid nitrogen, or a tissue lyser using 3mm tungsten carbide beads (Qiagen). The protein was extracted by one of two methods, the first added warm buffer (65°C) to the ground sample [100mM Tris-HCl pH 7.4, 150mM NaCl, 5mM EDTA, 10mM DTT 5% SDS, 4M Urea + Roche protease inhibitor cocktail], incubated at 90°C for 10 minutes, centrifuge maximum speed for 2 minutes, collect supernatant and mix with SDS sample buffer. The second method was as described by (Waadt et al., 2008). Protein extract produced by either method was mixed with SDS sample buffer 4x Laemmli sample buffer (Bio-Rad), heated
at 90°C for 5 minutes, loaded into a 10x SDS-PAGE Gel and ran at 150V for 1 hour 25 minutes.

2.15.2 Western Blot
Proteins were transferred to the membrane at 100V for ~1 hour. The membrane was blocked using 5% milk in TBS-T [10mM Tris-HCl pH7.4, 150mM NaCl, 0.1% Tween 20] for 1 hour. The membrane was washed in TBS-T and incubated overnight at 4°C with the primary antibody. The membrane was washed in TBS-T before incubation with the secondary antibody for 1 hour at room temperature. The membrane was washed in TBS-T 3 times for 10 minutes before treating with enhanced chemiluminescence (ECL) solution and exposing to an X-ray film for times between 30 seconds and 30 minutes. Total protein levels loaded were determined by Coomassie staining of protein gels or by Ponceau staining of the membranes used in the blot.
Chapter 3: Characterisation and expression of Receptor-Like Cytoplasmic Kinase 1 and 2

3.1 Introduction

*Medicago truncatula* (alongside *Lotus japonicus* and *Glycine max*) is one of the model plants used to study the legume symbioses with rhizobia and AMF. Transformation of *Medicago* by floral dip transformation as previously performed in Arabidopsis is not possible on a large scale (Bechtold and Pelletier, 1998). T-DNA mutagenesis is not suitable for generating a large mutant collection due to the relative difficulty in transforming *M. truncatula* compared to *Arabidopsis*. D'Erfurth *et al.* (2003) used the long terminal repeats (LTR) retrotransposon *Tnt1* from tobacco (*Nicotiana tabacum*) (*Grandbastien et al.*, 1989) to transform the *M. truncatula* R108 ecotype. They showed that during regeneration from tissue culture the *Tnt1* retrotransposon transposes at a higher frequency into transcriptionally active euchromatin than non-coding euchromatin regions and that they are stable during the life cycle of the plant. The retrotransposon can also be re-activated by tissue culture to generate more mutations. The *Tnt1* retrotransposon uses a “copy and paste” system leaving the transposon in the original position whilst a copy of the transposon is inserted somewhere else in the genome, usually far away from the original. *Tnt1* is 5.3 kb long and creates a 5 bp duplication at each end upon insertion. Each *Tnt1* line is estimated to contain between 6 and 59 new insertions (Tadege *et al.*, 2008). This means that in the case of reverse genetic studies, within a single line there can be many insertions additional to the one in the gene of interest. Hemizygous insertions that arise in the initial generation (R0) segregate in subsequent generations (R1, R2 etc.) and so can be randomly lost or fixed as homozygous as the material is advanced, resulting in genotypically distinct sibling lines at later generations.

At the Samuel Roberts Noble Foundation a *Tnt1* insertion mutant collection has been generated that in 2011 was estimated to cover 85% of the *M. truncatula* genome (D'Erfurth *et al.*, 2003; Tadege *et al.*, 2005; Tadege *et al.*, 2008; Cheng *et al.*, 2011). A Flanking Sequence Tag (FST) database, which was generated by TAIL-PCR showing the sequences surrounding *Tnt1* insertions in each line, can be BLAST searched using the genomic sequence of a gene of interest to find useful *Tnt1* lines. The *Tnt1* collection can also be reverse screened via PCR using gene-specific primers if an FST cannot be found. In this way, this collection can be used to identify mutant lines segregating for a desired mutation.
Another extremely useful resource open to the Medicago research community is the Medicago Gene Expression Atlas (MtGEA). It is compiled by the Samuel Roberts Noble Foundation and collates data from Affymetrix Medicago Gene Chip® experiments for M. truncatula under a variety of biological and chemical conditions (Benedito et al., 2008; He et al., 2009; Hogekamp et al., 2011; Czaja et al., 2012; Gaude et al., 2012; Seabra et al., 2012). Based on interesting expression profiles that show gene expression during, and induced by, the presence of symbiotic partners, and a high sequence homology to each other (76 % at the nucleotide level and 69 % at the amino acid level), two kinase genes were selected for investigation. Based on their lack of transmembrane domains and close homology to a member of the Rice Receptor-like cytoplasmic Kinase Family XV (Jung et al., 2010) these genes were named Receptor-like Cytoplasmic Kinase 1 (RLCK1) and RLCK2.

RLCKs have been shown to interact with membrane bound receptor kinases and phosphorylate downstream signalling components in the PAMP-triggered immunity and brassinosteroid signalling pathways (Tang et al., 2008; Kim et al., 2009; Lu et al., 2010; Zhang et al., 2010a; Shi et al., 2013a). There are several RLKs required for symbiosis and currently for the symbiosis signalling pathways we do not know how the signal travels from the plasma membrane to the nuclear envelope to activate DMI1. It is possible that RLCKs are part of the signal relay.

During the course of this study, Damiani et al. (2012) published that in M. truncatula expression of RLCK1 was in nodule zones I and II as part of their study into the similarities between the development of a nodule and a pathogenic nematode gall. Andrio et al. (2013) later identified RLCK1 as part of their transcriptome analysis investigating the role of reactive oxygen species (ROS) in nodulation, naming it MtSpk1 (Symbiotic Protein Kinase 1). They showed that RLCK1 was upregulated both in response to NF and hydrogen peroxide treatment. Using promoter-GUS analysis they showed that RLCK1 was expressed throughout rhizobial infection, early nodule development and in the nodule infection zone (Zone II) confirming the earlier study (Damiani et al., 2012). Andrio et al. (2013) used an artificial micro-RNA (amiRNA) to reduce the gene expression of RLCK1 to an average of 40% of the control expression and was able to see a significant reduction of 2 nodules/plant on the knock-down plant roots. They also had a preliminary look at the expression of known symbiotic genes in the amiRNA roots and saw a reduction in the expression of transcription factors MtNIN and NF-YC1 (formerly MtHap2.1). However, Andrio et al. (2013) did not have any stable mutants for RLCK1 and RNAi silencing can be subject to off-target gene silencing. I discuss the limitations of RNAi in more detail in Chapter 4. They also did not look at RLCK2 which has a 76% nucleotide sequence
homology with \textit{RLCK1}. Here I show that my work on the RLCKs is complimentary to and expands upon the work already published using stable mutants to characterise the mutant phenotypes.

3.2 Identification of Mutants for \textit{RLCK1} and \textit{RLCK2}

Using the tools available on the Medicago Insertion Database FSTs were identified for insertions in the second intron (447 bp genomic), third exon (621 bp genomic) and third intron (878 bp genomic) of \textit{RLCK1} in Tnt1 insertion lines NF12390 (\textit{rlck1-1}), NF10796 (\textit{rlck1-2}) and NF11296 (\textit{rlck1-3}) respectively. R1 generation seeds were obtained from the Samuel Roberts Noble Foundation and homozygote mutants for each insertion were identified using gene specific primers #1 and #3 and primers #1 and #10 for Tnt1 identification (see appendix I) (fig. 3.2.1a). \textit{rlck1-3} produced a very low number of viable seed and so was not studied further. The expression of \textit{RLCK1} in the \textit{rlck1-1} and \textit{rlck1-2} alleles was tested using semi-quantitative PCR on cDNA generated from 21dpi nodulated root tissue (fig. 3.2.1b). \textit{RLCK1} expression was detected in \textit{rlck1-1} but not in \textit{rlck1-2} at 28 cycles. The amplified cDNA band for \textit{RLCK1} in \textit{rlck1-1} was sequenced and was not different from WT (R108). This allele was not used for further investigation.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure321.jpg}
\caption{Allele identification for \textit{RLCK1}.}
\end{figure}

(a) \textit{RLCK1} comprises of 4 exons (blue) and 3 introns. Tnt1 insertion alleles at nucleotides 447 (\textit{rlck1-1}), 621 (\textit{rlck1-2}), 878 (\textit{rlck1-3}) of the genomic sequence. Gene specific (solid arrows) and Tnt1 specific (dashed arrows) primer positions shown beneath the gene structure. (b) Semi-quantitative RT-PCR of \textit{RLCK1} and \textit{EF1\alpha} (positive control) in 21 dpi nodulated root. \textit{RLCK1} band is 1122bp. \textit{EF1\alpha} is a 100 bp fragment. 28 cycles.
For RLCK2, an FST was identified in the fourth exon (1342 bp genomic) in line NF5270 and the allele was designated rck2-1. Homozygous mutants for rck2-1 were identified by PCR using the gene-specific primers #4 and 6 and primers #4 and #10 for the Tnt1 identification (See Appendix I). A second allele, designated rck2-2, was identified by TILLING (RevGen UK, Norwich). The mutation is a C to T change at base pair 1407 (genomic) causing a Q to STOP change at amino acid 269, creating a premature stop codon at the protein level (fig. 3.2.2). RLCK2 expression was tested using semi-quantitative PCR on cDNA generated from 21dpi nodulated root tissue (fig. 3.2.2b). Some rck2-1 individuals had some detectable expression at 28 cycles, but the band detected contained a 147bp deletion (478-625) within the cDNA which was confirmed by sequencing of the cDNA. This could occur by the splicing out of the Tnt1 transcript from the rck2-1 mRNA removing some of the RLCK2 sequence in the process. The resulting protein from the mis-spliced mRNA would have a missense amino acid sequence resulting in a missing RD domain and a premature stop codon that would remove the majority of the kinase domain. rck2-2 still has detectable levels of RLCK2 expression at 28 cycles, however the C to T mutation was confirmed by sequencing at base pair 805 in the cDNA. rck2-1 is in the R108 background and rck2-2 is in the A7 background.

**Figure 3.2.2.** Allele identification for RLCK2.

(a) RLCK2 comprises of 4 exons (blue) and 3 introns. rck2-1 is an exonic Tnt1 insertion at 1342 bp (genomic). rck2-2 is a single base pair mutation at 1407 bp (genomic) causing a premature stop codon. Gene specific (solid arrows) and Tnt1 specific (dashed arrows) primer positions shown beneath the gene structure. (b) Semi-quantitative RT-PCR for RLCK2 and EF1α in A17 and rck2 mutants in 21 dpi nodulated root. rck2-1 bands (963 bp) are 147 bp smaller than WT (1110 bp). rck2-2 has a full length cDNA and WT expression but contains a premature stop codon. EF1α is a 100 bp fragment. 28 cycles.
3.3 Nodulation phenotypes of *rlck1* and *rlck2*

3.3.1 Rhizobial Infection

To score the numbers of infection events *rlck1* and *rlck2* mutant seedlings alongside WT R108 and A17 were inoculated with LacZ-expressing Sm1021. Plants were scored for the number of microcolonies, infection threads (ITs) and nodule primordia at 6 dpi. *rlck1-2* was found to have a significantly higher number of microcolonies than WT (R108) (p=0.029 and p=1.59x10^-6 respectively) (fig. 3.3.1.1). The infection threads in the *rlck1-2* mutants sometimes appeared to be thicker than WT (fig. 3.3.1.2a,c,e) but progressed to the nodule primordia as normal (fig. 3.3.1.2f).

The *rlck 2-1* mutant is not significantly different from WT (R108) for any of the infection events (fig. 3.3.1.1) and the infection threads and nodule primordia look normal (fig. 3.3.1.3a). However, *rlck2-2* had significantly fewer infection events at all stages compared to WT (A17) – microcolonies (p=3.53x10^-5), ITs (p=2.19x10^-6) and fewer nodule primordia (p=7.38x10^-7) (fig. 3.3.1.1). The root hairs on the *rlck2-2* mutant appeared shorter and infection threads (where viewed) appeared thicker and misshapen compared to WT (A17) (fig. 3.3.1.3b).

![Figure 3.3.1.1. The number of infection events per plant in the *rlck* mutants and WT.](image)

Plants were harvested at 6dpi Infections were visualised by LacZ-expressing *S. meliloti* 1021 (Sm1021). The subscript next to each allele refers to the background of the allele. A 2-tailed t-test was used to compare means between each allele and their respective control. The data is an average of 16 to 20 individual plants with the bars representing standard error. *p<0.05 ***p<0.001
Figure 3.3.1.2. Differences between rick1 and WT (R108) infection threads.

The infection threads in rick1-2 (a) appear thicker and less regular in shape than WT (c). The rick1-2 mutants are also hyper-infected, infection threads are indicated by arrowheads (b) compared to WT (R108) (c). Infection threads in the rick1-2 mutants are able to progress to the nodule primordia (d). Infections were visualised by using LacZ-expressing S. meliloti 1021. Scale bars in (a) is 50µm, all other scale bars are 100 µm. Images taken at 6 dpi.
Figure 3.3.1.3. Differences between *rlck2* and WT (R108 and A17) infection threads.

