Molecular insights into the biochemical functions and signalling mechanisms of plant NLRs

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1 | INTRODUCTION

Plants are constantly exposed to a wide variety of microbial pathogens, including bacteria, fungi, oomycetes, and viruses. To resist infections, plants have evolved an active two-layered immune system. The first layer involves recognition of pathogen-associated molecular patterns (PAMPs) by plant cell surface pattern recognition receptors (PRRs), known as PAMP-triggered immunity (PTI). Successful pathogens secrete effectors into plant cells to suppress the PTI response and cause infections. Plants respond to pathogen effectors specifically with cytoplasmic NLR (nucleotide-binding leucine-rich repeat) protein receptors and activate effector-triggered immunity (ETI), which constitutes the second layer of the plant immune system. The hallmark of ETI is localized plant cell death at infection sites, also known as the hypersensitive response, to restrict pathogen spread (Jones et al., 2016).

Knowledge of activation and signalling mechanisms of PRR and NLR receptors has greatly advanced (DeFalco & Zipfel, 2021; Saur et al., 2021). Increasing evidence suggests that the two layers of immunity are more intimately connected than previously thought (Ngou et al., 2021; Yuan et al., 2021). Research on the interdependency between PRR- and NLR-mediated immunity is rapidly developing (Pruitt et al., 2021; Tian et al., 2021). NLR protein signalling mechanisms are also under intensive investigation after recent breakthrough studies revealing the unexpected biochemical functions of NLR proteins (Wang et al., 2020). Structural biology, specifically through advancements in cryoelectron microscope (cryo-EM) techniques, has significantly contributed to our understanding of NLR protein biology. This review will focus on current knowledge and future perspectives on plant NLR protein activation and signalling.

2 | NLR DOMAIN STRUCTURE AND SUBFAMILIES

In addition to the central nucleotide-binding (NB) domain and the C-terminal leucine-rich-repeat (LRR) domain, plant NLR proteins have a variable N-terminal domain that can be a Toll/interleukin-1 receptor (TIR) domain or a coiled-coil (CC) domain or an RPW8 (Resistance to Powdery Mildew 8)-like CC (CC8) domain. NLR proteins with...
N-terminal TIR, CC, or CC8 domains are known as TNLs, CNLs, or RNLs, respectively (Duxbury et al., 2021). TNLs and CNLs recognize pathogen effectors directly or indirectly to activate cell death and defense responses, thus are regarded as sensor NLRs (Jubic et al., 2019). TNLs or CNLs also function as pairs to perceive pathogen effectors and confer resistance. For instance, in Arabidopsis two TNLs, RPS4 and RRS1, located in a head-to-head orientation in the genome, cooperate to resist infection by at least three different pathogens (Narusaka et al., 2009). In rice, two CNLs, RGA4 (Resistance Gene Analogue 4) and RGA5 (Resistance Gene Analogue 5), function together to resist Magnaporthe oryzae carrying the effector Avr-Pia or Avr1-CO39 (Cesari et al., 2013). In the NLR pair system, one NLR possesses an additional domain called the integrated decoy (ID) domain for direct effector interaction functioning as a sensor, while the other NLR functions as an executor for immune signalling (Le Roux et al., 2015; Williams et al., 2014). Downstream of sensor NLRs or sensor/executor NLR pairs, higher plants also use helper NLRs to transduce signals. There are three described families of helper NLRs: NLR Required for Cell death (NRC), Activated Disease Resistance 1 (ADR1), and N Required Gene 1 (NRG1) (Jubic et al., 2019). In terms of domain structure, NRCs belong to the CNL class, while ADR1s and NRG1s belong to the RNL class. In solanaceous species such as Nicotiana benthamiana, NRCs are required for phylogenetically related CNLs to activate cell death and immune responses (Wu et al., 2017). In N. benthamiana, NRC2, NRC3, and NRC4 function downstream of different CNLs to positively regulate immunity, while NRCX negatively regulates NRC2 and NRC3-mediated immunity (Adachi et al., 2021). In Arabidopsis thaliana, three ADR1s, NRG1.1 and NRG1.2 are full-length RNLs (CC8-NB-LRR) functioning as positive regulators, while the truncated NRG1.3 (NB-LRR) negatively regulates AtNRG1.1- and AtNRG1.2-mediated immunity (Wu et al., 2021). In dicots, ADR1s and NRG1s are required downstream of TIR-only receptors, TNLs and TNL pairs to activate defense responses, and ADR1s are also required for the full function of CNLs (Jubic et al., 2019). ADR1s, but not NRG1s, are also present in monocots with uncharacterized function. Some NLRs that function as both sensors of effectors and helpers directly activating cell death are regarded as NLR singletons (Adachi et al., 2019) (Figure 1).

