REVIEW ARTICLE

Receptor Kinases in Plant-Pathogen Interactions: More than Pattern Recognition

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Short title: Receptor kinases in disease resistance

One-sentence summary: A current view of the role and regulation of plant receptor kinases that act as innate immune receptors or an Achilles’ heel targeted by phytopathogens.

ABSTRACT

Receptor-Like Kinases (RLKs) and Receptor-Like Proteins (RLPs) play crucial roles in plant immunity, growth and development. Plants deploy a large number of RLKs and RLPs as Pattern Recognition Receptors (PRRs) that detect microbe- and host-derived molecular patterns as the first layer of inducible defense. Recent advances have uncovered novel PRRs, their corresponding ligands, and mechanisms underlying PRR activation and signaling. In general, PRRs associate with other RLKs and function as part of multi-protein immune complexes at the cell surface. Innovative strategies have emerged for the rapid identification of microbial patterns and their cognate PRRs. Successful pathogens can evade or block host recognition by secreting effector proteins to “hide” microbial patterns or inhibit PRR-mediated signaling. Furthermore, newly identified pathogen effectors have been shown to manipulate RLKs controlling growth and development by mimicking peptide hormones of host plants. The ongoing studies illustrate the importance of diverse plant RLKs in plant disease resistance and microbial pathogenesis.
INTRODUCTION

A longstanding question in plant biology is how plants perceive and ward off attack by numerous potential pathogens. Two parallel paths were taken to address this question in the last century. The first path follows Harold Flor’s famous gene-for-gene hypothesis, which predicts that resistance gene products in the host plant specifically recognize cognate “avirulence” gene products in the pathogen to confer disease resistance. This led to the discovery that plant resistance proteins broadly fall into two classes according to their protein sequences and subcellular localization, cytoplasmic Nucleotide binding domain Leucine-rich repeat domain-containing Receptors (NLRs; Jones et al., 2016) and cell surface-localized receptors. The latter are proteins belonging to large families of Receptor-Like Kinases (RLKs) and Receptor-Like Proteins (RLPs) (Jones and Dangl, 2006). Accordingly, their corresponding “avirulence” proteins were identified as effector proteins targeted to the plant cytosol or apoplast. The second path follows the findings that microbe-derived and host-derived elicitors, also called Pathogen/microbe-Associated Molecular Patterns (PAMPs) and Damage-Associated Molecular Patterns (DAMPS), respectively, can act as danger signals to trigger defenses in plants, which culminated in the identification of RLKs and RLPs as Pattern Recognition Receptors (PRRs) for direct recognition of the patterns (Boller and Felix, 2009). These combined efforts led to the consensus that the plant surveillance system is primarily composed of NLRs and PRRs. Thus, NLRs are responsible for the recognition of cytoplasmic effectors, whereas PRRs are responsible for the detection of apoplastic effectors or patterns.

Unlike animals, which employ receptor tyrosine kinases (RTKs), seven transmembrane G protein-coupled receptors (GPCRs) and Toll-Like Receptors at the plasma membrane to perceive growth hormones, environmental signals and danger signals derived from pathogens, plants rely on RLKs (~410 in Arabidopsis) and RLPs (~170 in Arabidopsis) to fulfill these diverse roles (Shiu and Bleecker, 2001; Shiu et al., 2004; Li et al., 2016a). A plant RLK is analogous to an animal RTK and contains an ectodomain (ECD), a single pass transmembrane domain and a cytoplasmic kinase domain, whereas an RLP is essentially an RLK lacking a cytoplasmic kinase domain. The ECDs of RLKs and RLPs are highly variable, providing means to recognize a wide range of ligands, including steroids, peptides, polysaccharides, and lipopolysaccharides. While some RLKs and RLPs are known to act as PRRs that perceive danger signals (Table 1), others regulate plant growth and development, reproduction, symbiosis, and tolerance to abiotic stresses (Breiden and Simon, 2016).
Pathogenic microbes, including bacteria, fungi, oomycetes, and nematodes deliver a large number of effector proteins into the plant apoplast or cytosol (Dou and Zhou, 2012). Analyses of these effectors have uncovered a variety of mechanisms that pathogens use for infection and colonization of host plants. Often, these proteins provide means to evade PRR-mediated surveillance by camouflage or blockage of PRR-induced signaling. Others can mimic host growth factors and manipulate specific RLK-mediated host processes for their benefit. On the other hand, perturbation by these effectors imposes a selection pressure on host plants, which ultimately leads to the evolution of NLRs or PRRs recognizing these effectors as danger signals and conferring disease resistance in plants.

A major effort in plant–pathogen interaction research has been to identify PRRs, their corresponding ligands, and mechanisms by which different PRRs activate defenses. While genetic and reverse genetic studies have proved straightforward in the identification of a large number of RLKs and RLPs involved in disease resistance to diverse pathogens, identification of molecular patterns and their corresponding PRRs has been much more challenging. Nonetheless, conventional biochemical studies and genetic analyses have identified important founding members of microbial and plant molecular patterns and their corresponding PRRs. The advent of microbial and plant genomics has enabled the development of novel strategies for the discovery of new microbial molecular patterns and PRRs. Furthermore, molecular genetics, biochemistry, and structural biology studies are uncovering generalities and differences in mode of action among different types of PRRs, how PRRs activate downstream defenses, cross-talk between different receptor kinase pathways, and mechanisms by which various pathogens perturb receptor kinase signaling for pathogenesis.

In this review, we provide an update on PRRs involved in disease resistance and mechanisms by which different types of PRRs recognize ligands to form active receptor complexes. We then discuss how the knowledge gained has led to the development of strategies for identifying PAMPs and PRRs. For PRR-triggered defenses, we focus on early signaling mediated by PRRs. Readers are referred to several excellent reviews on the regulation of downstream signaling, such as transcriptional controls (Bigeard et al., 2015; Couto and Zipfel, 2016; Li et al., 2016b). We also discuss how pathogen effectors interfere with early PRR signaling to inhibit defenses and how pathogen effectors enhance virulence by manipulating RLKs that do not function as PRRs.
RLKs AND RLPs INVOLVED IN PLANT IMMUNITY

Numerous RLKs and RLPs have been shown to function in plant disease resistance, but only a handful of them are confirmed to function as pattern recognition receptors (PRRs) with known ligands (Wu and Zhou, 2013; Bohm et al., 2014; Zipfel, 2014; Couto and Zipfel, 2016). In the following text, RLKs and RLPs known to directly bind ligands are referred to as RKs and RPs, respectively. An extended list of known PRRs and other RLKs and RLPs involved in plant immunity is presented in Tables 1 and 2.

PRRs can be classified into different subfamilies according to domains or motifs in their ECDs: Leucine-Rich Repeat (LRR) domain, Lysine Motifs (LysM), Lectin domain, or Epidermal Growth Factor (EGF)-like domain (Wu and Zhou, 2013; Bohm et al., 2014; Zipfel, 2014; Couto and Zipfel, 2016). All known LRR-containing PRRs, including LRR-RKs and LRR-RPs, bind proteins or peptides. For example, the Arabidopsis LRR-RKs FLAGELLIN SENSING2 (FLS2) and EFR recognize a conserved 22-amino acid epitope (flg22) of the N-terminus of the bacterial flagellin and a conserved N-terminal epitope (elf18) of the bacterial elongation factor Tu (EF-Tu), respectively (Gomez-Gomez and Boller, 2000; Bauer et al., 2001; Kunze et al., 2004; Zipfel et al., 2006). The Arabidopsis LRR-RKs PEP RECEPTOR1 (PEPR1) and PEPR2 perceive proteinaceous DAMPs called Peps (Plant elicitor peptides), which are conserved epitopes of a small family of pro-peptides (PROPEPs) (Huffaker et al., 2006; Huffaker and Ryan, 2007; Krol et al., 2010; Yamaguchi et al., 2010; Liu et al., 2013a; Yamada et al., 2016a). ProPep proteins are produced and released through an unknown mechanism into the extracellular space upon pathogen infection, in a manner comparable to the production and secretion of inflammatory cytokines in animals after pathogen infection (Yamada et al., 2016a). In Oryza sativa (rice), LRR-RK XA21 recognizes RaxX, a highly conserved protein in many Xanthomonas species, to trigger immune responses. RaxX21-sY, a sulfated, 21-amino acid synthetic peptide derived from RaxX, is sufficient for XA21 activation (Pruitt et al., 2015).