The infection threads (arrows) in *rlck2*-1 mutant (a) are normal compared to WT (R108) (c), however the *rlck2*-2 mutant (b) has shorter root hairs and deformed infections compared to their WT (A17) control (d). Very few threads at 6 dpi progressed to nodule primordia in the *rlck2*-2 mutant. Infections were visualised by using *LacZ*-expressing *S. meliloti*. Scale bars in (a) and (c) are 100 µm, scale bars in (b) and (d) are 50 µm. Images taken at 6 dpi.
3.3.2 Nodule formation

*rlck1-2* and *rlck2* mutants were tested for their ability for form nodules at 21 dpi with *S. meliloti* strain Sm1021 using R108 and A17 as controls. *rlck1-2* had significantly fewer nodules than WT (R108) (p=0.038) with a specific reduction in the number of pink nodules (p=0.015) (fig. 3.3.2.1a). *rlck2-1* had significantly more nodules than WT (R108) (p=0.048) with more white nodules (p=0.044). *rlck2-2* had significantly fewer nodules than WT (A17) (p=6.11x10^{-5}) with significantly fewer pink nodules (p=3.22x10^{-6}) (fig. 3.3.2.1b).

To more closely examine the symbiotic phenotype, mature pink nodules were sectioned. Nodule sections showed that in *rlck1-2* mutants the bacteria are able to be released into the cortical cells of the nodule. In younger nodules the infection threads within the infection zone (Zone 2) are clearly visible in both the *rlck1-2* mutants and in WT (R108) (fig. 3.3.2.2 a-d). However, as the nodule matures, this zone becomes diminished in the WT (R108) nodules but stays prominent in *rlck1-2* with ITs seeming to reach into zone I (fig. 3.3.2.2e-f). This, along with the hyper-infected phenotype of the *rlck1-2* mutant at the early stages of nodulation, suggests a role for RLCK1 during infection.

Nodule sections of *rlck2* mutants show that the bacteria were able to be released into the cortical cells of the nodules. *rlck2-1* nodules look as WT (R108) (fig. 3.3.2.3a-d). At 28 dpi *rlck2-2* had only produced small nodules compared to WT (A17) but zones I, II and II-III were visible and normal in appearance (fig. 3.3.2.3e-f).
Figure 3.3.2.1. Nodulation in WT and rick mutants.

(a) Number of nodules for the rick1 mutant alleles and WT (R108). The data are averages of 13 to 19 individual plants with error bars displaying the standard error. (b) Number of nodules for the rick2 mutant alleles and respective control WT (R108 and A17). A 2-tailed t-test was used to compare the means of each allele to their corresponding control. The data are averages of 12 - 20 plants. Bars display standard error. *p<0.05.
Figure 3.3.2.2. Nodule sections of rick1 mutants and R108 WT at different developmental stages.

In young nodules infection threads (arrows) can be seen in the apex of the nodule in both R108 WT (a and c) and in the rick1-2 mutant (b and d). However the infection threads can still be seen in mature rick1-2 mutant nodules (f) but not in the mature nodules of R108 WT (e). Images (g) and (h) are computationally magnified from images (e) and (f) respectively. Infections were visualised by using LacZ-expressing S. meliloti. Sections were taken at 28 dpi. 3 nodules sectioned per genotype. Scale bars are 100 µm.
Figure 3.3.2.3 Nodule sections of *rick2* mutants and WT (R108 and A17).

*rick2*-1 nodules (b and d) appear normal compared to WT (R108) (a and c). *rick2*-2 only produced small nodules (f and h) compared to WT (A17) (e and g) but these were colonised normally. Infections were visualised by using *LacZ*-expressing *S. meliloti* 1021. Sections were taken at 28 dpi. Scale bars are 100 µm.
3.4 Mycorrhizal phenotyping of *rlck1* and *rlck2*

*rlck1* and *rlck2* mutants were tested for their ability to be colonised by AMF *Rhizophagus irregularis* alongside WT R108 and A17. Colonisation was scored using the grid method (Giovannetti and Mosse, 1980) once WT colonisation was more than or equal to 50% (see materials and methods).

*rlck1-2* was found to be reduced in mycorrhizal colonisation compared to WT (R108). *rlck1-2* mutants show a significant reduction in internal hyphae (p=4.89x10^-5), vesicles (p=5.12x10^-7) and arbuscules (p=7.38x10^-7) (fig. 3.4.1a). Arbuscules in *rlck1-2* were small and did not fill the plant inner cortical cell as in WT (R108) (fig. 3.4.2).

*rlck2-1* and *rlck2-2* were also reduced in mycorrhizal colonisation compared to their respective WT controls (R108 and A17). The *rlck2* mutants were significantly reduced in internal hyphae (p=3.16x10^-8 and p=0.006, *rlck2-1* and *rlck2-2* respectively), vesicles (p=1.07x10^-5 and p=0.004, *rlck2-1* and *rlck2-2* respectively) and arbuscules (p=1.62x10^-7 and p=0.009, *rlck2-1* and *rlck2-2* respectively) (fig. 3.4.1b). Arbuscules in *rlck2-1* appear smaller and do not fill the plant inner cortical cells as in WT (R108). However, in *rlck2-2* arbuscules, although fewer in number, fill the plant inner cortical cell as seen in WT (A17) (fig. 3.4.3).
Figure 3.4.1. AMF (*R. irregularis*) colonisation is reduced in *rick* mutants.

(a) Percentage AMF colonisation for the *rick*1 mutant alleles and WT (R108). The data are averages of 18 – 25 individual plants. (b) Percentage of AMF colonisation of the *rick*2 alleles and WT (R108 and A17). The data are averages of 13- 25 plants. Plants were scored when WT (R108) plants reached ≥50% colonisation (see materials and methods). Bars display standard error. A 2-tailed t-test was used to compare the means of each allele to their corresponding control. **p<0.01 ***p<0.001
Figure 3.4.2. AMF colonisation phenotypes of rlck1 mutants and WT (R108). R108 (a, b) and rlck1-2 (c, d) mutant roots colonised by R. irregularis. Arbuscules in rlck1-2 were smaller, and did not fill the inner cortical cells (d). AMF is stained by ink. Phenotypes were scored when WT (R108) was colonised >50% by AMF. Scale bars on (a) and (c) are 100 µm. Scale bars on (b) and (d) are 25µm.
Figure 3.4.3. AMF colonisation phenotypes of *rick2* mutants and WT plants. 
(a,b) R108 and A17 (c,d) WT and (e,f) *rick2*-1 and (g,h) *rick2*-2 mutant roots colonised by *R. irregularis*. Arbuscules in *rick2*-1 did not fill the plant inner cortical cell as occurred in WT (R108). Arbuscules appeared normal in *rick2*-2 compared to WT (A17). AMF is stained by ink. Plants were scored when WT (R108 and A17) were colonised >50 % by AMF. Scale bars on (a), (c), (e) and (g) are 100 µm. Scale bars on (b), (d), (f) and (h) are 50 µm.
3.5 Expression of **RLCK1** and **RLCK2**

3.5.1 Expression of **RLCK1** and **RLCK2** by Microarray

The Medicago Gene Expression Atlas (MtGEA) was used to survey expression data for **RLCK1** and **RLCK2** under varying biological and chemical conditions. **RLCK1** expression is recorded via probe set Mtr.16214.1.S1_at (fig. 3.5.1.1). **RLCK1** expression is upregulated just 6 hours after inoculation with Nod Factors and AM “Myc Factors” S-LCOs or NS-LCOs (unpublished data). **RLCK1** expression is also high in young nodules (3, 4 and 6 dpi) and during AM infection. This upregulation at the early stages of both symbioses supports the *rlck1* phenotype seen during infection processes. Expression data acquired using laser capture of the AM colonised roots (Hogekamp et al., 2011) shows that **RLCK1** expression is higher in the cortical cells adjacent to the arbusculated cortical cells than in the cells that contain arbuscules. These adjacent cells are potentially preparing for AM colonisation which fits with the infection related expression of **RLCK1** seen at earlier time-points. Unpublished microarray data show that **RLCK1** is upregulated in root hairs of rhizobially infected plant at 3 and 5 dpi, the former time point corresponding to microcolony formation and the latter to when the infection thread is progressing through the root hair (Murray Lab)(fig. 3.5.1.2). The microarray data also suggests that the expression of **RLCK1** in response to Nod Factor is dependent on **NFP**, **DMI1** and **DMI3**. **RLCK1** expression seems to be lower than WT in the transcription factor mutants **nsp1**, **nsp2**, **ern1** and **nin** at 6 hours post inoculation (hpi) with NF. At 24 hpi with NF **RLCK1** expression is dependant also on **NSP1** and **NSP2**, but not **ERN1** or **NIN** (Oldroyd Lab).

**RLCK2** expression is recorded via probe set Mtr.24207.1.S1_at. **RLCK2** has a constitutive expression in root tissue and is upregulated in response to rhizobia or mycorrhization (fig. 3.5.1.3). **RLCK2** expression is upregulated in young nodules (3, 4 and 6 dpi) and during AM infection. Laser capture of the AM colonised roots (Hogekamp et al., 2011) shows that **RLCK2** is expressed higher in the root cortical cells that contain arbuscules than other cortical cells. Unpublished microarray data (fig. 3.5.1.4) shows that **RLCK2** has some constitutive expression in root hairs and is upregulated at 3 and 5 dpi (Murray Lab). It also shows that **RLCK2** upregulation in response to NF is dependent on **NFP**, **DMI1**, **DMI3**, **NSP2**, **ERN1** and **NIN** (Oldroyd Lab). These differing expression profiles suggest the two RLCKs could have differing but overlapping roles, or be working synergistically, during symbiosis and that the regulation their expression is under different genetic control.
**Figure 3.5.1.1.** Medicago Gene Expression Atlas (MtGEA) data for *RLCK1*.

*RLCK1* expression across a number of different experimental conditions. *RLCK1* is highly expressed during early stages of both symbioses – nodulation (4dpi) and mycorrhizal infection. *RLCK1* is also upregulated quickly after treatment with bacterial and mycorrhizal LCOs (NF and myc-LCOs). Laser capture dissection of AMF colonised roots (Root LCM) show that *RLCK1* expression is higher in plant inner cortical cells neighbouring arbusculated cells than the arbusculated cells.
Figure 3.5.1.2. Unpublished Medicago Gene Expression Atlas data for RLCK1. Affymetrix Medicago Gene Chip® data for RLCK1 in symbiosis mutants and in response to various conditions. RLCK1 is up regulated in root hairs during rhizobial. Early upregulation of RLCK1 in response to NF is dependent on the symbiosis genes NFP, DMI1 and DMI3. RLCK1 is not upregulated in response to P. infestans.
Figure 3.5.1.3 Medicago Gene Expression Atlas (MtGEA) data for **RLCK2**.

**RLCK2** expression in a number of different tissues and experimental conditions. **RLCK2** is constitutively expressed in root tissue. Expression of **RLCK2** is strongly upregulated during both symbioses – nodulation and mycorrhization. **RLCK2** is also upregulated after treatment with bacterial LCOs (Nod factor) but not with mycorrhizal LCOs. Laser capture dissection of AMF colonised roots (Root LCM) show that **RLCK2** expression is higher in arbusculated plant inner cortical cells than neighbouring cells.
Figure 3.5.1.4. Unpublished Medicago Gene Expression Atlas data for RLCK2.

Affymetrix Medicago Gene Chip® data for RLCK2 in symbiosis mutants and in response to various conditions. RLCK2 is up regulated in root hairs during rhizobial infection. Early upregulation of RLCK2 in response to NF is dependent on the symbiosis genes NFP and DMI3. RLCK2 is not upregulated in response to P. infestans.
3.5.2 Symbiotically enhanced expression of \textit{RLCK1} and \textit{RLCK2}

The \textit{MtRLCK1} promoter-GUS construct (pRLCK1:GUS) was obtained from Nicolas Pauly (\textit{SPK1}) (Andrio et al., 2013) and was transformed into \textit{M. truncatula} R108 via \textit{A. rhizogenes}. Roots were checked at 4, 7, 10, and 14 dpi with \textit{LacZ}-expressing Sm1021. The GUS expression was visualised using X-GlcA staining and the bacteria were counterstained using Magenta-Gal. Expression was confirmed for \textit{RLCK1} in infected root hairs, nodule primordia and in developing nodules as previously shown by Andrio et al. (2013). There was some non-symbiotic expression of \textit{RLCK1} in the root tips and lateral root primordia (fig. 3.5.2.1). Roots were also checked at 5, 10, 15 and 20 dpi with \textit{R. irregularis} (Chive inoculum). The GUS expression was visualised using X-GlcA staining and the AMF was visualised by WGA staining. \textit{RLCK1} is expressed in the early stages of mycorrhizal infection with GUS expression visible in cells surrounding AMF penetration. \textit{RLCK1} is expressed at later stages of mycorrhizal infection, with GUS visible in the plant inner cortical cells surrounding AMF progression through the root (fig. 3.5.2.2).

To compare the expression of \textit{RLCK2} to \textit{RLCK1} an \textit{RLCK2} promoter-GUS (pRLCK2:GUS) construct was made by PCR amplifying 2.3kb of the promoter region and cloning via Gateway into vector pKGWFS7 (Karimi et al., 2002). The construct was transformed into \textit{M. truncatula} R108 via \textit{A. rhizogenes}. Roots were checked at 4, 7, 10, and 14 dpi with \textit{LacZ}-expressing Sm1021 and was visualised as described above. \textit{RLCK2} was expressed throughout the root during the early stages of rhizobial symbiosis and at later stages in the developing nodule (fig. 3.5.2.3a-c). Roots were also checked at 5, 10, 15 and 20 dpi with \textit{R. irregularis} (Chive inoculum) as described above. \textit{RLCK2} was expressed throughout the root during the early stages of AM symbiosis and it was strongly expressed in the inner plant cortical cells that contain arbuscules (fig. 3.5.2.3e-f).
Figure 3.5.2.1. pMtRLCK1:GUS expression during rhizobial symbiosis.