### 3 | NLR RECOGNITION OF EFFECTORS

Plant sensor NLRs recognize pathogen effectors via direct interaction or perceiving the modifications by effectors (Duxbury et al., 2021). Well-known examples of direct NLR–effector recognition mostly come from TNLs including flax L567, Arabidopsis RPP1 (Recognition of Peronospora parasitica 1), and N. benthamiana ROQ1 (Recognition of XopQ 1). The structural bases of direct effector recognition, ATR1 by RPP1 and XopQ by ROQ1, have been reported (Ma et al., 2020; Martin et al., 2020). Interestingly, a previously undefined C-terminal jelly roll/ Ig-like domain (C-JID) was revealed in the post-LRR domain regions of RPP1 and ROQ1. Both C-JID and LRR domains are involved in effector recognition (Figure 2a,b). The structure-conserved but sequence-diversified C-JID is present in

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**Figure 1** Schematic diagram depicting the domain structures of sensor NLRs and helper NLRs and their interdependence for function. TIR, toll/interleukin-1 receptor; CC, coiled-coil; CC8, (Resistance to Powdery Mildew 8)-like CC; NB, nucleotide-binding; LRR, leucine-rich repeat; ID, integrated decoy.
many other TNLs from dicotyledonous plant species, but not in CNLs from diverse plant species (Ma et al., 2020).

In addition, well-studied NLR pairs use the ID domain of sensor NLRs to physically associate with effectors for activation (Duxbury et al., 2021). In the TNL pair of RPS4 (Resistance to *Pseudomonas syringae*) and RRS1 (Resistance to *Ralstonia solanacearum*), the integrated WRKY domain of RRS1 detects effectors AvrRps4 from *Pseudomonas syringae* pv. *pisi* and PopP2 from *Ralstonia solanacearum* (Le Roux et al., 2015; Narusaka et al., 2009). Interestingly, crystal structures of RRS1 WRKY in complex with AvrRps4 or PopP2 revealed that the two effectors target the same region of RRS1 WRKY (Mukhi et al., 2021, Zhang et al., 2017b) (Figure 2c). In rice, two CNL pairs, RGA4/RGA5 and Pik-1/Pik-2, use the integrated heavy metal-associated (HMA) domain in the sensors RGA5 and Pik-1, respectively, to directly recognize cognate effectors. A few structures of the Pik-1 HMA domain in complex with the effector AvrPik highlighting the recognition specificities and key residues for activation have been reported (Bialas et al., 2021; De la Concepcion et al., 2018, 2019) (Figure 2d).