Several LRR-RLPs have been shown to recognize proteinaceous patterns. Among the 57 LRR-RLPs encoded by the Arabidopsis genome (Wang et al., 2008), RLP23 specifically binds and recognizes nlp20, a conserved amino-acid peptide from necrosis and ethylene-inducing peptide1-like proteins (NLPs), which are widespread proteins among diverse group of microbes, including bacteria, fungi and oomycetes (Albert et al., 2015). In Solanum lycopersicum (tomato), the LRR-RLPs Cf-2, CF-4 and Cf-9 confer resistance to the fungal pathogen Cladosporium fulvum by recognizing the effectors Avr2, Avr4 and Avr9, respectively (Dixon et al., 1996; Kruger et al., 2002; Luderer et al., 2002; Rooney
et al., 2005). It is not known whether any of these apoplastic effectors directly bind to the RLPs. However, Avr2 is known to inhibit several cysteine proteases, including Rcr3 for virulence, and the Cf-2 protein indirectly recognizes Avr2 likely by sensing the modification of Rcr3 (Rooney et al., 2005; van Esse et al., 2008). In this case, the Avr2-modified Rcr3 is equivalent to a DAMP that activates immunity in plants carrying Cf-2.

LysM-RPs and LysM-RKs represent a major class of receptors for microbial N-acetylglucosamine (GlcNAc)-containing glycans, including fungal chitin and bacterial peptidoglycan (PGN), which trigger defenses, and Rhizobium- and arbuscular mycorrhiza-derived lipochitooligosaccharides (LCOs), which trigger symbioses (Gust et al., 2012). Three Arabidopsis LysM-RLKs, CHITIN ELICITOR RECEPTOR KINASE 1 (CERK1), LysM-CONTAINING RECEPTOR KINASE 4 (LYK4), and LYK5, are required for chitin signaling (Miya et al., 2007; Wan et al., 2008; 2012; Cao et al., 2014), whereas CERK1 and two Arabidopsis LysM-RLPs LysM DOMAIN-CONTAINING GPI-ANCHORED PROTEIN 1 (LYM1) and LYM3 are required for response to PGN (Willmann et al., 2011). Both CERK1 and LYK5 have been shown to bind chitin, with LYK5 reported to have greater affinity (Liu et al., 2012b; Cao et al., 2014), whereas LYM1 and LYM3 were shown to bind PGN (Willmann et al., 2011). The rice LysM-RP CHITIN OLIGOSACCHARIDE ELICITOR-BINDING PROTEIN (CEBiP) and the rice counterpart of CERK1 are essential for chitin-induced defenses in plants (Kaku et al., 2006; Shimizu et al., 2010). CEBiP binds chitin with high affinity, indicating that it is the chitin receptor in rice plants. The perception of chitin by the Arabidopsis LysM-RK LYK5 and rice LysM-RP CEBiP echoes the recent findings that bacterial cold shock protein epitope csp22 is perceived by the tomato LRR-RK CORE and Nicotiana tabacum (tobacco) LRR-RP NbCSPR (Saur et al., 2016; Wang et al., 2016b). Interestingly, the peptide hormone CLAVATA3 (CLV3p) is perceived by not only the LRR-RK CLV1, but also by a likely LRR-RP CLV2, although direct ligand binding has been shown only for CLV1 (Soyars et al., 2016). Thus, an RK and an RP can perceive the same ligand with a shared ECD.

Lectin-RLKs are uniquely important for plant disease resistance. The Arabidopsis lectin-RK LORE is a receptor for the lipid A moiety of bacterial lipopolysaccharides (LPS) (Ranf et al., 2015). Another Lectin-RK, DORN1 (Choi et al., 2014), is a high affinity receptor for ATP (eATP), which may act as a DAMP as it induces cytosolic Ca\(^{2+}\) elevation and MAPK activation (Tanaka et al., 2014). It remains to be determined whether eATP contributes to plant immunity in response to pathogens.
Upon ligand-binding, PRRs of the LRR-RK class recruit BRI1-ASSOCIATED RECEPTOR KINASE (BAK1), a LRR-RLK with only five LRRs and member of the SOMATIC EMBRYOGENESIS RECEPTOR KINASES (SERKs) (Figure 1) (Couto and Zipfel, 2016). Thus, the binding of FLS2 to flg22 recruits BAK1 to form a hetero-dimer, resulting in rapid phosphorylation of both FLS2 and BAK1 (Chinchilla et al., 2007; Heese et al., 2007; Schulze et al., 2010; Roux et al., 2011). Molecular and genetic studies showed that SERKs are also recruited to EFR, PEPRs, and Xa21 upon ligand binding and required for signaling (Roux et al., 2011; Chen et al., 2014; Yamada et al., 2016a). In addition, LRR-RKs for peptide hormones, including tracheary element differentiation inhibitory factor (TDIF), INFLORESCENCE DEFICIENT IN ABSCISSION (IDA), PHYTOSULFOKINE (PSK), and ROOT GROWTH FACTORS (RGFs), also require SERKs for function (Ma et al., 2016; Zhang et al., 2016). LRR-RKs generally contain many more LRRs (28 for FLS2 and 27 for PEPR1), which provide an interface for peptide-binding, than do SERKs. Crystal structures have been solved for the ECDs of FLS2, PEPR1, RGF receptor RGFR1, PSK receptor PSKR1, IDA receptor HAESA, and TDIF receptor PXY (phloem intercalated with xylem) in complex with their respective ligands and/or a SERK ECD (Sun et al., 2013; Tang et al., 2015; Song et al., 2016; Santiago et al., 2016; Wang et al., 2016a; Zhang et al., 2016). The FLS2-flg22-BAK1 complex showed that flg22 adopts a linear conformation and binds to FLS2 LRR3-16 in the same N-terminal to C-terminal orientation (Figure 1; Sun et al., 2013). Strikingly, Gly18 of flg22 fits in an “inner-curved loop” between Thr52 and Val54 at the N-terminus of the BAK1 ECD, and this interaction is further stabilized by hydrogen bonds between flg22 Leu19 and BAK1 Thr52 and Val54. This interaction effectively bridges the dimerization between FLS2 and BAK1, demonstrating that BAK1 is a co-receptor for flg22. FLS2 LRR23-26 additionally interacts with a cluster of bulky BAK1 amino acids, forming a stable complex. The binding of flg22 does not lead to a conformational change in the FLS2. The PEPR1-Pep1, HAESA-IDA-SERK1, PXY-TDIF-SERK2, and RGFR1-RGF1-BAK1 complexes also adopt a conformation strikingly similar to the FLS2-flg22-BAK1 complex (Figure 1; Tang et al., 2015; Santiago et al., 2016; Song et al., 2016; Zhang et al., 2016). The PSKR1-PSK-SERK1 structure showed a different mechanism for ligand-binding and receptor complex formation. PSK presents as a β-strand and forms an anti-parallel β-sheet conformation with the island domain of PSKR1 ECD (Wang et al., 2016a). This induces an allosteric change in PXY, allowing the recruitment of SERKs. Taken together, SERKs are likely common co-receptors for many LRR-RKs.
perceiving proteinaceous ligands.

Although LRR-RPs do not carry a cytoplasmic kinase domain, they associate with RLKs to transmit the signal to downstream components. The LRR-RLK SUPPRESOR of BIR1 (SOBIR1), originally identified as a component required for autoimmunity caused by loss-of-function of BAK1-INTERACTING RECEPTOR-LIKE KINASE 1 (BIR1) (Gao et al., 2009), functions as a common adaptor for multiple LRR-RP-type PRRs (Gust and Felix, 2014). In tomato, SOBIR1 interacts with a number of RLPs, including Ve1, EIX1 and Cf-4, and plays crucial roles in plant immunity against fungal infection (Liebrand et al., 2013, 2014). RLP23 forms a complex with SOBIR1 in the resting state and recruits BAK1 into the complex after nlp20 recognition to activate immune signaling (Albert et al., 2015). RLP30 associates with SOBIR1, and SCFE1-triggered immune responses are dependent on BAK1 and SOBIR1 (Zhang et al., 2013). In addition, SOBIR1 is required for PG-triggered, RBPG1-mediated immune responses (Zhang et al., 2014), and ELR associates with BAK1 for elicitin recognition (Du et al., 2015). The consensus is that LRR-RPs form a complex with SOBIR1 before ligand-binding and recruit BAK1 to form an active receptor complex upon ligand-binding.