The GUS activity (stained in blue) shows that RLCK1 is expressed in infected root hairs and cortical cells (a), and in nodules (b) as well as in root tips (c) and lateral root primordia (d). Scale bars in (a), (c) and (d) are 100 µm and 500 µm in (b).
Figure 3.5.2.2. pRLCK1:GUS expression during mycorrhizal symbiosis.

GUS activity (stained blue) shows that RLCK1 is expressed during the early stages of mycorrhizal infection (a), (c) and (e), and in cortical cells surrounding mycorrhizal progression through the root (b), (d) and (f). The AMF is stained in red with WGA, shown in red in images (c) to (f). Scale bars are 60 µm in (a), (c) and (e), and 200 µm in (b), (d) and (f).
Figure 3.5.2.3. pRLCK2:GUS expression during rhizobial and mycorrhizal symbioses. The GUS expression (shown in blue) shows that *RLCK2* is expressed throughout the root during the early stages of rhizobial (4 dpi) (a) and (b) and mycorrhizal symbiosis (10 dpi) (d). During the later stages of rhizobial symbiosis *RLCK2* expression is can be seen in the nodule (14 dpi) (c). *RLCK2* expression remains in the cortical cells at later stages of mycorrhizal symbiosis and is higher in arbusculated cells (starred) (e) to (g). The AMF is stained with WGA, shown in red in images (f) and (g). Scale bars are 100 µm in (a) and (b), 200 µm in (c) and (d), and 25 µm in (e), (f) and (g).
3.6 Phylogenetic analysis of RLCK1 and RLCK2 proteins

It has been shown that legumes have undergone several Whole Genome Duplication (WGD) events (Cannon et al., 2006; Young et al., 2011); many of these gene duplicates have been retained, and some have been recruited into the legume-rhizobia symbiosis. The high homology between RLCK1 and RLCK2 could be due to the WGD in *M. truncatula* or it is possible that these two genes evolved much earlier in plant evolution. In order to determine the evolutionary history of RLCK1 and RLCK2 a phylogenetic analysis of RLCK1 and RLCK2 amino acid sequence was carried out identifying orthologues in a variety of different species across the plant kingdom (fig. 3.6.1).

RLCK1 and RLCK2 have representatives in legumes (fig. 3.6.1 in blue) such as soybean and chickpea, and in monocots (fig. 3.6.1 in light brown) such as rice, switchgrass and maize known to symbiose with AMF. Interestingly orthologues are present in ancient species such as *Selaginella mollendorffii* (spikemoss) and *Physcomitrella patens* (moss), and the basal angiosperm *Amborella trichopoda*. It is also important to note that the non-symbiotic species *Arabidopsis thaliana* does not have orthologues of the RLCKs. The presence of both RLCK1 and RLCK2 in such diverse plant species suggests that RLCK1 and RLCK2 evolved before the WGD event in legumes. The fact that there seems to be at least one copy each of RLCK1 and RLCK2 in most species supports the phenotypic evidence that they are both required for properly functional symbiosis.

**Figure 3.6.1.** Phylogenetic tree for RLCK1 and RLCK2 proteins.

RLCK1 and RLCK2 are closely related proteins but have their own distinct subclades across a wide variety of plants including Soybean (*Glycine max*), Rice (*Oryza sativa* and *Oryza indica*), Poplar (*Populus trichocarpa*), and *Amborella trichopoda*. The tree is rooted using the moss (*Physcomitrella patens*) orthologues. There are no *Arabidopsis* orthologues for either RLCK1 or RLCK2. Legumes are coloured blue, monocots are coloured in light brown, and mosses are coloured in green. Figure over leaf.
3.7 Discussion

Evidence of RLCK1 and RLCK2 in ancient and basal land plants, such as mosses and *Amborella*, shows that these proteins evolved from an early gene duplication event. The persistence of these proteins in symbiotic plant species, and the lack of them in non-mycorrhizal plants such as *A. thaliana*, suggests that both RLCK1 and RLCK2 play an important role during plant-microbe symbiosis. Based upon available gene expression data, expression of *RLCK1* and *RLCK2* is increased during both rhizobial and AMF symbiosis and this increase in expression may be dependent upon NF perception and calcium spiking. *RLCK2* has some constitutive expression in the *Medicago* root which suggests *RLCK2* might have a role during root growth and that it has been recruited for symbiotic interactions. The constitutive expression *RLCK2* could also indicate that it is required for the very early stages of symbiosis and thus its expression may be required prior to signalling, an example of this is the Nod Factor receptors NFP and LYK3 (Amor et al., 2003; Smit et al., 2007).

Stable Tnt1 insertion mutants for *rlck1* and *rlck2* were isolated and found to have a reduction in AMF colonisation. Both *rlck1-2* and *rlck2-1* alleles had smaller, misshapen arbuscules compared to WT R108. The smaller arbuscules were not evident on the EMS generated *rlck2-2* allele. The small arbuscules seen are similar to those seen on *Mtpt4* and *Mt-ha1* (Javot et al., 2007; Krajinski et al., 2014; Wang et al., 2014). It is unclear if this phenotype is due to stunted development of arbuscules or premature senescence. A time course looking at arbuscule development will be able to differentiate these two things.

The reduction in nodulation with the *rlck1-2* allele agrees with the effects reported for RNAi knockdown against *RLCK1* carried out by Andrio et al. (2013). *RLCK1* does appear to have a role in rhizobial infection with the *rlck1-2* mutants showing a hyper infected phenotype which is sometimes seen on mutants that have a reduced nodule number (Murray et al., 2011). Promoter-GUS analysis showed *RLCK1* expression in infected root hairs which contributes to the evidence of a role for *RLCK1* during rhizobial infection.

The *rlck2-1* allele had an increased number of mature nodules compared to the WT (R108) control while in contrast the *rlck2-2* allele showed a decreased number of infection events and fewer mature nodules compared to the WT (A17) control. The insertion in the *rlck2-1* allele produces a cDNA that is missing 147bp from the middle of the sequence which encodes the RD domain, and results in a tract of missense amino acid sequence and a premature stop codon removing the majority of the kinase domain.
*rlck2-2* originated from a mutant population generated by EMS mutagenesis (RevGen, UK). The seeds obtained were 5th generation after mutagenesis (M5) and by this time many mutations would be fixed through self-pollination. It seems likely that background mutations in *rlck2-2* are contributing to the severe nodulation phenotype. A backcross of the *rick2-2* allele would reduce the number of background mutations and would help to clarify the *rick2* phenotypes. The promoter-GUS analysis for *RLCK2* revealed expression in developing nodules, however the constitutive expression of *RLCK2* in the root makes it hard to distinguish any symbiotically induced increases of expression in the root hairs and during early stages of rhizobial infection. However, the symbiotic induction of *RLCK2* expression is clear from available microarray data.

The discrepancy between the different *rlck2* alleles makes it difficult to conclude about the role of this gene in the symbioses. Further alleles or complementation of the *rlck1* and *rlck2* mutants are needed to better define their roles in symbiosis, however this study has shown that they both have a positive role in mycorrhizal symbiosis.
Chapter 4: Double Knockdown of RLCK1 and RLCK2

4.1 Introduction

Prokaryotes and eukaryotes contain many classes of RNAs with varying cellular functions, such as messenger RNAs (mRNA) and structural features within ribosomes (rRNAs). Eukaryotes contain a class of single-stranded small RNAs involved in endogenous gene silencing and post-transcriptional regulation of mRNA as well as protection from viral genomes by the destruction of viral RNA (Bazin et al., 2012). These single-stranded small RNAs are referred to as silencing RNAs (sRNAs). Two classes of sRNAs are microRNA (miRNA) and short interfering RNA (siRNA) (Hannon, 2002; Molnar et al., 2005; Ossowski et al., 2008).

miRNAs are derived from longer RNA transcripts that contain imperfect complementary sequences that cause fold backs in the RNA. miRNA create the accumulation of a unique sRNA that tends to bind to one specific target sequence (Ossowski et al., 2008). siRNAs are derived from long strands of RNA that contain perfectly complementary sequences separated by an intronic region. This creates a hairpin RNA (hpRNA) with a long region of double stranded RNA (dsRNA) (Chuang and Meyerowitz, 2000; Wesley et al., 2001). siRNA generate several species of sRNA that can bind to many different sequences of mRNA. For both miRNA and siRNA the double stranded RNA is processed by a member of the DICER family of RNases that cut the dsRNA into short pieces of dsRNA between 20 and 30 bp long with 5’ and 3’ overhangs (Depicker and Van Montagu, 1997; Zamore et al., 2000; Bernstein et al., 2001; Elbashir et al., 2001). The sRNAs act as part of the RNA-induced silencing complexes (RISCs) providing the target specificity for the Argonaute catalytic subunit (Hammond et al., 2000; Bernstein et al., 2001; Baumberger and Baulcombe, 2005). It has been shown that siRNA in plants can also cause transcriptional gene silencing (TGS) by interacting with matching DNA, and recruiting DNA- and histone-modifying proteins to put the locus into a silent chromatin state (Hamilton, 2002; Xie et al., 2004).

Jackson et al. (2003) showed in mammalian cells that siRNAs could affect the transcripts with partially complementary sequences as well as the intended target. This is known as off-target silencing. It is thought to be due to the way that Dicer and Dicer-like proteins process the hairpin RNA transcripts, and the generation of multiple different sRNAs. It was also shown by Palatnik et al. (2003) that miRNAs in Arabidopsis could have up to 5 mismatches with a target sequence and still be able to cause degradation. Off-target degradation is useful if trying to knockdown homologues or gene families, but is something to be mindful of when designing specific silencing constructs. The
effectiveness of silencing constructs can vary greatly, the reason for which is not greatly understood, but could be due to positive feedback systems where transcription rates increase due to the reduced levels of mRNA.

The best way of studying genes that are potentially functionally redundant is to cross stable single mutants to produce a double mutant. However, this is not always possible for example if the genes are closely linked on a chromosome or in cases where the double mutant is inviable. As an alternative, methods such as RNA interference (RNAi) can be used to investigate the double mutant phenotype. Despite many attempts to cross rck1-2 and rck2-1, in both directions, a viable seed was not produced. In this chapter I describe use of two different RNAi approaches to observe the nodulation phenotype of the rck1/rck2 double mutant.

4.2 Strategy 1: RNAi double knockdown

In the first approach an RNAi construct was designed to target the RLCK1 mRNA using 99 bp at the 3’ end of the RLCK1 CDS (fig.4.2.1) (see appendix I for construct map) was provided by Myriam Charpentier, John Innes Centre. It was made by TOPO® and Gateway® cloning of the fragment into a modified destination vector of pK7GW|WG2D(II) (Karimi et al., 2002) in which the eGFP has been replaced with dsRED as a plant marker - the new backbone is called pK7GW|WG2D(II)R (Capoen et al., 2011). This construct is named pK7GW|WG2D(II)R RLCK1 RNAi.

WT (R108) roots were transformed with the RNAi construct pK7GW|WG2D(II)R RLCK1 RNAi via Agrobacterium rhizogenes-mediated hairy root transformation (Boisson-Dernier et al., 2001). The transformed roots were inoculated with Sm1021 and examined at 21 dpi. The knockdown roots showed a significant reduction in the number of nodules at 21 dpi (p=0.019) compared to the empty vector controls (fig. 4.2.2a). qRT-PCR was used to determine the efficiency of the knockdown of the RLCK1 and RLCK2 transcripts. The construct knocks down RLCK1 by ~90% (p=0.030) compared to the empty vector control (fig. 4.2.2b). The construct also causes a reduction of RLCK2 expression compared to the empty vector control (p=0.13) (fig. 4.2.2b), this could be due to off-target gene silencing as the RLCK2 CDS has a 76% sequence homology to the RLCK1 CDS at the nucleotide level.
Figure 4.2.1. Clustal Omega alignment of RLCK1 and RLCK2 CDS with 5’ and 3’ UTR. Continued over page.
Figure 4.2.1. Clustal Omega alignment of RLCK1 and RLCK2 CDS with 5’ and 3’ UTR. RLCK1 and RLCK2 transcripts aligned using Clustal Omega. Estimated 5’ UTR 161 bp upstream of start codon and 3’ UTR 100 bp downstream of stop codon. pK7GW|WG2D(II)R RLCK1 RNAi target region highlighted in blue. pK7GW|WG2D(II)R RLCK2 RNAi target region highlighted in yellow. Asterisks signify matching alignment.
Figure 4.2.2. pK7GW|WG2D(II)R RLCK1 RNAi nodule number and fold change of expression of RLCK1 and RLCK2 relative to empty vector controls.

(a) Knockdown plants show on average a significant reduction in the number of nodules compared to the empty vector control. Bars represent standard error. (b) RLCK1 and RLCK2 expression in the empty vector and RLCK1 knockdown roots was analysed by qRT-PCR. Knockdown plants have a reduced expression of RLCK1 and RLCK2 by approximately 90%. Bars represent relative standard error. Averages are composed of 10 to 11 transformed roots. Data shown is representative of 2 experiments. A 2-tailed t-test was used to compare the means to the empty vector control * = p < 0.05
**4.3 Strategy 2: RLCK2 knockdown in the RLCK1-2 mutant**

In order to confirm if the phenotype seen with pK7GW|WG2D(II)R RLCK1 RNAi was due to the rlck1/rlck2 knockdown and not due to off-target silencing of an alternative gene, a new RNAi construct was made using a 161 bp region of the estimated 5’ UTR of RLCK2 immediately upstream of the start codon (fig. 4.2.1). This construct was designed to be more specific to RLCK2 and named pK7GW|WG2D(II)R RLCK2 RNAi (see appendix I for plasmid map). As a stable Tnt1 mutant for RLCK1 was available, construct pK7GW|WG2D(II)R RLCK2 RNAi alongside an empty vector control, was transformed into WT (R108) and rlck1-2 roots via A. rhizogenes-mediated transfer (Boisson-Dernier et al., 2001). Plants were inoculated with Sm1021 and roots observed at 14 dpi.