Well-studied examples of the indirect effector-recognition mode come from CNLs. In *Arabidopsis*, RPM1-interacting protein 4 (RIN4) serves as a guardee physically bound to two CNLs, Resistance to *Pseudomonas syringae* pv. *maculicola* 1 (RPM1). Effector-induced cleavage or phosphorylation of RIN4 activates RPS2 and RPM1, respectively (Duxbury et al., 2021). Another well-demonstrated example of indirect recognition is the *Arabidopsis* CNL ZAR1 (Hopz-Activated Resistance 1) and the *Xanthomonas campestris* effector AvrAC. Two receptor-like cytoplasmic kinases, RKS1 and PBL2, are involved in the recognition. RKS1 constitutively interacts with the LRR domain of ZAR1. AvrAC uridylates PBL2 and the uridylated PBL2 (PBL2UMP) is then recruited to the RKS1–ZAR1 complex to activate ZAR1 (Wang et al., 2015). The structures of the inactive RKS1-ZAR1 complex (ADP bound) and the intermediate PBL2UMP–RKS1-ZAR1 complex (ADP free) revealed effector-triggered allosteric ADP release as a key step to prime NLR for ATP binding and activation (Wang et al., 2019b) (Figure 2e).

4 | N-TERMINAL SIGNALLING DOMAIN: DEFINING THE BIOCHEMICAL FUNCTIONS OF NLRS

The N-terminal TIR, CC, or CC<sup>R</sup> domains of NLRs are the signalling domains, overexpression of which in *N. benthamiana* is sufficient to trigger cell death (Bernoux et al., 2011; Collier et al., 2011; Maekawa et al., 2011; Swiderski et al., 2009). Structural biology studies on plant NLRs were limited to the N-terminal signalling domains before the advancement of cryo-EM as a technique to solve full-length NLR structures.
Crystal structures of a few plant TNL TIR domains have been reported, including Arabidopsis RPS4, RRS1, SNC1 (suppressor of npr1-1, constitutive 1), RPP1, flax L6, and grape RUN1 (Resistance to Uncinula necator 1) (Bernoux et al., 2011; Horsefield et al., 2019; Williams et al., 2014; Zhang et al., 2017a). Plant TIR domains feature a global fold consisting of a five-stranded parallel \( \beta \) sheet \((\beta A-\beta E)\) surrounded by five \( \alpha \)-helical regions \((\alpha A-\alpha E)\). The global fold is similar to animal and bacterial TIR structures. The signalling incompetent TIR domain of RRS1 functioning as a sensor in the RPS4/RRS1 pair system harbours a deletion in the \( \alpha D \) helical region compared to the signalling competent TIR domain of RPS4 functioning as an immune executor (Williams et al., 2014). A well-characterized dimerization interface identified from the crystal contacts is the one mainly involving the \( \alpha A \) and \( \alpha E \) helices with a conserved serine and histidine motif at the core of this AE interface (Williams et al., 2014). Another dimerization interface mainly involving the \( \alpha D \) and \( \alpha E \) helices, termed the DE interface, is positioned in slightly different orientations in different TIR crystal structures and thus is less well defined (Zhang et al., 2017a). TIR self-association via the AE and DE interfaces is required for the cell death function of plant TIR domains. AE and DE interfaces are not mutually exclusively and could potentially mediate the formation of TIR filament structures (Nishimura et al., 2017; Zhang et al., 2017a) (Figure 3a). Animal TIR proteins have been shown to form filament structures in vitro (Ve et al., 2017).