Chitin-binding can induce homo-dimerization of the LysM-ECDs of CEBiP and CERK1, which is essential for chitin-induced defenses in Arabidopsis (Shimizu et al., 2010; Liu et al., 2012b; Hayafune et al., 2014). Two competing models have been proposed for chitin-induced dimerization of LysM-ECDs. Chitin oligomers longer than six sugars are required for both triggering defenses and inducing dimerization of CERK1 or CEBiP, whereas only three sugar residues are directly bound to CERK1-ECD in structural analyses (Liu et al., 2012b). Liu et al., 2012b thus proposed a “cross-linking” model in which the two ends of long chitin oligomers each bind a LysM-ECD to form a dimer. In a separate NMR study assisted by synthetic chemistry, N-acetyl groups of alternating sugars were found to be located on two opposite sides of the chitin chain and directly bound by the LysM-ECD (Hayafune et al., 2014). Depleting N-acetyl groups from one side of the chitin chain did not affect chitin binding to a CEBiP-ECD, but abolished dimerization. This led to the proposal of a "sandwich" model in which two CEBiP-ECDs bind the chitin chain, one on each side (Hayafune et al., 2014). However, this does not explain why shorter chitin oligomers do not induce dimerization. Whether CEBiP possesses significantly different affinities toward short and long chitin oligomers remains controversial, in part because different fragments of CEBiP-ECD were used in different studies (Hayafune et al., 2014; Liu et al., 2016a). Moreover, the sandwich model predicts that five chitin sugars are required for binding
to one CEBiP, whereas the recent crystal structure data clearly showed only three are bound (Liu et al., 2016a). Liu et al. (2016a) thus proposed a third “sliding model” in which two CEBiP-ECDs slide along a longer chitin chain involving alternating N-acetyl groups from opposite sides for optimum interaction (Liu et al., 2016a), a model explaining both structural data and N-acetyl depletion experiments.

In rice plants, chitin-binding to CEBiP recruits CERK1 to form a CEBiP-CERK1 hetero-complex (Shimizu et al., 2010). In Arabidopsis, chitin treatment was reported to induce CERK1-dimerization only in the presence of LYK5 (Cao et al., 2014). LYK5 is present as homodimer before and after chitin perception. LYK5 kinase activity is not required for chitin signaling, but the intact kinase domain is essential for chitin-induced LYK5/CERK1 association (Cao et al., 2014). How CERK1 and LYK5 work together to recognize chitin remains to be determined. In sum, these findings support that CEBiP and LYK5 are chitin receptors and recruit CERK1 to form active receptor complexes (Figure 1). The requirement of CERK1 for PGN signaling in Arabidopsis (Willmann et al., 2011) and AM symbiosis in rice (Zhang et al., 2015) suggests that CERK1 is similarly recruited to PGN and LCO receptors. Thus, CERK1 appears to be a universal component for the perception of chitin, PGN, and LCOs. CERK1 may be functionally analogous to BAK1 and may act as a co-receptor for these highly related ligands, a possibility that remains to be tested by structural studies.

INNOVATIVE APPROACHES TO IDENTIFY MICROBIAL PATTERNS/APOPLASTIC EFFECTORS AND THEIR COGNATE RECEPTORS

Identification of microbial patterns, apoplastic effectors and PRRs through which they activate immunity remains challenging. While most of the microbial patterns and immune-triggering apoplastic effectors reported to-date were identified through classical biochemical purification and genetics, these approaches are time consuming and require extensive expertise. For proteinaceous microbial patterns and apoplastic effectors recognized by host PRRs, their coding genes often display a positive selection on individual amino acids as a result of adaptation. For instance, *X. campestris campestris* isolates frequently accumulate mutations in the flagellin-coding gene that evade detection by FLS2 (Sun et al., 2006). This unique feature can be exploited to rapidly identify new effectors or microbial patterns by interrogating microbial genome sequences. As such, a second 28-amino acid epitope (flgII-28) of bacterial flagellin and Ave1 of *Verticillium dahliae* were found to be subject to strong positive selection (Cai et al., 2011; de Jonge et al., 2012). These findings directly led to the discoveries of flgII-28 as a
pattern recognized by LRR-RK FLS3 (see below) and of Ave1 as an effector recognized by LRR-RLP Ve1 in tomato plants. Most recently this approach was applied to identify six new peptides, including xup25, from *Pseudomonas syringae* that trigger immune responses in Arabidopsis (Mott et al., 2016). This important work further illustrates the power of the comparative genomics approach in revealing novel microbial patterns and in isolating their corresponding PRRs.

Like microbial patterns and pathogen effectors, plant PRRs are also evolving rapidly. FLS2, EFR, FLS3, XPS1, and Xa21 all belong to LRR-RLK XII, a subfamily that has undergone significant gene expansion (Shiu et al., 2004). Functional EFR and Xa21 occur only in Brassicaceae and certain rice cultivars, respectively (Boller and Felix, 2009; Lacombe et al., 2010). Similarly, flgII-28 only elicits immune responses in a few species of the Solanaceae, such as tomato, *Solanum tuberosum* (potato), and *Capsicum annuum* (pepper) (Clarke et al., 2013; Hind et al., 2016). Natural variations in flgII-28 responses among tomato varieties were exploited to isolate FLS3 as the flgII-28 receptor. Lineage-specific expansion also appears to occur in LRR-RLP types of PRRs. Ve- and Cf-mediated resistance only occurs in some tomato cultivars, whereas RLP23, RLP30 and ReMax/RLP1 occur only in Brassicaceae species (Jehle et al., 2013; Zhang et al., 2013; Albert et al., 2015). A recent success story is the identification of CORE as a receptor of csp22 (Wang et al., 2016b). csp22 sensitivity was observed in many, but not all, Solanaceae species. Exploitation of natural variation among cultivated and wild tomato species allowed the isolation of CORE, which is required for csp22 perception. Transformation of CORE into Arabidopsis, which lacks CORE, confers csp22 sensitivity and disease resistance to *P. syringae*. Thus, the lineage-specificity of PRRs will not only facilitate the future identification of PRRs, but also allow deployment of disease resistance in crop plants by transgenic expression of PRRs from different plant species (Lacombe et al., 2010; Albert et al., 2015).

The discovery of BAK1 as a common co-receptor for LRR-RKs and LRR-RPs is particularly useful for the identification of receptors for protein ligands. csp22-induced defenses require BAK1, suggesting that an LRR-RK or LRR-RP is involved. BAK1 was thus used as a bait to identify the hypothetical PRR in *Nicotiana benthamiana* that interacts with BAK1 only in the presence csp22 (Saur et al., 2016). Indeed, mass spectrum analysis identified NbCSPR as a novel LRR-RLP associated with BAK1 in csp22-treated tissue. Subsequent genetic analyses supported NbCSPR as a likely LRR-RP for csp22. The findings that both CORE and NbCSPR perceive csp22 suggest that the two types of PRRs evolve independently for the perception of a single PAMP. The approach using BAK1 as a bait is
particularly useful in the rapid identification of new PRRs, as BAK1 has been shown to be required for defenses triggered by many pathogens, including aphids (Heese et al., 2007; Chaparro-Garcia et al., 2011; Korner et al., 2013; Larroque et al., 2013; Prince et al., 2014).

In addition to using BAK1 as bait for receptor identification, recent advances in LRR-ECD-ligand complex structures also provide novel means for receptor identification. A PEPR-Pep structure suggests that a ligand-recognition motif consisting of Arg-x-Arg in the ECD of PEPR directly recognizes the C-terminal residue Asn of Pep1. This signature motif is conserved in subfamily XI of the LRR-RLKs in Arabidopsis. Interestingly, several members of this subfamily have been shown to bind peptides ending with an Asn or His. Members of a rather large family of Arabidopsis peptide hormones carries an Asn or His residue at their C-termini, suggesting that additional members of the XI subfamily could be receptors for these peptides. Song et al. (2016) tested if the ECD of a previously uncharacterized member of this subfamily could bind a specific peptide from a pool of synthetic peptides. Mass-spectrum analysis indicated that this LRR-RK specifically binds RGFs. Subsequent genetic analysis showed that this LRR-RK and four additional members of this clade, namely RGF RECEPTORS 1–5 (RGFR1–5), are required for RGF-regulated root growth. Structural studies showed that the specificity of RGFRs in RGF-binding lies in a unique signature motif Arg-x-Gly-Gly that specifically recognizes the sulfated tyrosine in the RGF peptide. This elegant work illustrates how knowledge gained from structural studies can guide future identification of receptor–ligand pairs (Song et al., 2016).