*rlck1-2* empty vector (EV *rlck1-2*) roots had significantly (p=0.04) fewer nodules than the R108 empty vector control (EV R108) (fig. 4.3.1a). This is consistent with the nodule phenotype seen previously for *rlck1-2* (fig. 3.3.2.1). RLCK2 knockdown roots (RNAi R108) resulted in a significant reduction in the number of nodules compared to EV R108 (p=0.007). Knockdown of RLCK2 in *rlck1-2* roots (RNAi rlck1-2) have a significant reduction in nodule number compared to EV R108 (p=3.57x10^{-6}), but also a further reduction in nodule number compared to EV *rlck1-2* (p=0.002) and RNAi R108 (p=0.05) (fig. 4.3.1a). The RLCK2 specific RNAi construct significantly reduces the expression of RLCK2 in both the R108 and *rlck1-2* backgrounds by an average of approximately 50% (p=0.03 for both) (fig. 4.3.1b).
Figure 4.3.1. pK7GW|WG2D(II)R RLCK2 RNAi nodule number and relative expression of RLCK2 in RLCK2-specific knockdown roots.

(a) WT (R108) or rlck1-2 roots transformed with empty vector (EV) or RLCK2 RNAi knockdown (RNAi) constructs. Roots transformed with the RLCK2 knockdown construct had a significantly reduced nodule number compared to their EV control. Nodulation scored at 14 dpi. Bars represent standard error. (b) RLCK2 expression was analysed by qRT-PCR. RLCK2 expression was significantly reduced in in the knockdown roots. Bars represent relative standard error. Averages are from of 7 to 14 transformed roots. Data shown is representative of 2 experiments. A 2-tailed t-test was used to compare the means to the R108 empty vector control *p < 0.05, **p<0.01 ***p<0.001
4.4 Discussion

A stable *rck1/rck2* double knockout mutant would be the best material to determine quantitative symbiotic phenotypes. In the absence of a stable double mutant, the effect of an *rck1/rck2* double mutant on rhizobial symbiosis was investigated using RNAi.

*pK7GW|WG2D(II)R RLCK1 RNAi* creates a hairpin RNA that is identical in sequence to a region of to *RLCK1* mRNA and highly similar to *RLCK2* mRNA. *pK7GW|WG2D(II)R RLCK1 RNAi* was designed to work via PTGS to directly reduce the level of *RLCK1* transcript and to reduce the levels of *RLCK2* transcript by off-target gene silencing. The second construct *pK7GW|WG2D(II)R RLCK2 RNAi* was designed to be more specific to *RLCK2*, creating a hairpin RNA identical to the 5' UTR of *RLCK2* immediately upstream of the start codon.

Both RNAi constructs caused a reduction of *RLCK2* expression, however *pK7GW|WG2D(II)R RLCK2 RNAi* is less effective in reducing *RLCK2* transcript levels than *pK7GW|WG2D(II)R RLCK1 RNAi*; This could be due to the 5'UTR being AT rich and thus less efficient at binding the target transcript. When levels of *RLCK1* and *RLCK2* expression were simultaneously reduced, either by double knockdown or by selective targeting of *RLCK2* in the *rck1* mutant, transformed roots showed a near complete reduction in the number of nodules.

The strong reduction in nodulation in the RNAi-mediated double knockdown roots supports the idea that *RLCK1* and *RLCK2* may have partially overlapping or synergistic functions during symbiotic interactions. *rck1* and *rck2* single mutants had relatively stronger phenotypes during AMF interactions than with rhizobial interactions, therefore it would be interesting to see how the double knockdown roots interact with AMF. I would expect that in the absence of *RLCK1* and *RLCK2* arbuscular mycorrhization would be massively reduced compared to WT and single mutants.
Chapter 5: Work towards identification of interacting partners of RLCK1 and RLCK2 Proteins

5.1 Introduction
In PAMP triggered immunity and brassinosteroid signalling cytoplasmic kinases have been shown to interact with RLKs at the plasma membrane and relay the signal through the cytoplasm to activate gene expression in the nucleus (Tang et al., 2008; Zhang et al., 2010a; Kim et al., 2011; Shi et al., 2013a; Shi et al., 2013b; Wang et al., 2013). As RLCK1 and RLCK2 are predicted to be cytoplasmic kinases and are expressed during both symbioses it is possible that they interact with the other kinases of the CSP such as NFP, LYK3, DMI2 or DMI3 or themselves (Stracke et al., 2002; Lévy et al., 2004; Mitra et al., 2004). Potential interactions could be tested through yeast-2-hybrid assays and co-immunoprecipitation (Co-IP) experiments. In addition, fluorescent fusions of these proteins could be used for co-localization studies and to determine their subcellular localization. Towards accomplishing these goals a large suite of constructs were made using different epitope tags and fusions that could be used for localization, yeast-2-hybrid and Co-IP approach to find interacting partners. In order to make the large set of multigene constructs needed for Co-IP a more advanced cloning approach was taken using the new Golden Gate cloning system (Engler et al., 2008; Engler et al., 2009; Weber et al., 2011).

Traditionally molecular cloning was a long process comprised of multiple steps, cutting and sticking DNA with restriction enzymes and ligases until the desired construct was complete. Many attempts have been made to streamline the process of molecular cloning and make DNA construction a standardised process. In 1996 Rebatchouk et al published the NOMAD strategy where modules could be combined sequentially using the StyI restriction enzyme to form a composite module that can then be subcloned into alternate preconstructed expression vectors (Rebatchouk et al., 1996). Knight (2003) proposed the BioBrick standard of assembly where each biological part (promoter, gene, tag etc.) is sequentially combined using restriction enzymes, such that when two or more pieces are joined there is still the same restriction sites on the ends as before making the basic and composite parts idempotent. The beauty of the BioBrick system was that any of the basic or composite parts could be joined to any other and an ever growing library of parts was available (Shetty et al., 2008). The Gateway® cloning system by Invitrogen™ remains a popular cloning system, using recombination sites rather than restriction enzymes, to place fragments of interest into destination vectors in frame with tags or signals already present in the destination vector (Hartley et al., 2000; Karimi et al., 2002). This reduces the cloning process to 3 steps; amplification, BP reaction and LR reaction. The compatibility between entry and destination vectors means that the same cloned
fragment within an entry vector can be inserted into multiple different destination vectors reducing the time taken to create new constructs. However, there are limitations with all of these cloning systems. For NOMAD and BioBrick the limitations lie in only being able to assemble fragments in single sequential steps. In Gateway® the limits are the limited number of destination vectors available, the maximum size of the vector and the creation of unavoidable linker nucleotides at the recombination site.

The Golden Gate cloning system allows directional multi-fragment assembly in one reaction (Engler et al., 2008; Engler et al., 2009). Golden Gate cloning is based on Type IIS restriction enzymes which cleave outside of their recognition site. This allows the creation of many different cleavage sites and when the recognition sites are placed flanking a DNA fragment of interest the cleavage sites are removed from the ligated product. The design of the cleavage sites can be such that the pieces can be ligated without additional nucleotides or linkage sequences i.e. the restriction enzyme cut and ligation sequence can be the last base pair of the CDS sequence and the first 3 bases of a tag allowing for the seamless ligation of CDS to tag (Engler et al., 2008; Engler et al., 2009; Weber et al., 2011).

The Golden Gate system works by having 3 levels of construction. Level 0 modules (L0) are the basic components, such as promoters, signal peptides, 5'UTRs Terminators, 3' Tags, and CDSs. These modules can be plasmid inserts, PCR amplicons or synthesised double stranded DNA. Each type of component would be flanked by the same cleavage sites and these sites would be compatible with the flanking module. For example, all promoter sequences would have a 5’ overhang of GGAG and 3’ overhang of TACT, all 5’ UTR sequences have 5’ overhangs of TACT and 3’ overhangs of AATG, and so on. In this way any combination of basic modules from the L0 library can be used as long as the overhangs match up. The combined L0 constructs combine using the Type IIS restriction enzyme BsaI and a T4 DNA ligase to create a Level 1 (L1) module which is a complete transcriptional unit. Multiple L1 modules can then be combined using a different Type IIS restriction enzyme BpiI and a T4 DNA ligase to create multigene constructs. Up to 7 L1 modules can be combined into Level 2 (L2) constructs (Engler et al., 2008; Engler et al., 2009; Weber et al., 2011). If an L2 construct is required with more than 7 transcriptional units, an additional reaction can be performed using the L2i destination vector which can then accommodate an extra 6 constructs (13 transcriptional units in total), as demonstrated by Weber et al. (2011).
In this chapter I have used both the Gateway® and Golden Gate systems to create a suite of destination vectors that can be used for subcellular localisation and biochemical analysis of RLCK1 and RLCK2 using a yeast 2-hybrid and Co-IP approach.

5.2 In silico analysis of RLCK1 and RLCK2 proteins

The 373 and 374 amino acid sequences of RLCK1 and RLCK2 were analysed for potential protein domains initially using InterProScan4 (Hunter et al., 2012) and the ExPASy Bioinformatics Portal (Artimo et al., 2012) (accessed in 2011) and then again later using InterProScan5 (Jones et al., 2014) and the Calmodulin target database (http://calcium.uhnres.utoronto.ca/ctdb/ctdb/home.html) (accessed in 2014).

Initial predictions suggested that RLCK1 and RLCK2 were predominantly kinase domain with an ATP binding site and an RD active site. A Concanavalin A-like lectin domain and a calmodulin binding domain at the N-terminal region of the protein before the RD active site were identified using the updated protein prediction software (table 5.2.1). No signal sequences or transmembrane domains are predicted suggesting cytoplasmic localisation. Therefore the full length proteins can be used in the yeast 2-hybrid system as they are suggested to be soluble proteins.

<table>
<thead>
<tr>
<th>Domain</th>
<th>Amino acid position in RLCK1</th>
<th>Amino acid position in RLCK2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concanavalin A-like Lectin domain</td>
<td>12-54</td>
<td>27-59</td>
</tr>
<tr>
<td>Calmodulin binding domain</td>
<td>55-119</td>
<td>60-124</td>
</tr>
<tr>
<td>ATP binding domain</td>
<td>48-71</td>
<td>53-75</td>
</tr>
<tr>
<td>RD active site</td>
<td>164-176</td>
<td>169-181</td>
</tr>
<tr>
<td>Kinase domain</td>
<td>23-313</td>
<td>31-329</td>
</tr>
</tbody>
</table>

Table 5.2.1. Positions of predicted protein domains in RLCK1 and RLCK2.
**5.3 Preliminary Yeast 2-Hybrid Analysis**

A preliminary yeast 2-hybrid experiment was conducted testing RLCK1 and RLCK2 for interactions with symbiosis pathway kinases DMI2 and DMI3, as well as with themselves and each other. Yeast 2-hybrid constructs were made for RLCK1 and RLCK2 by the TOPO® and Gateway® cloning systems into both bait (pGADT7) and prey (pGBK7) vectors. DMI2 and DMI3 constructs were obtained from the lab construct library.

Interactions with AD-T or BD-53 were used as negative controls and the AD-T/BD-53 interaction was used as a positive control. When grown on the selection plate only the positive control yeast grew (fig. 5.3.1). These preliminary results suggest that the RLCKs do not interact with these components of the Nod factor signalling pathway. This approach was not pursued further.

![Yeast 2-Hybrid assay](image)

**Figure 5.3.1.** Yeast 2-Hybrid assay for RLCK1 and RLCK2 protein interactions. RLCK1 and RLCK2 were tested for interactions with RLCK1, RLCK2, DMI2 and DMI3.

(a) AD-DMI2, BD-RLCK1, (b) AD-DMI2, BD-RLCK2, (c) AD-DMI2, BD-53, (d) AD-DMI3, BD-RLCK1, (e) AD-DMI3 BD-RLCK2, (f) AD-DMI3, BD-53, (g) AD-T, BD-RLCK1, (h) AD-T, BD-RLCK2, (i) AD-T, BD-53 (positive control), (j) AD-RLCK1, BD-RLCK1, (k) AD-RLCK1, BD-RLCK2, (l) AD-RLCK1, BD-DMI2, (m) AD-RLCK1, BD-DMI3, (n) AD-RLCK1, BD-53, (o) AD-RLCK2, BD-RLCK1, (p) AD-RLCK2, BD-RLCK2, (q) AD-RLCK2, BD-DMI2, (r) AD-RLCK2, BD-DMI3, (s) AD-RLCK2, BD-53.
5.4 Protein localisation
Protein localisation constructs were made by cloning the CDS of RLCK1 and RLCK2 fused to mCherry behind the Ubiquitin promoter. RLCK1 and RLCK2 were fused at the 5' end to mCherry through PCR and then cloned into vector pUB-GW-GFP (Maekawa et al., 2008) by the Gateway® cloning system. This destination vector contains a constitutively expressed GFP to allow the easy identification of transgenic roots. To check that the fused proteins were expressed the constructs were infiltrated into N. benthamiana leaves. Leaf discs were taken at 1, 2 and 3 dpi and checked on a confocal microscope for protein localisation. No mCherry expression was seen. GFP expression was seen at 2 dpi which suggests that the constructs were transformed into N. benthamiana. The lack of mCherry fluorescence suggests that the constructs are not correctly expressing the fusion protein or that the fusion protein was being degraded. Due to uncertainty over the functionality of the constructs they were not transformed into M. truncatula roots.