Structures of plant TIRs and their dimeric forms do not implicate the biochemical functions. Well-characterized TIR domains of animal cell-surface toll-like receptors dimerize on activation to recruit downstream signalling proteins (Fitzgerald & Kagan, 2020). Hence scientists thought plant TIRs would also function as scaffold proteins. In 2017 and 2018, some bacterial TIR domains and the TIR domain of animal SARM1 involved in the cell death pathway of neuron degeneration were demonstrated to possess enzymatic activity, cleaving nicotinamide adenine dinucleotide (NAD\(^+\)) (Essuman et al., 2017, 2018). Follow-up investigations confirmed that plant TIRs indeed function as NADase to trigger cell death dependent on the conserved catalytical glutamate and the integrity of the two dimerization interfaces (Horsefield et al., 2019; Wan et al., 2019). Different from the more active SARM1 enzyme that causes NAD\(^+\) depletion and subsequent cell death, plant TIRs domains are weak NADase without causing NAD\(^+\) depletion and potentially produce signalling molecules such as nicotinamide (NAM), adenosine diphosphate ribose (ADPR), and variant-cADPR (an isomer of cyclic ADPR) to bind and activate proteins downstream (Wan et al., 2019).
pore-forming proteins, such as animal Mixed Lineage Kinase Domain-Like (MLKL) protein and fungal HeLo and HeLo-like proteins (Bentham et al., 2018; Daskalov et al., 2016; Jubic et al., 2019; Mahdi et al., 2020). The crystal structure of AtNRG1.1 CC8 confirmed the 4HB fold of CC8 (Jacob et al., 2021) (Figure 4a). AtNRG1.1 CC8 harbours an extra region N-terminal to the 4HB fold compared to canonical CC domains, which is consistent with previous structural predictions of CC8 and RPW8. The extra region N-terminal to the 4HB fold of CC8 and RPW8 is predicted to be responsible for membrane anchoring (Zhong & Cheng, 2016). A recent study showed that deletion of the extra region N-terminal to the 4HB fold impairs the cell death function of auto-active AtNRG1.1 but not its membrane localization (Jacob et al., 2021). It has also been shown that RNLs interact with anionic plasma membrane (PM) phospholipids, and depletion of phosphatidylinositol-4-phosphate from the PM causes mislocalization of RNLs and abolishes their cell death activities (Saile et al., 2021).

MLKL is a well-studied membrane pore-forming protein (Murphy et al., 2013). Current modeling suggests that MLKL remains cytosolic and monomeric in the resting state, but starts to oligomerize, move to the PM, and form pores at the PM, leading to loss of membrane integrity and cell death (Huang et al., 2017). The exact oligomerization state and pore size of activated MLKL are still unknown. It has also been shown that MLKL functions as a cation channel to induce cell death, implicating the possible channel activities of plant CNLs and RNLs (Xia et al., 2016).

7 | NLR RESISTOSOMES AND SIGNALLING

Self-association is known to be required for NLR function. The NB domain is responsible for the self-association and oligomerization of NLRs and functions as a molecular switch bridging immune stimulus perception by the LRR domain and immune signalling by the N-terminal domain (Burdett et al., 2019; Saur et al., 2021).

8 | CNL ZAR1 PENTAMERIC RESISTOSOME FUNCTIONING AS A Ca^{2+} CHANNEL

In 2019, cryo-EM studies of full-length ZAR1 in both resting and active states for the very first time revealed the structure of interdomain interactions for autoinhibition and oligomerization for immune activation of a plant NLR (Wang et al., 2019a, 2019b). The structure of monomeric ADP-bound ZAR1 in complex with RKS1 represents the resting state. RKS1 interacts exclusively with the ZAR1 LRR domain to maintain ZAR1 in the resting state (Wang et al., 2019b). The structure of dATP-bound ZAR1 in complex with RKS1 and effector-produced PBL2UMP represents the active state. On effector recognition, ZAR1 undergoes dramatic conformational changes from an inactive monomeric state to an active pentameric state with the first helix in the 4HB fold of its CC domain being flipped, potentially forming a pore at the PM. In addition to the first helix α1 being flipped, the fourth helix α4A becomes disordered in the active state. Previously identified sequences C-terminal to the 4HB domain that are critical for self-association and cell death function are largely disordered in the resting state but become a helix, termed α4B, in the active state (Figure 4b,c). These conformational changes in the CC domain and pentamerization driven by the NB domain feature the dynam- ics of ZAR1 resistosome formation. The ZAR1 resistosome displays a channel pore of 1 nm in diameter (Wang et al., 2019a) (Figure 4d). A follow-up study confirmed that the effector-activated ZAR1 resistosome protrudes into the PM and forms a calcium-permeable cation channel leading to calcium influx and further activation of cell death and defence responses (Bi et al., 2021) (Figure S1a).