**DYNAMIC CONTROL OF PRR COMPLEXES**

Besides the co-receptors mentioned above, receptor-like cytoplasmic kinases (RLCKs), are the major components of PRR complexes, which are involved in transducing signals from extracellular ligand perception into downstream signaling by phospho-relay (Wu and Zhou, 2013; Bohm et al., 2014; Macho and Zipfel, 2014; Couto and Zipfel, 2016). Several Arabidopsis RLCKs associate with PRRs and play important roles in PTI. For example, BOTRYTIS-INDUCED KINASE 1 (BIK1) (Veronese et al., 2006), a member of Arabidopsis RLCK subfamily VII, associates with FLS2 and BAK1, in the absence of ligand elicitation (Lu et al., 2010; Zhang et al., 2010a). Upon flg22 elicitation, BAK1 associates with FLS2 and phosphorylates BIK1 (Lu et al., 2010; Zhang et al., 2010a). BIK1 then dissociates from the PRR complex to activate downstream signaling. In addition, BRASSINOSTEROID-SIGNALING KINASE 1 (BSK1), a member of the RLCK XII subfamily and
previously reported to be a substrate of BRI1 (Tang et al., 2008), also associates with FLS2 to regulate flg22-induced immune responses (Shi et al., 2013).

Immune receptor complexes are subject to dynamic regulation to allow tight control of the intensity and duration of plant immune responses, as inappropriate or unnecessary activation of PRR-mediated immune responses will lead to developmental and growth defects (Couto and Zipfel, 2016). PRR complexes are actively maintained in a resting state in the absence of patterns. For example, BIR2, a LRR-RLK with a pseudokinase domain, associates with BAK1 to prevent BAK1-FLS2 complex formation in the resting state (Halter et al., 2014). Upon perception of flg22, BIR2 release BAK1 to facilitate FLS2-BAK1 interaction. BIR2 can be phosphorylated by the BAK1 kinase domain \textit{in vitro} (Halter et al., 2014). BIR1 is a BIR2-related RLK, and also functions as a negative regulator in plant immunity (Gao et al., 2009). The \textit{bir1} mutant displays constitutive cell death and defense responses, which are partially dependent on BAK1 and SOBIR1. Interestingly, BAK1 and SOBIR1 constitutively associate with each other only in seedlings lacking BIR1, suggesting that BIR1 prevents the interactions between BAK1 and SOBIR1 in the resting state (Liu et al., 2016b; Gao et al., 2009).

The phosphorylation of PRR complex components is critical for receptor activation and is subject to tight regulation (Couto and Zipfel, 2016). In Arabidopsis, several protein phosphatases negatively regulate PRR activation. The Arabidopsis Ser/Thr phosphatase type 2A (PP2A) negatively regulates EFR-mediated defenses by controlling the phosphorylation levels of BAK1 (Segonzac et al., 2014), although it is not clear whether PP2A directly dephosphorylates BAK1. Similarly, the partially redundant Arabidopsis protein phosphatases type 2C (PP2C) POLTERGEIST-LIKE 4 (PLL4) and PLL5, associate with EFR to negatively regulate PRR-mediated responses and bacterial resistance (Holton et al., 2015). PLL4 and PLL5 are close homologs of rice XANTHOMONAS RESISTANCE 21 (XA21) BINDING PROTEIN15 (XB15), which was previously shown to dephosphorylate XA21 \textit{in vitro} and negatively affect XA21-mediated immunity to the bacterium \textit{X. oryzae pv. oryzae} (Xoo) (Park et al., 2008). These results also indicate conserved regulatory mechanisms for Xa21 and EFR, a notion supported by XA21-EFR domain swapping experiments (Holton et al., 2015; Schwessinger et al., 2015).

Protein accumulation of the PRR immune components is also under tight control. Two closely related Arabidopsis U-box (PUB) family of E3 ubiquitin ligases, PUB12 and PUB13, were reported to associate with BAK1 in the resting state. Upon flg22 elicitation, BAK1 phosphorylates PUB12 and
PUB13, which then associate with and promote degradation of FLS2 through ubiquitination. Loss-of-function mutations in \textit{PUB12} and \textit{PUB13} enhance immune responses to flg22 and disease resistance (Lu et al., 2011).

Protein accumulation of PRRs is also affected by endocytosis (Ben Khaled et al., 2015; Frescatada-Rosa et al., 2015; Couto and Zipfel, 2016). Flg22 induces FLS2 internalization and degradation in Arabidopsis (Robatzek et al., 2006). Clathrin-dependent endocytosis and subcellular compartmentalization of FLS2 is critical for pattern-triggered immunity, and requires flg22-induced recruitment of BAK1 (Chinchilla et al., 2007; Beck et al., 2012; Mbengue et al., 2016). Flg22-induced internalization and degradation of FLS2 coincides with signal desensitization, which was proposed to play an important role in turning over ligand-occupied FLS2 by replenishing newly synthesized signal-competent FLS2 during subsequent infections (Smith et al., 2014). PEPR1/2 and EFR are also internalized in a ligand-dependent manner with a similar regulation mechanism (Mbengue et al, 2016; Ortiz-Morea et al 2016; Yamada et al 2016a). Similarly, Cf-4- and Cf-9-mediated resistance and endocytosis in tomato plants require ligand-induced recruitment of BAK1, indicating that diverse classes of PRRs share similar requirements for initiation of resistance and endocytosis (Postma et al., 2016). Readers are referred to recent reviews for detailed discussion on the endocytic regulation of plant RKs and RPs (Ben Khaled et al., 2015; Frescatada-Rosa et al., 2015).

Pathogen infection often induces the rearrangement of host plasma membrane constituents and the formation of a membrane microdomain, a compartment distinct from the surrounding membrane in composition, structure and biological function (Malinsky et al., 2013). Ligand-induced receptor endocytosis leads to plasma membrane compartmentation or microdomain formation. For instance, flg22 induces FLS2 internalization, which forms distinct vesicles underneath the plasma membrane (Robatzek et al., 2006). Consistent with this observation, flg22 induces rapid and profound changes of protein composition in detergent-resistant membranes (DRM), i.e., the non-solubilized fraction of membranes, in which FLS2 is enriched (Keinath et al., 2010). By contrast, FLS2 is depleted in the soluble fraction of membranes, indicting a re-localization of FLS2 (Keinath et al., 2010). Ligand-induced accumulation of PRRs in defined microdomains may play an important role in receptor regulation.

In addition to PRRs, BIK1 protein levels are also finely-regulated by protein turnover. Arabidopsis CPK28 constitutively associates with and promotes proteasome-dependent turnover of BIK1.
The \textit{cpk28} mutant accumulates higher BIK1 protein levels and displays enhanced PTI responses, whereas overexpression of \textit{CPK28} leads to reduced BIK1 protein levels and decreased responses to PAMPs (Monaghan et al., 2014).

Plant heterotrimeric G proteins play important roles in PRR-mediated immunity. Arabidopsis GUANINE NUCLEOTIDE-BINDING PROTEIN SUBUNIT-\(\beta\) (AGB1), NUCLEOTIDE-BINDING PROTEIN SUBUNIT-\(\gamma\) and AGG2 are required for FLS2-, EFR-, and CERK1-mediated immunity as well as cell death and defenses triggered by the \textit{bir1} mutation (Liu et al., 2013b; Lorek et al., 2013; Torres et al., 2013). Recently, it was shown that a heterotrimeric G protein complex, consisting of EXTRALARGE GUANINE NUCLEOTIDE-BINDING PROTEIN 2 (XLG2), AGB1 and AGG1 or AGG2, associates with the FLS2 complex to stabilize BIK1 and promotes immune signaling (Zhu et al., 2009; Maruta et al., 2015; Liang et al., 2016). It remains unknown which ubiquitin E3 ligase is responsible for BIK1 turnover and whether this E3 is subject to regulation by CPK28 and the heterotrimeric G protein complex.