5.5 Construction of Golden Gate constructs for future protein interaction studies
Using the Golden Gate cloning system constructs were made that can be used to look for interaction partners by Co-IP in Medicago roots or targeted interactions in N. benthamiana (table 5.5.1; more detail in appendix 1). Interactions of the RLCKs would help us elucidate a role for these proteins during symbiosis. N. benthamiana leaves were infiltrated with A. tumefaciens transformed with the constructs and leaf discs were taken at 1, 2 and 3 dpi. Proteins were detected using Western blots and α-myc or α-eGFP antibodies. Expected protein sizes are listed in table 5.5.2.

C-terminally myc tagged RLCK1 and RLCK2 were detectable at 2 and 3 dpi with most constructs with the exception of constructs EC67044 to EC67047. N-terminally tagged RLCK1 was detectable at 1, 2 and 3 dpi for most constructs but myc-RLCK2 was not very well expressed and was only detectable with the EC67035 construct at 1 and 3 dpi. Out of all the eGFP tagged proteins MtHMGR1-GFP was the easiest to detect via Western Blot and was expressed with all its respective constructs being detectable at 2 and 3 dpi. Otherwise MtDMI2-GFP was weakly detected at 2dpi with construct EC67022 (fig. 5.5.1).
<table>
<thead>
<tr>
<th>Construct</th>
<th>Protein 1</th>
<th>Protein 2</th>
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</thead>
<tbody>
<tr>
<td>EC67007</td>
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<td>-</td>
</tr>
<tr>
<td>EC67008</td>
<td>RLCK2-myc</td>
<td>-</td>
</tr>
<tr>
<td>EC67014</td>
<td>RLCK1-myc</td>
<td>MtNFP-eGFP</td>
</tr>
<tr>
<td>EC67015</td>
<td>RLCK1-myc</td>
<td>MtLYK3-eGFP</td>
</tr>
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<td>EC67016</td>
<td>RLCK1-myc</td>
<td>MtHMGR-eGFP</td>
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<td>EC67017</td>
<td>RLCK1-myc</td>
<td>MtDMI2-eGFP</td>
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<td>MtLYK3-eGFP</td>
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<td>MtDMI2-eGFP</td>
</tr>
<tr>
<td>EC67034</td>
<td>myc-RLCK1</td>
<td>-</td>
</tr>
<tr>
<td>EC67035</td>
<td>myc-RLCK2</td>
<td>-</td>
</tr>
<tr>
<td>EC67038</td>
<td>myc-RLCK1</td>
<td>MtNFP-eGFP</td>
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<td>EC67047</td>
<td>myc-RLCK2</td>
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Table 5.5.1. A list of Golden Gate constructs and the proteins they express.
<table>
<thead>
<tr>
<th>Protein</th>
<th>Size (kD)</th>
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</tr>
<tr>
<td>myc</td>
<td>3.67</td>
</tr>
</tbody>
</table>

**Table 5.5.2.** Expected protein sizes for the proteins expressed by the Golden Gate constructs.
Figure 5.5.1. continued overleaf
Figure 5.5.1. Expression of Proteins in *N. benthamiana* from the Golden Gate constructs.

Expression of tagged proteins in *N. benthamiana* infiltrated with Golden Gate constructs using *A. tumefaciens* was checked at 1, 2 and 3 dpi by Western blot. RLCK1-myc and RLCK2-myc bands could be detected at 2 and 3 dpi (~45kD). myc-RLCK1 can be seen with some constructs at 1, 2 and 3 dpi. myc-RLCK2 could only be detected with one construct (EC67035) at 1 and 3 dpi. HMGR-GFP could be detected at 2 or 3 dpi (~86kD). DMI2-GFP was weakly detected at 2 dpi with construct EC67022 (~131.18kD). See table 5.4.1 for construct details and table 5.4.2 for protein sizes. Protein levels were checked using Coomassie (blue) or Ponceau (Red) staining; Rubisco bands shown beneath the western blot.
5.6 Discussion

An attempt was made to visualise protein localisation using mCherry tagged overexpressed RLCK1 and RLCK2. In preliminary experiments in *N. benthamiana* no mCherry fluorescence was seen in the leaf discs under the confocal microscope at 1, 2 or 3 dpi. A Western blot is needed to be sure that the proteins are being properly expressed. It may be that no mCherry was seen because expression was being suppressed in the *N. benthamiana* leaves. Co-infiltration with a silencing suppressor plasmid such as p19 would improve expression of the tagged proteins in *N. benthamiana* (Voinnet *et al.*, 2003). Ideally localisation of the RLCKs would be best performed in *M. truncatula* roots where these proteins are naturally expressed. Alternate constructs using the ubiquitin and the native promoters in combination with the genomic and CDS sequences could be used to maximise the construct expression in *Planta*. It would also be good to check the localisation of the RLCKs with and without rhizobia and AMF in *M. truncatula* roots as the localisation may be different upon interaction with symbiotic partners.

Based on the expression patterns of RLCK1 and RLCK2 and the symbiotic phenotypes observed, members of the Nod factor signalling pathway were chosen to be investigated as possible interacting partners. Yeast 2-hybrid constructs have been made for RLCK1 and RLCK2. The yeast 2-hybrid constructs described here can be used in the future with a yeast 2-hybrid library to screen for interacting partners. A western blot should be performed to check that the proteins were correctly expressed in yeast. Yeast 2-hybrid uses the soluble regions of proteins, so for transmembrane proteins such as MtNFP1 and MtLYK3 and MtDMI2 this may not be the best method to use for the receptor kinases as they may require the full length protein for activation, or activation by a ligand such as Nod Factor, before they are able to interact. For this reason interaction studies of PAMP RKs and RLKs such as FLS2 tend to be investigated using a Co-IP method in *N. benthamiana* leaves.

Several constructs have been made for expressing RLCK1 and RLCK2 tagged to a myc tag, at either N or C termini, on their own or in combination with GFP tagged components of the symbiosis pathway. Preliminary analysis showed that only a handful of these constructs have given clearly expressed tagged proteins of the correct size on the Western blots. The proteins which were detected by Western blot analysis were the RLCKs and MtHMGR1. The RLCK proteins are smaller and are predicted to be localised within the cytoplasm which would make them easier to extract from the plant material than the larger membrane bound proteins. Further testing is needed to optimize expression of these constructs. Once expression is confirmed, these constructs can be used for a targeted Co-IP to see if RLCK1 and RLCK2 interact with MtNFP, MtLYK3, MtHMGR1 or
MtDMI2 in *N. benthamiana*. They can also be transformed into *M. truncatula* roots by *A. rhizogenes*-mediated hairy root transformation to look for interacting partners *in planta* and to confirm any positive interactions observed in *N. benthamiana*.

Additionally in the analysis of RLCK1 and RLCK2 is to look at the protein functions and to discover interacting partners. RLCK1 and RLCK2 are predicted to contain a Concanavalin A lectin or lectin-like domain. Lectins have been shown to bind sugars and oligosaccharides. It would be worth testing these proteins with a carbohydrate-chip to attempt to determine their potential carbohydrate substrate(s) (Houseman and Mrksich, 2002; Park *et al.*, 2004).
Chapter 6: Identification of a mycorrhizal phenotype in *M. truncatula* Tnt1 insertion line NF5270

6.1 Introduction
Alongside the main project of *RLCK1* and *RLCK2*, *Tnt1* insertion seeds were obtained from the Samuel Roberts Noble Foundation for a third predicted kinase, the gene Medtr7g116650. The protein was predicted to contain a transmembrane domain and a cytoplasmic kinase domain, but have no extracellular domain, and as such the gene was named *receptor-like cytoplasmic kinase 3* *RLCK3*. The MtGEA profile for *RLCK3* showed that it was upregulated during AMF colonisation and during nodule senescence, though interestingly not during nodule development (fig. 6.2.1) (Benedito et al., 2008; He et al., 2009; Hogekamp et al., 2011; Czaja et al., 2012; Gaude et al., 2012; Seabra et al., 2012). It was possible that *RLCK3* could provide insight to, and play a role in, the differentiation between rhizobial and mycorrhizal symbiosis.

6.2 Identification of the *RLCK3* mutant and Characterization of its Nodulation and Mycorrhizal Phenotypes
A gene of interest was selected based on the expression profile of probe set Mtr.20292.1.S1_at on the Medicago Gene Expression Atlas (MtGEA) (Benedito et al., 2008; He et al., 2009; Hogekamp et al., 2011; Czaja et al., 2012; Gaude et al., 2012; Seabra et al., 2012). The gene named as *RLCK3* (Medtr7g116650) is up-regulated in later stages in nodulation and in nodules treated with phosphinothricin (PPT), a glutamine synthetase inhibitor. It is also up-regulated during mycorrhizal infection by *R. irregularis* and in root inner cortical cells that contain arbuscules (fig. 6.2.1). To study this gene a *Tnt1* line NF5270 was identified that contained a *Tnt1* insertion in the second exon of *RLCK3* at 1238 bp. This allele was named *rlck3*-1 (fig. 6.2.2 a). R1 generation seeds were obtained from the Samuel Roberts Noble Foundation and homozygote mutants for *rlck3*-1 were identified by PCR using the gene specific primers #11 and #12 and primers #10 and #11 for *Tnt1* identification within *RLCK3* (fig. 6.2.2b and 7.2.2c). See Appendix 1 for primer details.
Figure 6.2.1. Medicago Gene Expression Atlas data for RLCK3.

The graph shows the expression level of RLCK3 under a number of different experimental conditions. Expression of RLCK3 is highly induced at later stages of nodulation and mycorrhization. RLCK3 is specifically upregulated in arbusculated inner cortical cells during AMF colonisation and in nodules treated with a source nitrogen or a Glutamine synthetase (GS) inhibitor (Phosphinothricin (PPT)).
6.2.1 Preliminary Nodulation and Mycorrhizal phenotypes of rlck3-1

Two homozygous mutant sibling lines 9_NF5270 and 15_NF5270 were chosen for further study (fig 6.2.2). Preliminary nodulation and mycorrhizal phenotypes were examined in the progeny of these lines (fig. 6.2.3). Plant were inoculated with Sm1021 and examined at 21 dpi. Unexpectedly, line 15_NF5270 has significantly fewer nodules than WT (R108) (p=0.002) at 21 dpi, however only 5 plants were scored and so requires confirmation. Line 9_NF5270 was not significantly different from WT (R108) (p=0.3) (fig. 6.2.3a).

Plants were inoculated with the PlantWorks mycorrhizal inoculum and scored for percentage arbusculation at 8 wpi. A few progeny for 9_NF5270 was checked whilst seeds for 9_NF5270 and 15_NF5270 were being bulked for further phenotypic analysis. rlck3-1 line 9_NF5270 has significantly fewer arbuscules at 8wpi than WT (R108) (p=3.43x10⁻⁵) (fig. 6.2.3b).

Figure 6.2.2. Identification of RLCK3 Tnt1 insertion mutants in line NF5270. (a) Gene structure for RLCK3. Arrows represent exons. The position of the Tnt1 insertion is show as rlck3-1. Gel electrophoresis image showing (b) WT RLCK3 and (c) Tnt1 insertions amplified by PCR using DNA from individual NF5270 R1 plants. WT (R108) DNA was used as a control. Plants number 9 and 15 were selected as homozygote mutants and plant number 14 was chosen as a heterozygote for further study.
Figure 6.2.3. Preliminary nodulation and AMF colonisation phenotypes of *rck3-1*
(a) The NF5270 Tnt1 sibling line 15 has fewer nodules than line 9 and WT (R108). Data is an average of 5 - 10 plants. Plants were scored at 21 dpi. A 2-tailed t-test was used to compare the means of each line and the WT (R108) (*p*<0.01). (b) Percentage of the root with arbuscules at 8 wpi with the PlantWorks inoculum. The 9_NF5270 line has significantly fewer arbuscules than WT (R108). Data is an average of 11 (R108) and 17 (NF5270) plants. A 2-tailed t-test was used to compare the means (*p*<0.001). Bars are standard error.
6.2.2 Segregation population of *rlck3*-1

To test whether the mutation in *RLCK3* cosegregated with the mycorrhizal phenotype observed in 9_NF5270, progeny from an *rlck3*-1 heterozygote sibling line 14 (14_NF5270) was used to generate a segregating population. The *rlck3/rlck3* plants were not significantly different from the *RLCK3/RLCK3* plants for levels of arbusculation in the root (Fig. 6.2.4).

These results strongly suggest that the symbiotic phenotypes observed in NF5270 were not due to the insertion in *RLCK3*, and instead resulted from alternative mutations arising in the original R0 NF5270 *Tnt1* insertion mutant line. It also indicates that the gene responsible for the mycorrhizal phenotype present in 9_NF5270, is not in the sibling line 14_NF5270 used to generate the segregating population for *rlck3*-1, and that the gene responsible for the mycorrhizal phenotype was segregating in the parental generation (R2). The unknown gene conferring the strong mycorrhizal phenotype was named *SCOOBY*. 
Figure 6.2.4. AMF colonisation of progeny for different genotypic classes from the 14_NF5270 segregating population for rick3-1.

(a) Average percentage of arbusculation in the rick3-1 segregating population generated from plant 14_NF5270. Roots were scored at 3 wpi with the Symplanta inoculum (see materials and methods). The data are averages of 11, 20 and 12 plants respectively. The means were not significantly different from each other (2-tailed t-test, p>0.05). The bars represent standard error. (b) A box plot to show the distribution of arbusculation in each rick3-1 genotype.
6.3 *scooby* mycorrhizal phenotype

A further investigation into the *scooby* mycorrhizal phenotype was carried out. Lines 9_NF5270 and 15_NF5270 were grown with AMF chive inoculum for 5 weeks, until the WT colonisation was ≥ 50%. Both the 9_NF5270 and 15_NF5270 line showed a reduced percentage of root that contained arbuscules, with a mean average of 7.2% and 5.2% respectively compared to WT (R108) mean average of 81.4% and 77.33% respectively. A 2-tailed t-test shows the colonization in both lines was significant (P=1.08x10^{-13} and P=4.92x10^{-16} for 15_NF5270 and 9_NF5270 respectively) (fig. 6.3.1). Microscopic examination of the *scooby* roots shows that as well as having a reduced number of arbuscules in the inner cortical cells, the arbuscules that were formed did not fill the cells and were misshapen (fig. 6.3.2). The AMF was able to penetrate the root, grow intracellular hyphae and form vesicles in the *scooby* mutant as in WT (R108).

![Figure 6.3.1. AMF colonisation of NF5270 Tnt1 lines](image)

(a) Percentage of the root with arbuscules with the chive inoculum when colonisation of WT (R108) is >50%. Both 9_NF5270 and 15_NF5270 have significantly fewer arbuscules than WT (R108). Data is an average of 5 15_NF5270, 5 9_NF5270 and 9 and 10 WT (R108) plants respectively. A 2-tailed t-test was used to compare the means of the mutant to the WT (R108) control (p<0.001).
Figure 6.3.2. Mycorrhizal phenotypes of *scooby* and WT (R108).

Light microscope images of (a, c) *scooby* mutant and (b, d) WT (R108) roots colonised by *R. irregularis*. AMF hyphae are able penetrate and grow through the root cortex of NF5270 (a) and form vesicles (black arrowhead) as the WT (R108) (white arrowhead). Although the mutant can form structures resembling arbuscules (asterisks) they are much fewer in number and are misshapen compared to WT (R108) (b and d, starred). Images were taken at 5wpi. Scale bars are 200µm (a), 100µm (b) and 50µm (c and d).
6.4 Discussion
The regeneration of *M. truncatula* containing the *Tnt1* retrotransposon can cause many insertion mutations (Tadege *et al.*, 2008). When taking a reverse genetics approach, the phenotype seen in one of the *Tnt1* mutant lines may not necessarily be caused by the gene of interest but by a different background mutation. The Samuel Roberts Noble Foundation lists the mutant population by generating FSTs which show the genomic regions around the *Tnt1* insertion. However, due to the possible number of insertions and limitations to the approach not all FSTs in a regeneration line are identified.

In the case of the mutant line NF5270, it was selected based on a known FST for the gene *RLCK3* (Medtr7g116650), a predicted cytoplasmic kinase. Initial phenotyping in a sibling line homozygous for this gene showed a strong mycorrhizal phenotype. However, a segregating population for *rlck3-1*, which was generated from a heterozygote sibling line (14_NF5270), showed that the mycorrhizal phenotype was not caused by the *rlck3* mutation. It also indicates that the mutation responsible for the mycorrhizal phenotype seen in 9_NF5270 and 15_NF5270 (*scooby*), was hemizygous at R0 and was segregating at R2 (fig. 6.4.1).
Figure 6.4.1. *Tnt1* insertion line NF5270

A diagram of the *Tnt1* insertion line NF520. Generation R0 are hemizygous for many *Tnt1* insertions. Seeds from generation R1 are obtained and generation R2 plants which were genotyped (fig 6.2.2) – green are WT, blue are homozygous mutants and purple are heterozygous for the *rlck3-1 Tnt1* insertion. Another insertion mutation in the line responsible for reduced mycorrhization, symbolised by the yellow circles, is also segregating in the R2 population and is homozygous in plant 9 and 15 but WT in plant 14.
As the mycorrhizal phenotype, *scooby*, is strong in the two sibling lines 9_NF5270 and 15_NF5270, and is sustained over generations, it is worthy of further investigation. The gene, or genes, responsible for the phenotype can be investigated through genetic mapping and additional TAIL-PCR to look for more FSTs. Single base pair mutations, deletions and frame-shift mutations can also occur during the regeneration process. Whole genome sequencing such as Illumina sequencing, or transcriptome sequencing via RNA-seq could be used to identify these mutations.

Further characterisation of the *scooby* mycorrhizal phenotype should be carried out. A time-course following AMF colonisation of mutant roots overtime could help to establish at which stage the AMF is inhibited. Lack of arbusculation could be caused by inhibition of the AMF at the epidermis or at a cortical level. Growth with nurse plants would establish if 9_NF5270 and 15_NF5270 are missing signalling molecules causing the *scooby* phenotype. Differential gene regulation and genetic markers could shed light on the function or position of *SCOOBY* relative to the symbiosis pathway.

*SCOOBY* may prove to be a mycorrhizal specific gene, of which there are only a few currently known for *M. truncatula* including a GRAS-type transcription factor (*RAM1*), a glycerol-3-phosphate acyltransferase (*RAM2*) an H⁺-ATPase (*Mt-HA1*), and a phosphate transporter (*MtPT4*) (Javot *et al.*, 2011; Gobbato *et al.*, 2012; Wang *et al.*, 2012; Krajinski *et al.*, 2014; Wang *et al.*, 2014). In *ram1* AMF is inhibited at the epidermal level (Gobbato *et al.*, 2012). In *ram2*, the AMF appears to be inhibited at the epidermal level through a lack of hyphopodia formation; however with increased AMF levels some arbuscules can be seen (Wang *et al.*, 2012). *mt-ha1* and *mtpt4* have similar phenotypes, allowing epidermal entry of AMF but have premature senescence of the arbuscules resulting in fewer arbuscules visible in the root cortex (Javot *et al.*, 2011; Wang *et al.*, 2014). To confirm that *SCOOBY* is not caused by *RAM2*, *Mt-HA1* or *MtPT4* these genes could be checked for *Tnt1* insertions by PCR. Alternatively complementation of the *scooby* phenotype could be checked by transformation of 9_NF5270 and 15_NF5270 root with WT *RAM2*, *Mt-HA1* or *MtPT4*. 
Chapter 7: Discussion

Mutant studies have provided insight into many of the genetic components required for the establishment and maintenance of the plant-microbe symbioses. However, many of the genetic aspects of the plant-rhizobial and plant-mycorrhizal symbioses remain to be identified. It is as yet unknown how the signalling is transduced from the RLKs at the plasma membrane to the nucleus through the CSP, or how parallel signalling pathways help to modulate each symbiosis. It also remains unclear how the plant is able to establish the difference between friend and foe when both carry microbe-associated molecular patterns (MAMPs) that activate plant defence. The control of ROS levels during development and defence are tightly controlled, and whilst there is some evidence to suggest roles for ROS during symbiosis, it is not yet known how ROS levels are regulated during symbiosis. We are at an exciting time in the study of mycorrhizal symbiosis with the genome of *R. irregularis* being recently published (Tisserant et al., 2013). This will provide the community with better tools to study this symbiosis in the future.

*scooby* is a novel mycorrhizal mutant in *M. truncatula*

The *scooby* mutant appears to be a novel mycorrhizal mutant in *M. truncatula*. *scooby* plants had fewer arbuscules (fig. 6.3.1) that appeared to be arrested between the birdseye stage and full arbuscule development. Arbuscules are smaller in *scooby* than WT (R108) and do not fully fill the cortical cell (fig. 6.3.2). However, the branches of the arbuscules are well defined and appear different than the early senescent arbuscules seen in *Mpt4* or *Mt-ha1* (Javot et al., 2007; Krajinski et al., 2014; Wang et al., 2014). AMF are able to penetrate the *scooby* roots as normal, which rules out *ram1* and *ram2* as these mutants are blocked at the epidermis, and *scooby* has a large amount of intraradical hyphae seen which is different to the phenotypes of *Mt-HA1* (Gobbato et al., 2012; Wang et al., 2012; Krajinski et al., 2014; Wang et al., 2014). Mutation or down regulation in the half –ABC transporters *STR* and *STR2* have been identified in *M. truncatula*, rice and *L. japonicus* (Zhang et al., 2010b; Gutjahr et al., 2012; Kojima et al., 2014), however they have a lower total level of AMF colonisation compared to *scooby* despite similar stunted arbuscules in these mutants. Groth et al. (2013) recently described two mycorrhizal mutants in *L. japonicus*, SL0181-N mutant had two arbuscule phenotypes (type I and II). Type II arbuscules were severely stunted and appeared similar to the *scooby* phenotype. However segregation analysis revealed that the severe arbuscule stunting is likely to result from the combination of two distinct loci.
Further analysis of the mycorrhizal phenotype is needed to fully characterise the scooby mutant. A time-course following AMF colonisation of mutant roots over time could help to establish at which stage the AMF is inhibited. The phenotypes of some mycorrhizal mutants that are inhibited at the early stages of mycorrhization, such as pre-mycorrhizal infection 1 (PMI1) in tomato, are able to be rescued when grown alongside WT “nurse” plants in the presence of AMF (David-Schwartz et al., 2001). In cases such as that of Mtpt4 and Mt-ha1, nurse plants are unable to rescue the shrunken arbuscule phenotype, however the number of these type of arbuscules seen per root is increased (Javot et al., 2007; Wang et al., 2014); this is possibly due to the AMF being provided a source of carbon from the nurse plants and as such AMF growth is not inhibited as it would be with the Mtpt4 and Mt-ha1 mutants alone. Growth with nurse plants would establish if 9_NF5270 and 15_NF5270 are missing signalling molecules, such has been hypothesised for pmi1 or are lacking in nutrient exchange as in Mtpt4 or Mt-ha. Single base pair mutations, deletions and frame-shift mutations can also occur during the regeneration process and could be identified by whole genome sequencing, such as Illumina sequencing, or transcriptome sequencing via RNA-seq.

Two Receptor-like kinases required for nodulation and mycorrhization

RLCK2 is a novel symbiotic gene

RLCK2 encodes a putative receptor-like cytoplasmic kinase expressed constitutively in root tissue with up-regulation during symbiosis. rlck2 mutants were significantly reduced in AMF colonisation (fig. 3.4.1) with smaller arbuscules that are similar in morphology to those seen in M. truncatula mutants Mtpt4 and Mt-ha1 (fig. 3.4.3) (Javot et al., 2007; Wang et al., 2014). RLCK2 may have a nodulation phenotype, however due to disagreeing allelic phenotypes it is not clear if lack of RLCK2 results in an increased (rlck2-1), or decreased (rlck2-2), nodule number compared to WT (fig. 3.3.2.1). On the other hand, due to the relative positions of the premature stop codons, it is more likely that rlck2-1 is the stronger allele and thus it could be argued has the more reliable phenotype. The EMS mutagenesis method used to generate the rlck2-2 allele is likely to have created mutations at many sites in the genome; one of these alternate mutations could be responsible for the nodulation phenotype. Complementation, backcrossing the two alleles or a third allele will be needed to confirm which the correct rlck2 nodulation phenotype is.
**RLCK1/SPK1** is required for correct mycorrhizal development as well as nodulation in *M. truncatula*

**RLCK1/SPK1** has shown by Damiani *et al.* (2012) and Andrio *et al.* (2013) to be expressed during nodulation and upon ROS application. Andrio *et al.* (2013) showed via RNAi that a reduction of **RLCK1/SPK1** expression leads to a reduction in nodule number. Using a stable *Tnt1* insertion mutant, this study has confirmed that the absence of **RLCK1/SPK1** does lead to a reduction in the number of nodules. Additionally, the *rlck1/spk1* mutant had an increased number of infection events (fig. 3.3.1.1), with infection threads being misshaped compared to WT (R108) (fig. 3.3.1.2). This study also found that in mature nodules of the *rlck1/spk1* mutant, the infection threads that would normally be advancing only into the infection zone (Zone II) were advancing into the meristematic zone (Zone I) (fig. 3.3.2.2). There is a delicate balance between carbon provision and N acquisition which the plant controls by regulating the number of successful infections leading to nodule organogenesis. The fewer nodules on the *rlck1/spk1* plants may be a consequence of a feedback loop driven by the hyper-infection of root hairs and within Zone II of the nodule.

In addition to the nodulation phenotypes demonstrated, *rlck1/spk1* mutants had a significantly reduced level of colonisation by the AMF *R. irregularis* compared to WT (R108) (fig. 3.4.1). The arbuscules on *rlck1/spk1* were smaller and were similar in morphology to *Mtpt4* and *Mt-ha1* (Javot *et al.*, 2007; Wang *et al.*, 2014), suggesting that they may be senescing early. A time course checking the progression of the AMF through the root and the speed of the advancement of the mycorrhizal symbiosis is needed to determine if the arbuscules are stunted in growth or are prematurely senescing. This could be achieved by checking plant roots at earlier time points for the number of the arbuscules at each stage of development. If the mycorrhiza do not form mature arbuscules then this would suggest they are stunted before reaching maturity, however if mature arbuscules are seen at these early time points then this would suggest the arbuscules are collapsed and prematurely senescing.