**REGULATION OF DOWNSTREAM SIGNALING BY PRRs**

A number of RLCKs have emerged as central components linking PRRs to downstream defenses (Figure 2; Lin et al., 2013a). Among these, BIK1 is a central component integrating signals from multiple PRRs by directly interacting with FLS2, EFR, PEPRs, and CERK1 (Lu et al., 2010; Zhang et al., 2010; Liu et al., 2013a). Additional Arabidopsis RLCK VII members, including PBS1-LIKE 1 (PBL1), PBL27, PCRKs, and one RLCK XII member, BSK1, have been shown to play a role in pattern-triggered immunity by directly interacting with PRRs (Zhang et al., 2010; Liu et al., 2013a; Shi et al., 2013; Shinya et al., 2014; Sreekanta et al., 2015; Shinya et al., 2016). The rice RLCK VII members RLCK185 and RLCK176 have also been reported to mediate PGN- and chitin-induced defenses by interacting with CERK1 (Yamaguchi et al., 2013; Ao et al., 2014). Likewise, rice RLCK107 and RLCK118 were reported to interact with XA21, and silencing of these genes compromised XA21-dependent resistance (Hu et al., 2016).

Within minutes, pattern recognition triggers a number of cellular events, including increase of cytoplasmic calcium ([Ca\(^{2+}\)\textsubscript{cyt}]), cation and anion effluxes, extracellular alkalization, production of reactive oxygen species (ROS), and activation of MAP kinase (MPK) cascades (Boller and Felix, 2009). Pharmacological experiments indicated that calcium influx from the apoplast and calcium release from
internal stores are required for the pattern-triggered \([\text{Ca}^{2+}]_{\text{cyt}}\) increase (Klüsener et al., 2002; Thor and Peiter, 2014). Calcium influx is required for ROS burst, cation and anion effluxes, MPK activation, and downstream defense gene expression, suggesting that elevation of \([\text{Ca}^{2+}]_{\text{cyt}}\) is central for pattern-triggered immunity (Jeworutzki et al., 2010; Segonzac et al., 2011; Ranf et al., 2011). BIK1 and PBL1 are required for flg22-triggered \([\text{Ca}^{2+}]_{\text{cyt}}\) increase, suggesting that BIK1 and PBL1 directly or indirectly regulate a calcium channel (Figure 3). Identification of the pattern-triggered calcium channel remains a challenging task, but several plasma membrane-localized calcium permeable channels such as GLUTAMATE-LIKE RECEPTORS (GLRs), CYCLIC NUCLEOTIDE GATED CHANNELS (CNGCs), and REDUCED HYPEROSMOLARITY-INDUCED \([\text{Ca}^{2+}]\) INCREASEs (OSCA) exist in plants (Dodd et al., 2010; Hou et al., 2014; Yuan et al., 2014). For instance, GLRs are required for \([\text{Ca}^{2+}]_{\text{cyt}}\) increase in the pollen tube during self-incompatibility (Iwano et al., 2015). OSCA1 is required for \([\text{Ca}^{2+}]_{\text{cyt}}\) increase during osmotic stress (Hou et al., 2014; Yuan et al., 2014). CNGC2 has been shown to be a calcium channel for LPS- and pep-induced calcium influx (Ali et al., 2007, Qi et al., 2010), whereas GLRs have been implicated in calcium burst following flg22, elf18 and chitin treatment (Kwaaitaal et al., 2011). A more recent study, however, failed to lend support for GLRs and CNGCs in pattern-induced calcium burst (Thor and Peiter, 2014). During symbiosis, calcium oscillation in the nucleus is controlled by nuclear membrane-localized CNGCs and \(\text{Ca}^{2+}\)-dependent adenosine triphosphatase (\(\text{Ca}^{2+}\)-ATPase) (Capoen, et al., 2011; Charpentier, M., et al., 2016). Flg22 treatment also leads to calcium oscillation in the guard cell (Thor and Peiter, 2014), suggesting that pattern-induced calcium burst may be similarly controlled by both calcium channels and calcium pumps. Indeed, Arabidopsis \(\text{Ca}^{2+}\)-ATPase 8 (ACA8) and ACA10 are directly associated with FLS2 and are required for optimum defenses (Frei dit Frey et al., 2012). Future challenges will be to identify the calcium channels involved, to decipher their regulation by BIK1 and PBL1, and to determine how they act together with \(\text{Ca}^{2+}\)-ATPases to regulate cytosolic calcium oscillation.

In Arabidopsis, the NADPH oxidase RESPIRATORY BURST OXIDASE HOMOLOG D (RbohD) is essential for pattern-triggered ROS production (Zhang et al., 2007). Pattern-triggered ROS production is required for stomatal closure and callose deposition (Zhang et al., 2007; Mersmann et al., 2010; Macho et al., 2012). BIK1, PBL1, BSK1, and RLCK185 have been shown to positively regulate pattern-triggered ROS production in Arabidopsis and rice (Figure 3; Zhang et al., 2010; Shi et al., 2013; Yamaguchi et al., 2013). RbohD constitutively associates with BIK1 and FLS2 in plants (Kadota et al.,
2014; Li et al., 2014). Upon activation by flg22, BIK1 and PBL1 phosphorylate RbohD at multiple sites required for ROS production. This phosphorylation, however, is not sufficient for the activation of the NADPH oxidase activity of RbohD, indicating that additional regulation is required. Indeed, CALCIUM-DEPENDENT PROTEIN KINASEs (CPKs), such as CPK5, are necessary to phosphorylate additional sites in RbohD required for activation (Dubiella et al., 2013). Furthermore, RbohD contains EF-hand motifs for calcium ion binding, which likely provide further regulation. XLG2 and PBL13 may also positively or negatively regulate RbohD activity, respectively, although the underlying mechanisms are unknown (Figure 3; Liang et al., 2016; Lin et al., 2015).

It is not known how pattern recognition leads to extracellular alkalinization, or what the biological function of extracellular alkalinization is. FERONIA (FER), a Lectin-RK, interacts with a family of Arabidopsis peptides termed RAPID ALKALINIZATION FACTORs (RALFs) to trigger extracellular alkalinization in plants and plays multiple roles in the regulation of plant growth and development (Pearce et al., 2001; Haruta et al., 2014; Tavorrmina et al., 2015). The activation of FER by RALF results in the phosphorylation of Arabidopsis plasma membrane H⁺ ATPase 2 (AHA2) at Ser899, which inhibits the H⁺ ATPase activity and increases extracellular alkalinization. Interestingly, flg22 treatment also induces AHA2 Ser899 phosphorylation, suggesting that the phosphorylation-regulated H⁺ ATPase activity may be responsible for flg22-induced extracellular alkalinization (Figure 3; Filex et al., 1999; Benschop et al., 2007). Although it is unknown whether AHA2 or other AHAs are required for pattern-triggered immunity, constitutive activation of AHA1 prevents stomatal closure in response to flg22 treatment (Liu et al., 2011). The Arabidopsis RPM1-INTERACTING 4 (RIN4) protein originally identified as a protein guarded by the NLR protein RESISTANCE TO PSEUDOMONAS SYRINGAE PV. MACULICOLA1 (RPM1), has been shown to interact with and promote the activity of AHA1 to induce stomatal opening, which favors bacterial pathogen invasion. Interestingly, induced phosphorylation of RIN4 at Thr-21, Ser-160, and Thr-166 by P. syringae effector AvrB enhances AHA activity and promotes stomatal opening (Lee et al., 2015), whereas flg22 treatment specifically induces RIN4 phosphorylation at Ser141 to enhance FLS2-mediated immunity (Chung et al., 2014), although it remains unknown whether this is associated with a regulation of AHA activity.

Pattern-triggered immunity activates two MPK cascades that play crucial roles in regulating downstream defense gene expression and phytoalexin biosynthesis (Meng and Zhang, 2013). One cascade is composed of MAP Kinase Kinase Kinase 1 (MEKK1), MAP Kinase Kinase 1 (M KK1) and
MKK2, and MAP Kinase 4 (MPK4). The second MAP kinase cascade is composed of an unknown MAPKKK, MKK4 and MKK5, and MPK3 and MPK6. Increasing evidence supports that PBLs mediate pattern-triggered MPK activation (Figure 3). The X. campestris campestris effector protein AvrAC, a specific inhibitor of BIK1 and related PBL kinases, strongly inhibits MPK3, MPK4, and MPK6 activity in flg22-treated seedlings, suggesting that PBLs collectively are required for MPK activation (Feng et al., 2012). The bik1 phl1 double mutant is normal in flg22-triggered but impaired in Pep-triggered MPK activation (Feng et al., 2012; Yamada, et al., 2016a). A recent report showed that perk1 perk2 plants are slightly reduced in flg22-induced MPK activation (Kong et al., 2016). PBL27 was shown to interact with CERK1, and the pbl27 mutant was reported to exhibit impaired MPK activation in response to chitin, but elevated MPK activation in response to flg22 (Shinya et al., 2014). These results suggest differential contribution of RLCK family members to MPK activation by different patterns. Most recently, MAPKKK5 was reported to be a specific substrate of PBL27 and mediate chitin-triggered MPK activation (Yamada, et al., 2016b). Paradoxically, MAPKKK5 appears to negatively impact flg22-triggered MPK activation. MAPKKK5 activation was shown to activate MKK4 and MKK5, which is consistent with the activation of MPK3 and MPK6 (Yamada, et al., 2016b). The finding that PBL27 directly links a PRR to MPK activation is potentially exciting. It is not clear, however, how MAPKKK5 activates MPK4, which is known to be activated by MEKK1, MKK1 and MKK2. Another MAPKKK, MAPKKK7, has been shown to interact with FLS2 to negatively regulate flg22-triggered MPK activation, but the underlying mechanism remains unknown (Mithoe et al., 2016). Moreover, a bacterial protease called PrpL was shown to trigger MPK activation potentially through an unknown PRR (Cheng et al., 2015). PrpL-triggered MPK activation requires heterotrimeric G proteins AGB1, GPA1 and AGG1 and AGG2, although heterotrimeric G proteins are not necessary for MPK activation by known PRRs. One possibility is that different PRRs activate MPKs through distinct mechanisms. It will be important to determine whether additional MAPKKKs are responsible for MPK3 and MPK6 activation by other PRRs. More importantly, the aforementioned RLCK mutants are only partially affected in MPK activation in response to various patterns, suggesting that multiple RLCKs have redundant functions. Higher order mutants for RLCKs and MAPKKKs must be carefully analyzed to test whether different RLCKs and MAPKKKs are recruited by different PRRs for MPK activation.

Salicylate (SA) is a key defense hormone whose biosynthesis is induced in plants upon pathogen infection. SA accumulation is required for resistance at the primary inoculation site (local resistance)
and elevated resistance at distal, un-inoculated sites, a phenomenon called systemic acquired resistance (SAR) (Fu and Dong, 2013). Pattern-triggered immunity has also been reported to contribute to SAR (Mishina and Zeier, 2007). Application of flg22 and LPS to local leaves induces salicylate accumulation, expression of SAR marker genes \textit{PATHOGENESIS RELATED 1 (PRI)} and \textit{FLAVIN-DEPENDENT MONOOXYGENASE 1 (FMO1)}, and disease resistance in systemic leaves (Mishina and Zeier, 2007). The pattern-induced systemic defenses require known SAR components such as SID2, PAD4, and NPR1, indicating that pattern-triggered immunity is integrated into the existing SAR signaling pathway. In addition, PEPRs are required for \textit{P. syringae}- and pattern-induced \textit{PRI} expression and disease resistance in systemic leaves, indicating that PEPRs mediate pattern-induced SAR (Ross et al., 2014). \textit{PROPEP2} and \textit{PROPEP3} transcripts are induced only in the primary leaves, and local application of Peps can induce SAR, suggesting that the PEPR pathway contributes to the onset of SAR at primary challenge sites. Recently, PCRK1 and PCRK2 were shown to act redundantly to mediate flg22-induced SA accumulation and disease resistance to pathogens (Figure 3; Kong et al., 2016), although it is not known whether the PEPRs are required for PCRK-mediated SA induction. Future work is needed to test whether PEPRs and PCRKs act in the same pathway to regulate SA signaling and SAR.

As discussed above, RLCKs collectively activate multiple signaling pathways downstream of PRRs (Figure 3). Often, loss of function of a RLCK only leads to partial impairment in downstream signaling, suggesting functional redundancy. Different PRRs can recruit both specific and common RLCKs. For example, PEPRs specifically require BIK1 and PBL1, but not other RLCKs (Liu et al., 2013a). Likewise, PBL27 was reported to be specifically required for defenses triggered by chitin, but not flg22. On the other hand, BIK1 and PBL1 are required for defenses mediated by FLS2, EFR, PEPRs, and CERK1. In addition, FLS2 has been shown to interact with multiple RLCKs including BIK1, PBL1, PCRKs, and BSK1. Furthermore, different RLCKs may be required for the regulation of different downstream signaling components (Figure 3). Thus BIK1 and PBL1 are required for flg22-, elf18-, and chitin-triggered \([Ca^{2+}]_{cyt}\) increase and ROS burst, but not MPK activation. PBL27 and PCRKs are required for chitin-triggered MPK activation and flg22-triggered SA accumulation, respectively, but appear to play a minimal role, if any, in ROS production in response to the same pattern. However, this assumption may be overly simplified, as BIK1 and PBL1 have been shown to contribute to Pep-triggered MPK activation (Yamada et al., 2016a), whereas PCRKs have been shown to contribute to flg22-triggered MPK activation (Kong et al., 2016). Given that RLCK VII is a large family containing
more than 40 members, many of which may play a role in pattern-triggered defenses, it will be important to empirically test both redundancy and specificity of these RLCKs in different PRR complexes and their role in different downstream signaling pathways.

**CROSS-TALK BETWEEN RLK-MEDIATED IMMUNE SIGNALING AND GROWTH/DEVELOPMENT**

Plants must prioritize immunity versus growth and development according to the presence or absence of pathogen-imposed danger (Belkhadir et al., 2014; Lozano-Duran and Zipfel, 2015). BRI1, an Arabidopsis LRR-RK that perceives the steroid hormones brassinosteroids (BRs), is essential for diverse growth and development processes (Li and Chory, 1997). BRs inhibit PTI responses, and activation of BR signaling increases susceptibility to bacterial pathogens (Lozano-Duran and Zipfel, 2015). The BR signaling pathway shares multiple components with FLS2 pathways. Although BAK1 is a co-receptor for both BRI1 and FLS2, it is not a limiting factor to modulate BR and PAMP signaling (Lozano-Duran et al., 2013). BSK1 associates with both BRI1 and FLS2 and plays a positive role in PTI and BR signaling, but does not appear to function in the cross-talk between BR and PTI signaling (Tang et al., 2008; Shi et al., 2013). By contrast, BIK1, a positive regulator in the FLS2 pathway, interacts with and phosphorylates BRI1 to negatively regulate BR signaling (Lin et al., 2013b). The transcription factors BRASSINAZOLE RESISTANT 1 (BZR1) and HBI1 are also shared components of the BRI1 and FLS2 pathways. BZR1 and HBI1 positively regulate BR signaling but suppress immunity upon BR treatment (Bai et al., 2012; Lozano-Duran et al., 2013; Fan et al., 2014; Malinovsky et al., 2014). A more recent study suggests that the transcriptional cascade formed by BZR1 and HBI1 integrates BR and immunity cross-talk with multiple environmental inputs to fine tune the trade-off between growth and immunity (Lozano-Duran and Zipfel, 2015).

The second example of cross-talk between RLK pathways controlling immune signaling and growth/development is exemplified by the study of FER, which plays multiple roles in the regulation of plant growth and development (Tavormina et al., 2015). Early studies showed that FER-mediated signaling can negatively impact disease resistance and pattern-triggered immune responses (Kessler et al., 2010) and that FER becomes phosphorylated following flg22 treatment (Benschop et al., 2007). The recent finding that RALFs negatively regulate PRR complex formation (Stegmann et al., 2017) provides a mechanism for such cross-talk. A previous study suggested that FLS2 and BAK1 exist in a
preformed complex prior to flg22 treatment (Sun et al., 2013). A recent study identified FER as a modulator of flg22, elf18, and chitin signaling, as a loss-of-function fer mutant displays reduced PAMP responses and increased susceptibility to P. syringae (Stegmann et al., 2017). FER is also enriched in microdomains after flg22-treatment (Keinath et al., 2010) and weakly interacts with both FLS2 and BAK1 (Stegmann et al., 2017). The flg22-induced FLS2-BAK1 and elf18-induced EFR-BAK1 interactions are compromised in fer seedlings, indicating that FER promotes ligand-induced dimerization of the receptors and co-receptor (Stegmann et al., 2017). Furthermore, co-treatment of seedlings with RALF23 peptide or overexpression of RALF23 decreases PAMP-induced responses and FLS2-BAK1 and EFR-BAK1 interactions (Stegmann et al., 2017). Together, the study supports that FER can act as a scaffold protein mediating PAMP-induced PRR complex formation and that RALFs actively regulate this process. Similar to the FLS2-FER interaction, the Arabidopsis Malectin-like/LRR-RLK IMPAIRED OOMYCETE SUSCEPTIBILITY1 (IOS1) was recently shown to interact with FLS2, EFR and CERK1, and contributes to pattern-triggered immunity (Yeh et al., 2016). Furthermore, a cysteine-rich RLK, CPK28, was recently shown to interact with FLS2 and positively regulate disease resistance and cell death (Yadeta et al., 2017). A direct interaction between different receptor complexes may facilitate cross-talk between different pathways. It will be extremely interesting to test whether this is a common feature for RKs and RPs.