**RLCK1 and RLCK2** are partially redundant during symbiosis

**RLCK1/SPK1** and **RLCK2** have overlapping expression patterns in the root which was visualised using promoter-GUS analysis. **RLCK2** was constitutively expressed in the root and was further upregulated upon symbiotic interactions (figs. 3.5.1.3, 3.5.1.4, and 3.5.2.3). Promoter-GUS analysis and MtGEA data shows that **RLCK1/SPK1** expression is more symbiosis specific than **RLCK2** with expression being highly upregulated upon
interaction with a symbiont or NF application (figs. 3.5.1.1, 3.5.1.2, and 3.5.2.1). 

*rlck1/spk1* and *rlck2* single mutants had very similar mycorrhizal phenotypes (fig. 3.4.1 to fig. 3.4.3). Knocking down *RLCK1/SPK1* and *RLCK2* by RNAi in *M. truncatula* roots results in a large reduction in the nodule number compared to the empty vector control (figs. 4.2.2 and 4.3.1).

Despite their high level of homology, orthologues of RLCK1 and RLCK2 are present in species across the plant kingdom, including ancient plant lineages such as moss and *Amborella*. Given the similarities of these two genes at both nucleotide and amino acid level, it is possible that *RLCK1* and *RLCK2* have partially redundant or synergistic roles during symbiotic development. In the single mutants, one RLCK may be able to partially compensate for the lack of the other. Considering the ubiquitous expression levels of *RLCK2* in the root, *RLCK2* could be required for a more general developmental process and is recruited during symbiosis whilst the expression and role of *RLCK1/SPK1* could be symbiosis specific. However, such a housekeeping role is contradicted by the loss of these genes in the *Brassicaceae*. It is also possible that *RLCK2* is required at the very early stages of symbiont perception, when a response is needed faster than there is time for new protein synthesis. An example of two genes being required partially redundantly whilst having overlapping yet different expression patterns is *ERN1* and *ERN2*, two gene encoding nodulation expressed transcription factors. *ERN1* and *ERN2* can independently activate symbiosis marker gene *ENOD11* (Andriankaja et al., 2007; Cerri et al., 2012) but the *ern1* single mutant is still able to form misshaped infection threads, and nodules which can be complemented by *ERN2* expressed under the *ERN1* promoter (Middleton et al., 2007). Symbiotic gene *DMI3* has been shown to have different roles in cortical cells and epidermal cells when it’s expression is driven via tissue specific promoters in a *dmi3* mutant (Rival et al., 2012). It is possible that *RLCK1* and *RLCK2* have tissue specific roles during symbiosis.

Numerous attempts to cross pollinate *rlck1-2* and *rlck2-1* mutants have not produced any seed. It is possible that *RLCK1* and *RLCK2* also play redundant roles in seed development. Related to this, a rice gene belonging to the same subfamily as *RLCK1/2* (RLCK-XV) Os06g07070 has been shown to be upregulated in embryo and endosperm development (Gao and Xue, 2011). The *ram2* mycorrhizal mutant produces seeds with a dark seed coat that can be permeated by dyes (Wang et al., 2012), demonstrating a link between the two developmental processes. Examples of mutants that are involved in both developmental and symbiotic processes are *sickle* (*ski*) an ethylene insensitive mutant named after the sickle shape of the hypocotyl, and *super numeric nodules* (*sunn*) a long-distance auxin transport mutant of *M. truncatula* that are
both hyper-nodulating and have root architecture phenotypes (Penmetsa and Cook, 1997; Prayitno et al., 2006; van Noorden et al., 2006). A review from Evangelisti et al. (2014) highlights the extensive cross-talk that exists between developmental processes and plant-microbe interactions.

Possible roles of RLCK1/RLCK2 in symbiosis

RLCK1 and RLCK2 are part of the RLCK-XV subfamily of the receptor-like cytoplasmic kinases. This subfamily does not have representatives in A. thaliana but is present in the model legumes M. truncatula, L. japonicus and G. max, as well as many species known to interact with AMF (e.g. rice and tomato). This suggests that this subfamily of RLCKs is required for symbiotic interactions. The RLCK-XV sub-family has not been widely studied. The subfamilies RLCK-V and RLCK-VI that display roles in PAMP and BR signalling, have led to these two subfamilies being relatively well studied. I am now going discuss three possible roles for RLCKs in symbiotic interactions that are not necessarily exclusive: as part of NF signal transduction, as regulators helping to distinguish pathogen from symbiont, and as regulators of ROS mediated development.

Part of the NF signalling cascade

The PTI signalling pathway through FLS2 and the BR signalling pathway through BRI1 have been well studied. Both of these pathways recognise signalling molecules (flg22 and BR) at the cell surface through binding of the signal with RLK complexes. Both FLS2 and BRI1 associate with BAK1 and the RLCKs BIK1, BSKs, and CDG1 (Veronese et al., 2006; Lu et al., 2010; Zhang et al., 2010a; Kim et al., 2011; Sreeramulu et al., 2013). Recognition of the signalling molecules causes phosphorylation of the RLCKs by the RLKs. The RLCKs then phosphorylate targets such as phosphatase BSU1 or the NADPH oxidase RbohD (Mora-Garcia et al., 2004; Li et al., 2014). There is also some evidence that RLCKs BIK1 and BSKs are involved in the cross-talk between PTI and plant development (Lin et al., 2013; Shi et al., 2013a; Shi et al., 2013b).

Currently it is not known in the CSP how the cell surface localised RLKs, NFP and LYK3, pass on the signal to the other RLK DMI2. It is also not known how the signal travels from the plasma membrane to the nucleus. There is some evidence showing the interaction of a MAP kinase with DMI2 (Chen et al., 2012). MAP kinases, a class of RLCKs, are often involved in signal transduction during defence ultimately leading to gene expression (Nühse et al., 2000; Asai et al., 2002; Suarez-Rodriguez et al., 2007; Zhang et al., 2007). Using the PTI and BR signalling pathways as a guide, possible roles for the
RLCKs studied here could be the transactivation of the RLKs NFP and LYK3, and/or DMI2. Data from Antolín-Llovera et al. (2014) shows that L. japonicus SYMRK (MtDMI2) is able to interact with LjNFR5 (MtNFP) in N. benthamiana when the extracellular malectin-like domain (MLD) is cleaved (SYMRK-ΔMLD). SYMRK-ΔMLD is similar in structure to BAK1, which interacts with BRI1 and FLS2 to form receptor complexes in the BR and flagellin signalling pathways. However, the interaction between SYMRK-ΔMLD and NFR5 is able to occur without a ligand. The SYMRK-ΔMLD protein undergoes rapid endocytosis and degradation but low levels can still be detected in planta (Antolín-Llovera et al., 2014). It is possible that the SYMRK-ΔMLD/NFR5 complex in planta binds to an RLCK similar to BIK1 or BSK1 which continues the signal transduction.

The proteins required for NF perception and fast responses to NF have a constitutive level of expression in the roots. Other genes, such as Nodule inception (NIN) are required for the early stages of symbiosis and are up regulated quickly upon NF perception (Schauser et al., 1999). RLCK2 has constitutive expression in M. truncatula roots and both RLCK1/SPK1 and RLCK2 are quickly up-regulated upon NF application. The tagged protein constructs made during this study (Chapter 5) can be used to test RLCK1/SPK1 and RLCK2 for interaction with the known CSP components to identify if RLCK1/SPK1 and/or RLCK2 could be the RLCKs involved in the signal transduction during symbiosis. The non-symbiotic expression of RLCK2 is much higher than RLCK1/SPK1. This would suggest that if these proteins are involved in any initial signal transduction through the CSP that this would initially depend more on RLCK2, while RLCK1/SPK1 may serve mainly to amplify the signal as the infection progresses. On the other hand, our results suggest that in the absence of RLCK2, RLCK1 may still compensate and vice versa, suggesting that sufficient levels of RLCK1 are present to support signalling through the CSP. The available transcriptomics data from mutants suggests that inductions of both genes are CCaMK, and therefore CSP, dependent, so successful signalling through the CSP in either single mutant would lead to induction of the remaining RLCK and fulfil most of the signalling role. This model could be tested by monitoring the onset of calcium oscillations (typically 10 minutes from time of NF application) in roots of the rlck1, rlck2 and double knockdown roots and checking for delays in expression of early nodulation genes such as ENOD11 and NIN.

Alternately the RLCKs may act in pathways that occur in parallel to the CSP that also require NF perception and enable the plant to accommodate the symbionts (Murray, 2011). For example, root hair growth/curling that follows rhizobial attachment in response to NF can still occur in dmi2 and other downstream CSP mutants. Interestingly, root hair growth is associated with ROS production (Foreman et al., 2003).
Differentiating between Pathogen and Symbiosis

Rhizobia and AMF, although symbiotic partners of plants, carry MAMPs that activate PTI such as flagellin and chitin. It has long been under discussion as to how plants are able to distinguish between friend and foe (Ausubel and Bisseling, 1999; Parniske, 2000; Kogel et al., 2006; Paszkowski, 2006; Rey and Schornack, 2013). In both PTI and symbiosis there is an oxidative burst upon PAMP recognition (Alvarez et al., 1998; Ramu et al., 2002). In PTI the levels of ROS species, such as •O₂⁻ and H₂O₂, stay high whereas in symbiosis there is a gradual decline of ROS levels (Shaw and Long, 2003). Studies have shown that PTI activated roots then treated with NF also show a decrease in ROS levels (Shaw and Long, 2003). ROS in PTI leads to the strengthening of cell walls through cross-linking and plays a central role in programmed cell death (Levine et al., 1994). Contrastingly, ROS production induced by NF has been shown to be relatively short lived (Ramu et al., 2002). ROS can also be found to be increased 12 hours after NF or rhizobial treatment of Medicago roots (Shaw and Long, 2003), which is likely to be due to the induction of developmental processes. Interestingly, Medicago roots inoculated with a mutant rhizobia strain that is unable to produce NFs, the ROS response of the plant is the same as that seen in PTI (Bueno et al., 2001).

The defence pathways that are triggered by the MAMPs carried by rhizobia and AMF must be suppressed before the plant is able to accommodate a symbiotic partner. It takes 20 to 30 minutes after NF application until ROS levels in the root begin to decrease, suggesting that new proteins need to be synthesised first (Shaw and Long, 2003). RLCKs like BIK1 and the BSKs have been shown to act as cross-talk mediators between PTI and BR signalling (Lin et al., 2013; Shi et al., 2013a; Shi et al., 2013b). It is possible that RLCKs play a similar function between PTI and symbiosis signalling. Andrio et al. (2013) showed that RLCK1/SPK1 is upregulated upon H₂O₂ treatment and is expressed in nodule zones that also contain high levels of H₂O₂. It is possible that RLCK1/SPK1 and RLCK2 play a role in the differentiation between pathogen and symbiont, possibly regulating ROS levels to enable infection by rhizobia and AMF. To test this hypothesis the response of RLCK2 to ROS application should be investigated; it would be interesting to see if RLCK2 responds similarly to RLCK1/SPK1. However, RLCK2 was not identified by Andrio et al. (2013) in their microarray experiments.

RNAi knockdown rck1/rck2 mutants produced almost no nodules (Chapter 4). This may be due to higher than normal ROS levels in the mutant than WT. ROS levels should be checked in the single mutants and in the double knockdown roots, this could be achieved through nitro blue tetrazolium (NBT) staining for •O₂⁻ or a ROS sensitive dye.
such as HyPer probe for $H_2O_2$ could be used to track dynamic changes during infection (Belousov et al., 2006).

The regulation of genes known to be induced during PTI should be checked in the \textit{rlck1} and \textit{rlck2} mutants and the \textit{rlck1/rlck2} knockdown roots after treatment with a symbiont such as AMF and a pathogen such as \textit{P. infestans}. Good candidates for this analysis are \textit{PR4} (Medtr1g080800) and \textit{PR5} (Medtr5g010635) which are pathogen-responsive genes which are repressed by NF and during infection by rhizobia, are initially induced and then repressed (Murray lab unpublished results).

**Regulation of developmental processes by ROS**

Data by Andrio et al. (2013) showing that \textit{RLCK1/SPK1} is upregulated within one hour of $H_2O_2$ application to the root, may suggest a role for \textit{RLCK1/SPK1} in the regulation of ROS during symbiosis promoting the correct development of symbiotic structures. Promoter-GUS analysis showed that \textit{RLCK1/SPK1} expression was correlated with rhizobial entry through the infection thread, in nodule primordia and in Zones I and II of the nodule. \textit{pRLCK2:GUS} expression shows that \textit{RLCK2} is higher in arbusculated cortical cells. All of the regions that promoter-GUS expression was seen for both \textit{RLCK1/SPK1} and \textit{RLCK2} have been previously identified as being rich in ROS. Misregulation of ROS, which is finely controlled in many plant processes, could cause disruption to the development of plant-microbe symbioses.