Another recent example of such cross-talk comes from the study of PHYTOSULFOKINEs (PSKs). PSKs and PLANT PEPTIDE CONTAINING SULFATED TYROSINE 1 (PSY1) are related tyrosine-sulfated peptide hormones that promote root growth and xylem trachea development in plants (Matsubayashi and Sakagami, 2006; Amano et al., 2007). These peptides are perceived by the related LRR-RKs PSK RECEPTORs (PSKRs) and PSY1 RECEPTOR (PSY1R), respectively (Matsubayashi et al., 2006; Amano et al., 2007). Similar to fer mutants, Arabidopsis mutants lacking PSKR1 or TYROSINE PROTEIN SULFOTRANSFERASE (TPSP), the latter of which is required for sulfation of PSKs, exhibit enhanced defense gene expression in response to elf18 and flg22 and increased disease resistance to P. syringae, indicating a negative role of PSKR1 in pattern-triggered immunity (Igarashi et al., 2012). Similarly, PSKR1 and PSY1R play an additive role in the negative regulation of disease resistance to the fungal pathogen Alternaria brassicicola in addition to P. syringae (Mosher et al., 2013). The pskr1 pskr2 psy1r triple mutant is strongly enhanced in flg22-induced defense gene expression. Conversely overexpression of PSKs and PSKR1 increases disease susceptibility to the bacterial and
fungal pathogens. These findings strongly suggest that PSKR1- and PSY1R-mediated signaling inhibits pattern-triggered immunity.

Several LRR-RLKs, including ERECTA (ER), ER-like 1 (ERL1) and ERL2, and LRR-RLP TOO MANY MOUTH (TMM), regulate stomatal patterning in response to epidermal pattern factor peptides (EPFs). Recent findings indicate that these proteins have prominent roles in disease resistance. Overexpression of ER lacking the kinase domain dominantly inhibits ER family receptor function, resulting in increased susceptibility to the fungal pathogen *Plectosphaerella cucumerina* (Jordá et al., 2016). The *er erl1 erl2* triple mutant and *er erl1 erl2 tmm* quadruple mutant also display increased susceptibility to *P.  cucumerina*, indicating that ER, ERL1, ERL2, and TMM contribute to disease resistance. *er* mutant plants are diminished in defense gene expression in response to the fungal spore extract, suggesting that ER positively regulates plant immunity in response to an unknown PAMP. Overexpression of EPFs fails to affect disease resistance to *P. cucumerina*, suggesting that ER, ERL1, ERL2, and TMM regulate disease resistance independently of EPF signaling. Future studies are needed to determine whether these RLKs and RLP regulate plant immunity by cross-talk with unknown PRRs that perceive fungal PAMPs or DAMPs.

**MANIPULATION OF RLP AND RLK SIGNALING BY PATHOGEN EFFECTORS**

While PRRs are powerful in sensing and triggering effective defenses against potential pathogenic microbes, adapted pathogens have evolved a variety of effectors to evade detection or subvert the pattern-triggered immunity (Figure 4). Many effectors impede PRR functions by targeting microbial patterns, PRRs, or early signaling components of pattern-triggered immunity (see Deslanders and Rivas, 2012; Feng and Zhou, 2012; Dou and Zhou, 2012; Macho and Zipfel, 2015 and references therein). Recently, *P. syringae* was found to deliver a novel protease called HopB1 to destroy BAK1 (Li et al., 2016c). Perturbation of BAK1 is known to trigger the plant surveillance system and elevates defenses (Yamada et al., 2016b). Surprisingly, HopB1 goes undetected during infection (Li et al., 2016c). Close examination of the biochemical mode of action showed that HopB1 does not cleave normal BAK1. Instead, it only attacks an flg22-induced, phosphorylated form of BAK1. This highly selective action minimized perturbation to the host, allowing evasion of the host surveillance system. The analyses of these effectors not only elucidated mechanisms of pathogenesis of various pathogens, but also advanced our understanding of key signaling mechanisms mediated by PRRs. For instance, the
search for virulence targets of the *P. syringae* effector AvrPphB led to the identification of BIK1 and PBLs as key signaling components downstream of multiple PRRs (Zhang et al., 2010). The investigation of targets of the *Xanthomonas campestris* effector AvrAC not only re-enforced the importance of BIK1 in pattern-triggered immunity (Feng et al., 2012), but also led to the identification of PBL2 as a decoy for NLR-mediated recognition of AvrAC (Wang et al., 2015). The finding that the *P. syringae* effector HopAO1 acts as a tyrosine phosphatase targeting EFR and FLS2 helped uncover a key role of Tyr836 phosphorylation in EFR immune activation (Macho et al., 2014).

In addition to pathways mediated by PRRs, increasing evidence shows that pathogens also actively manipulate RKs that regulate plant growth and development to increase plant susceptibility (Figure 4). It has long been known that cyst nematodes secrete peptides that are structural and functional mimics of plant CLAVATA3/ENDOSPERM SURROUNDING REGION-related (CLE) peptides (Wang et al., 2001, 2005; Gao et al., 2003; Lu et al., 2009), with the latter being peptide hormones required for meristem maintenance and vascular differentiation (Fiers et al., 2007). The nematode CLE effectors are required for virulence (Patel et al., 2008), apparently by stimulating both the CLV2 and the CLV1 pathways (Replogle et al., 2011, 2013). Similarly, phytopathogenic root-knot nematodes encode peptides similar to plant C-TERMINAL ENCODED PEPTIDEs (CEPs) that are involved in the regulation of cell expansion (Bobay et al., 2013; Tavormina et al., 2015), although their roles in pathogenesis remain unknown. An outstanding question remains how CLE-mediated signaling benefits nematode pathogenesis. It is possible that CLE effectors may reprogram root development for infection and/or feeding. It is equally possible that CLE signaling is engaged in cross-talk with immunity. Indeed, it has been shown that the Arabidopsis coryne (crn) mutant displays an enhanced defense response gene expression in roots (Miwa et al., 2008). Future studies are needed to elucidate the underlying mechanisms for CLE effector-mediated nematode virulence. The CLV1 and CLV2 pathways may be targeted by additional pathogens, as clv1 and clv2 plants are more resistant to *Ralstonia solanacearum* (Hanemian et al., 2016), although the underlying mechanism remains unknown.

A search of fungal genomes identified numerous RALF-like peptides (Masachis et al., 2016; Thynne et al., 2016). Strikingly, all fungi encoding RALF-like sequences are plant pathogens. Among these, a *Fusarium oxysporum* RALF-like peptide (F-RALF) has been shown to stimulate extracellular
alkalinization in plants. A loss-of-function fer mutant exhibits increased resistance to *F. oxysporum* (Masachis et al., 2016). Of note, hemitrophic and biotrophic fungal pathogens are known to cause extracellular alkalinization in plants, an important feature of their life styles (Prusky et al., 2001; Prusky and Yakoby, 2003). *F. oxysporum* mutants lacking F-RALF are unable to induce extracellular alkalinization and are less virulent in plants (Masachis et al., 2016). The F-RALF-induced pH elevation triggers rapid phosphorylation of the fungal MPK Fmk1 essential for invasive hyphal growth (Di Pietro et al., 2001; Masachis et al., 2016). The fungal pathogen may have taken advantage of the cross-talk between FER-mediated signaling and pattern-triggered immunity to promote pathogenesis (Figure 4). Indeed, *F. oxysporum* F-RALF mutants are less capable of inhibiting pattern-induced defense gene expression (Masachis et al., 2016).

The observation that PSKR1- and PSY1R-mediated signaling negatively impacts pattern-triggered immunity suggests that pathogens could target these proteins for virulence. Indeed, the *X. oryzae oryzae* peptide RaxX-sY has high levels of amino acid sequence similarity to PSY1 from plants (Pruit et al., 2015), raising the tantalizing possibility that RaxX evolved to mimic PSY1, thereby suppressing immunity (Figure 4). It will be interesting to determine whether RaxX-sY indeed mimics PSY1 in the regulation of plant growth and development and plant defenses. If so, an outstanding question would be how Xa21 differentiates endogenous PSY1 peptide from RaxX-sY so that defenses do not get activated by the endogenous peptide hormone.

Filamentous pathogens secrete hundreds of effectors, many of which are targeted to the apoplast. It is tempting to speculate that additional effectors inhibit PRRs or stimulate RKs that control growth and development. Indeed, the *C. fulvum* effector Ave1 is a virulence factor and shares sequence homology with proteins from diverse plant species, suggesting a plant origin of Ave1 (de Jonge et al., 2012). Although the function of these plant proteins is not known, the findings raise the possibility that Ave1 enhances virulence by mimicking these plant proteins. As peptide sequences are often too short for simple BLAST analysis, advanced computational methods will likely uncover additional pathogen effectors that mimic plant peptides.

**CONCLUDING REMARKS**

RLKs and RLPs are at the core of the plant early warning system in the wake of pathogen attacks. Biochemical and structural studies are rapidly advancing our understanding of the mechanisms of PRR
action in immunity. Ligand-induced recruitment of co-receptors is crucial for LRR- and LysM-containing PRRs. Future studies will test whether this mode of action can be extrapolated to PRRs containing other classes of ECDs. In addition to receptors and co-receptors, PRR complex components are also associated with RLKs, RLCKs, heterotrimeric G proteins, PUBs and PP2Cs. These proteins allow dynamic and tight regulation of PRR complexes before and after pattern recognition. RLCKs associated with the RK type of PRRs are central for the activation of bifurcating downstream signaling pathways, while it remains to be determined whether this is also true for the RP type of PRRs. Understanding how RLCKs activate downstream signaling components will continue to yield exciting mechanistic insights for plant immunity.

The number of microbial patterns recognized by different plant species is likely to be enormous. Our understanding of host recognition of microbial patterns is limited by our ability to identify microbial patterns and their corresponding receptors. Nonetheless, proteinaceous microbial patterns are subject to positive selection at individual residues as a result of host–pathogen co-evolution. With the help of computational tools it is now possible to exploit microbial genome sequences to expedite the discovery of microbial patterns. Likewise, PRRs often belong to the RLK and RLP subfamilies that display lineage-specific expansion. This feature not only helps the identification of PRRs, but also allows deployment of disease resistance by cross-species/genus transfer of PRR-coding genes. Indeed, transgenic overexpression of EFR confers elf18 sensitivity and elevated resistance to cereal bacterial pathogens in rice and Triticum aestivum (wheat) plants, which do not encode EFR (Schwessinger et al., 2015; Schoonbeek et al., 2015).

Extensive studies in the past decade have demonstrated that microbial effectors enhance virulence often by directly interfering with PRR-mediated signaling. Emerging evidence indicates that pathogen effectors can also promote virulence by mimicking plant peptide hormones to indirectly interfere with immune signaling through cross-talk between different signaling pathways. The interactions between microbial effectors and signaling pathways mediated by plant RKs will continue to be a fertile ground of investigation.

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AUTHOR CONTRIBUTIONS

All authors contributed to writing the article.
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</table>

Known PRRs refer to genetically confirmed receptors whose binding to patterns has been biochemically demonstrated, whereas probable PRRs refer to likely receptors with genetically confirmed roles in pattern recognition, but for which direct binding to patterns remains to be shown.
Table 2. Other RLKs and RLPs involved in plant immunity*.

<table>
<thead>
<tr>
<th>Name</th>
<th>Family</th>
<th>Function</th>
<th>Plant</th>
<th>References</th>
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<td>Gao et al., 2009</td>
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<td>SOBIR1</td>
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<td>FER</td>
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<td>RALF receptor, Scaffold for PRR</td>
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<td>CRK28</td>
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</table>

*Includes RLKs/RLPs that regulate plant immunity through an unknown molecular function or function other than PRRs.
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Figure 1. Ligand-induced hetero-dimerization between LRR-RKs and SERKs. Structural alignment of FLS2-flg22-BAK1, PEPR1-Pep1, PXY-TDIF-SERK2, and HAESA-IDA-SERK1 complexes. The structure of FLS2 ECD was used as the template for the alignment. Note only ECDs were included in the structural studies. N and C denote N-termini and C-termin of LRR-RK ECDs and peptides.
Figure 2. RLCKs differentially mediate immune signaling from different PRRs.
Different RLCKs known to interact with PRRs are colored in blue. BIK1 and PBL1 directly interact with multiple PRRs or co-receptor, including LRR-RKs (FLS2, EFR, and PEPRs), LRR-RLK (BAK1) and LysM-RLK (CERK1), to mediate immune signaling. The RLCK PBL27 was reported to specifically associate with CERK1 to mediate chitin-triggered immune signaling. In addition, RLCKs BSK1 and PCRKs are also capable of interacting with FLS2 to mediate flagellin-triggered immune signaling. The rice RLCK185 directly interacts with CERK1 to mediate chitin-triggered signaling. FLS2, EFR, and PEPRs perceive bacteria flagellin, EF-Tu, and endogenous DAPMP Peps, respectively, and recruit co-receptors BAK1/SERKs to form active receptor complexes, whereas the rice LysM-RLP CEBiP and Arabidopsis LysM-RK LYK5 bind chitin and recruit LysM-RLK CERK1 to form active receptor complexes. The activated receptor complexes phosphorylate RLCKs to activate downstream signaling.
Figure 3. RLCKs differentially regulate downstream signaling.
RLCKs including BIK1, PBL1, RIPK, PCRKs, BSK1, and PBL27 are colored in blue. BIK1 and PBL1 positively regulate multiple downstream responses including RbohD-dependent ROS burst, \([Ca^{2+}]_{cyt}\) increase, and activation of heterotrimeric G proteins. BIK1 and RIPK are required for phosphorylating distinct residues in RIN4, a plasma membrane proton ATPase-interacting protein, to positively or negatively regulate immunity. Proton efflux further regulates other ion channels, such as K⁺ channel. BSK1 positively regulate ROS burst downstream of FLS2. PCRKs positively regulate SA biosynthesis upon flg22 treatment. PBL27 positively regulates MAPKKK5 to specifically mediate chitin-triggered MPK activation. The activation of RbohD also requires \([Ca^{2+}]_{cyt}\) increase and CPK5. Solid lines indicate direct phosphorylations or protein-protein interactions, whereas dashed lines indicate that direct interactions/phosphorylations are unknown.
Figure 4. Pathogen effector proteins enhance by interfering PRR-mediated signaling or mimicking peptide hormones. Different effectors and molecular patterns are colored in purple. The fungal apoplastic effectors ECP6 and SLP1 compete with plant receptors for chitin-binding. Bacterial cytoplasmic effectors can block PRR-mediated signaling by physically inhibiting (AvrPto and AvrPtoB) or dephosphorylating (HopAO1) PRR kinases, proteolysis of co-receptors (HopB1), proteolysis (AvrPphB) or uridylation (AvrAC) of RLCKs, ADP-ribosylation of MAPKK or BAK1 (HopF2), or dephosphorylation of MPKs (HopAI1). AvrB induces phosphorylation of RIN4 to stimulate plasma membrane proton ATPase activity. Fungal apoplastic effector F-RALF mimics plant RALF peptide to induce FER-mediated signaling pathway, which inhibits PRR-specified immunity through cross-talk. The bacterial protein RaxX may mimic plant peptide PSY1 to stimulate PSY1R-mediated signaling, which is known to inhibit PRR-mediated immunity.
Receptor kinases in plant pathogen interactions: more than pattern recognition
Jian-Min Zhou, Dingzhong Tang and Guoxun Wang
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