During symbiosis the plant undergoes many different developmental processes which require changes in hormone and ROS levels. During rhizobial symbiosis root hair cells change the direction of growth to grow towards the source of NF (Esseling et al., 2003). ROS production is required for apical growth of root hairs (Foreman et al., 2003). It is believed that ROS generation drives a calcium influx into the tip of the root hair creating a calcium gradient (Foreman et al., 2003). This calcium gradient is thought to give direction to tip growth by regulating exocytosis at the plasma membrane and actin reorganisation (Rato et al., 2004; He et al., 2006). Several ROS regulatory proteins such as NADPH oxidase, ROP GTPases and RopGEFs have been identified to be essential for correct root hair tip growth (Foreman et al., 2003; Craddock et al., 2012). This tip growth machinery is thought to be altered to enable the typical curling of the root hair around the rhizobia, which is supported by recent data from Shailes (2014) that showed a RopGAP mutants (\textit{gap1}) with a decreased number of ITs, which were thicker than wildtype and sometimes ramified, and had abnormal root hair curling. The formation of the IT in the root hair during rhizobial infection could be thought of as an internally directed version of
polar tip growth (Oldroyd et al., 2011). ROS has also been shown to be present in ITs (Santos et al., 2001; Ramu et al., 2002; Nanda et al., 2010). ROS in this case could be acting in two roles. Firstly ROS is required for the cross-linking of cell wall material (Gapper and Dolan, 2006) and could make the IT structure more rigid, creating a clear route for the rhizobia. Secondly, it could act as another layer of defence whilst letting the rhizobia into the root; rhizobia that are mutant in catalase and superoxide dismutase genes, that convert ROS into H$_2$O and O$_2$, are unable to successfully traverse the infection threads (Santos et al., 2000; Jamet et al., 2003). Interestingly, the overexpression of the rhizobial catalase gene KatB also leads to compromised infection (Jamet et al., 2007), which could be due to a lower level of ROS in the IT suggesting that a balance of ROS is needed for symbiotic infection. In the rlck1/spk1 mutant the infection threads are misshaped; it is possible that this could be caused by lower levels of ROS present in the IT.

PIT and PPA formation are very similar developmental processes and the progression of the AMF through the root surrounded by a plant derived tube is much like the passage of rhizobia through ITs. It is possible that the regulation of ROS during mycorrhizal colonisation is as crucial for this process as for the correct development of the IT (Balestrini and Bonfante, 2014; Rich et al., 2014). H$_2$O$_2$ has also been shown to accumulate in arbuscule cells (Fester and Hause, 2005). Again ROS could be involved in the hardening of cell wall material in the matrix surrounding the arbuscule, preventing the AMF from diverting from the plant determined route into the cell. Alternatively, ROS accumulation, and especially ROS transfer into the arbuscule, correlates with arbuscule senescence (Fester and Hause, 2005). In rlck1/spk1 and rlck2 mutants the arbuscules appear to be senescing early.

During rhizobial colonisation cortical cells must switch from their differentiated state and restart cell division to create the nodule, and in indeterminate nodules a meristem must be created to drive the growth of the nodule. ROS production has been seen in cortical cells after inoculation with S. meliloti (Peleg-Grossman et al., 2007; Nanda et al., 2010). ROS may be acting to strengthen the new cell walls after cell division. Alternatively, ROS gradients are seen in growing root tips with ROS levels being the highest at root meristems (Tsukagoshi et al., 2010; Wells et al., 2010); it is possible that there is a ROS gradient from the nodule meristem, as in root meristems, which determines the differentiation of the nodule cells. ROS have been found in Zone II, or infection zone, of the nodule (Andrio et al., 2013). RLCK1/SPK1 promoter-GUS analysis shows RLCK1/SPK1 expression in Zone I and II of the nodule and could be contributing to the regulation of ROS in these zones.
The phenotypes seen in the *rck1* mutant - early senescent arbuscules, a large number of infection events and the progress of ITs into Zone I of the nodule - suggest that in the absence of *RLCK1/SPK1* the plant is progressing through the symbiosis faster than it would in WT. It is possible that this faster progression is due to misregulation of ROS in these developmental processes.

**Future Work**

This study has provided some insight into the expression and requirement of *RLCK1/SPK1* and *RLCK2* during rhizobial and mycorrhizal symbioses. It has also created several expression constructs that can be used to identify interaction partners of *RLCK1/SPK1* and *RLCK2* in yeast and *in planta*. As the RNAi knock down *rck1/rck2* roots had a significant reduction in the number of nodules, the effect of the double knockdown should be investigated with AMF to help determine the role of these genes during symbiosis. Given that *RLCK1/SPK1* expression is induced by H$_2$O$_2$, the expression of *RLCK2* in response to ROS should be investigated in order to identify if this is a shared response of the two RLCKs. Also the levels of ROS in the *rck* single mutants and the double knockdown roots should be investigated both with and without symbiotic interactions to see if they are part of the ROS regulatory machinery. Localisation of the RLCK1/SPK1 and RLCK2 proteins via fluorescent fusions would also help to define their role during symbiosis. These constructs can also be used to visualise co-localisation of the RLCK proteins with potential interaction partners. The recent suggestion from protein prediction software of a lectin-like domain and a putative calmodulin binding domain at the N-terminal end of the proteins would also be worthy of further investigation; this could be achieved via a carbohydrate chip to determine possible ligands, and an assay to confirm the calmodulin binding ability of the RLCKs with and without the presence of calcium – the existing expression constructs for the RLCKs could be used for this.

**Conclusion**

*SCOoby* is a novel mycorrhizal mutant phenotype in *M. truncatula* with stunted arbuscule growth; further investigation could provide additional insight into the plant-mycorrhizal symbiosis. *RLCK1/SPK1* and *RLCK2* have a high nucleotide and amino acid similarity, and possibly have partially redundant or synergistic functions. Severe symbiotic defects are not evident in the single mutants and the full effect of *RLCK1/SPK1* and *RLCK2* mutations on symbiotic processes within the plant may only be evident in the absence of both proteins. This is supported by the extremely low nodule number on the double knockdown RNAi roots whilst the single mutants can still make nitrogen fixing nodules. They may be acting together, or redundantly, to control ROS during NF signalling,
potential PTI responses or symbiotic developmental processes such as nodule, IT or arbuscule formation.
## Appendix I

### Primers

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<th>Description</th>
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</tr>
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<td>RLCK2 promoter forwards with attB recombination site (bold)</td>
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| **Table A1.** List of PCR primers for cloning or genotyping |
### Table A2. List of primers for qPCR

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<td>TTAAGGCCATTCCACCACCTC</td>
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<td>60</td>
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<td>61</td>
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**Table A3.** List of sequencing primers
**RNAi Plasmid Maps**

**Figure A1.** pK7GW|WG2D(II)R RLCK1 RNAi – RNAi vector designed on *RLCK1*. The vector map for RNAi Construct 1 designed on a 99bp region of *RLCK1*. The vector was made by TOPO® and Gateway® cloning. Destination vector modified from pK7GW|WG2D(II).
**Figure A2.** pK7GW|WG2D(II)R RLCK2 RNAi – RNAi vector designed on the 5' UTR region of *RLCK2*.

The vector map for RNAi Construct 2 designed on the 5' UTR region of *RLCK2*. The vector was made by TOPO® and Gateway® cloning. Destination vector modified from pK7GW|WG2D(II).
Golden Gate constructs

<table>
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<th>ENSA Standard name</th>
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<td>pLOM-PU-pMtRLCK2-67033</td>
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Table A4. List of Level 0 Golden gate modules

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Table A5. List of Level 1 Golden Gate modules
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**Table A6.** List of Level 2 Golden Gate constructs
Golden Gate Plasmid Maps

Figure A3. EC67001 plasmid map
Level 0 Golden Gate SC1 construct. Contains RLCK1 CDS with no stop codon.
Figure A4. EC67002 plasmid map
Level 0 Golden Gate SC1 construct. Contains *RLCK2* CDS with no stop codon.
Figure A5. EC67003 plasmid map
Level 0 Golden Gate SC1 construct. Contains MtNFP CDS with no stop codon.
Figure A6. EC67004 plasmid map
Level 0 Golden Gate SC1 construct. Contains MtLYK3 CDS with no stop codon.
Figure A7. EC67005 plasmid map
Level 0 Golden Gate SC1 construct. Contains *MtHMGR1* CDS with no stop codon.
Figure A8. EC67006 plasmid map

Level 0 Golden Gate SC1 construct. Contains MtDMI2 CDS with no stop codon.
Figure A9. EC67007 plasmid map

Level 1 module Golden Gate construct. 35S Promoter, RLCK1 CDS, 3x myc c-terminal tag, 35S terminator. Compatible with position R1 of Golden Gate Level 2 constructs.
Figure A10. EC67008 plasmid map

Level 1 module Golden Gate construct. 35S Promoter, RLCK2 CDS, 3x myc C-terminal tag, 35S terminator. Compatible with position R1 of Golden Gate Level 2 constructs.
Figure A11. EC67009 plasmid map
Level 1 module Golden Gate construct. *L. japonicus* Ubiquitin Promoter, *MtNFP* CDS, C-terminal GFP tag, NOS terminator. Compatible with position R2 of Golden Gate Level 2 constructs.
**Figure A11.** EC67009 plasmid map

Level 1 module Golden Gate construct. *L. japonicus Ubiquitin* Promoter, *MtNFP* CDS, C-terminal GFP tag, NOS terminator. Compatible with position R2 of Golden Gate Level 2 constructs.
**Figure A13.** EC67011 plasmid map

Figure A14. EC67012 plasmid map

Level 1 module Golden Gate construct. *L. japonicus Ubiquitin* Promoter, *MtDMI2* CDS, C-terminal GFP tag, NOS terminator. Compatible with position R2 of Golden Gate Level 2 constructs.
Figure A15. EC67013 plasmid map

Level 1 module Golden Gate construct. *L. japonicus* Ubiquitin Promoter, *MtRLCK2* CDS, C-terminal GFP tag, NOS terminator. Compatible with position R2 of Golden Gate Level 2 constructs.
Figure A16. EC67014 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, RLCK1 CDS, 3x myc c-terminal tag, 35S terminator. In position R2: L. japonicus Ubiquitin Promoter, MtNFP CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
**Figure A17.** EC67015 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, RLCK1 CDS, 3x myc c-terminal tag, 35S terminator. In position R2: *L. japonicus* Ubiquitin Promoter, *MtLYK3* CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
Figure A18. EC67016 plasmid map
Level 2 Golden Gate vector. In position R1: 35S Promoter, RLCK1 CDS, 3x myc c-terminal tag, 35S terminator. In position R2: *L. japonicus* Ubiquitin Promoter, *MtHMGR1* CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
Figure A19. EC67017 plasmid map

**Figure A20.** EC67018 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, RLCK1 CDS, 3x myc c-terminal tag, 35S terminator. In position R2: *L. japonicus Ubiquitin* Promoter, RLCK2 CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
**Figure A21.** EC67019 plasmid map

Figure A22. EC67020 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, RLCK2 CDS, 3x myc c-terminal tag, 35S terminator. In position R2: L. japonicus Ubiquitin Promoter, MtLYK3 CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
Figure A23. EC67021 plasmid map

**Figure A23.** EC67021 plasmid map

Figure A25. EC67030 plasmid map
Level 0 Golden Gate C construct. Contains \textit{RLCK1} CDS with a stop codon.
Figure A26. EC67031 plasmid map
Level 0 Golden Gate C construct. Contains *RLCK2* CDS with a stop codon.
Figure A27. EC67032 plasmid map
Level 0 Golden Gate PU construct. Contains *RLCK1* promoter region.
Figure A28. EC67033 plasmid map
Level 0 Golden Gate PU construct. Contains \textit{RLCK2} promoter region.
Figure A29. EC67034 plasmid map
Level 1 module Golden Gate construct. 35S Promoter, 3x myc N-terminal tag RLCK1 CDS, 35S terminator. Compatible with position R1 of Golden Gate Level 2 constructs.
Figure A30. EC67035 plasmid map

Level 1 module Golden Gate construct. 35S Promoter, 3x myc N-terminal tag RLCK2 CDS, 35S terminator. Compatible with position R1 of Golden Gate Level 2 constructs.
Figure A31. EC67036 plasmid map

Level 1 module Golden Gate construct. *L. japonicus* Ubiquitin Promoter, N-terminal GFP tag *RLCK2* CDS, NOS terminator. Compatible with position R2 of Golden Gate Level 2 constructs.
Figure A32. EC67037 plasmid map

Level 1 module Golden Gate construct. *L. japonicus Ubiquitin* Promoter, N-terminal GFP tag *RLCK1 CDS*, NOS terminator. Compatible with position R2 of Golden Gate Level 2 constructs.
Figure A33. EC67038 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, 3x myc N-terminal tag RLCK1 CDS, 35S terminator. In position R2: L. japonicus Ubiquitin Promoter, MtNFP CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
Figure A34. EC67039 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, 3x myc N-terminal tag RLCK1 CDS, 35S terminator. In position R2: *L. japonicus* *Ubiquitin* Promoter, *MtLYK3* CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
Figure A35. EC67040 plasmid map

Figure A36. EC67041 plasmid map

**Figure A37.** EC67042 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, 3x myc N-terminal tag $RLCK1$ CDS, 35S terminator. In position R2: *L. japonicus* Ubiquitin Promoter, $RLCK2$ CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
**Figure A38.** EC67043 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, 3x myc N-terminal tag RLCK2 CDS, 35S terminator. In position R2: *L. japonicus* Ubiquitin Promoter, N-terminal GFP tag, RLCK2 CDS, NOS terminator. Endlinker 2 in position F3.
Figure A39. EC67044 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, 3x myc N-terminal tag RLCK2 CDS, 35S terminator. In position R2: *L. japonicus* Ubiquitin Promoter, MtNFP CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
**Figure A40.** EC67045 plasmid map

Level 2 Golden Gate vector. In position R1: 35S Promoter, 3x myc N-terminal tag RLCK2 CDS, 35S terminator. In position R2: *L. japonicus Ubiquitin* Promoter, MtLYK3 CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
**Figure A41.** EC67046 plasmid map

Figure A42. EC67047 plasmid map
Level 2 Golden Gate vector. In position R1: 35S Promoter, 3x myc N-terminal tag RLCK2 CDS, 35S terminator. In position R2: *L. japonicus* Ubiquitin Promoter, MtDMI2 CDS, C-terminal GFP tag, NOS terminator. Endlinker 2 in position F3.
References


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