9 | TNL RPP1 AND ROQ1 TETRAMERIC RESISTOSOMES FUNCTIONING AS NADase

Recent cryo-EM studies of TNL/effector pairs ROQ1/XopQ and RPP1/ATR1 structures, followed by structure-guided biochemical studies in vitro and functional analyses in planta, greatly advanced our understanding of TNL activation mechanisms and functioning as a NAD\(^+\) cleavage enzyme (Ma et al., 2020; Martin et al., 2020). On effector recognition, ROQ1 and RPP1 assemble into homotetrameric resistosomes. An apparent difference is that the ROQ1 resistosome binds an ATP molecule in the NB domain as expected, while the RPP1 resistosome binds an ADP molecule although ATP was supplemented in the purification process. Further analyses indicated that plant CNLs and some TNLs, such as ROQ1, use a “TT/SR” motif for ATP binding, while some TNLs, such as RPP1, have a “TTE/Q” motif, losing the ATP binding ability and bind ADP molecules (Ma et al., 2020). Tetramerization in the NB domain drives the formation of the TIR tetramer as an active holoenzyme. The TIR tetramer contains two previously characterized symmetrical AE interfaces and two asymmetric interfaces that create the potential active sites for NAD\(^+\) cleavage catalysis (Figure 3b). Interestingly, each of the two active sites in the RPP1 resistosome binds an ATP molecule, which was supplemented during purification. The bound ATP probably acts as an analogue of NAD\(^+\) at the active site. More interestingly, structural analyses revealed that NADP\(^+\) binds to the TIR domain of RUN1 at a similar position to the ATP-occupied active site (Horsefield et al., 2019; Ma et al., 2020). These observations support the idea that active TNL resistosomes function as a holoenzyme cleaving NAD(P)\(^+\) to transduce the signal in the absence of NAD(P)\(^+\)-bound TNL resistosome structures. However, the identity of the TNL-produced signalling molecules remains elusive.

10 | NUCLEASE ACTIVITIES OF PLANT TIR FILAMENTS

Previous crystal structures of plant TIR domains identified two dimerization interfaces required for function (Zhang et al., 2017a). The functional relevance of the AE dimerization interface was demonstrated in TNL tetramer resistosomes for NADase activity. However, the DE interface
was not observed in TNL tetrameric resistosomes and its functional relevance remains unclear. Interestingly, a recent study showed that the TIR domain of flax TNL L7 can assemble into a filament using the AE and DE interfaces, as proposed previously (Figure 3a). Moreover, the L7 TIR domain was purified in complex with DNA. Cryo-EM studies of the structure of the L7 TIR filament wrapping DNA showed that filament–DNA interaction is critical for TIR nuclease activity and production of 2′,3′-cAMP (Yu et al., 2021). 2′,3′-cAMP molecules were identified in the NAD$^+$/NADH-binding pocket of L7 TIR domains in the TIR filament structure, suggesting that NADase and nuclease activities of TIR domains use similar catalytic sites. Further mutational studies confirmed that 2′,3′-cAMP synthetase activity correlates with TIR cell death function (Yu et al., 2021). It is not clear whether 2′,3′-cAMP functions as a signalling molecule. Thus, plant TIR domains could assemble into filaments and function as nuclease, but the possibility of TIR filament formation in the context of full-length TNLs and the distinct roles of TIR NADase and nuclease activities in TIR signalling remain unknown.

FIGURE 4 Conserved four-helical-bundle (4HB) fold structure of plant NLR CC and CC$^\beta$ domains, and the dynamics of ZAR1 resistosome formation. (a) Superimposition of AtZAR1 CC in green (PDB: 6J5W), Sr33 CC in teal (PDB: 2NCG), AtNRG1.1 CC$^\delta$ in yellow (PDB: 7L7W), and mouse MLKL 4HB in pink (PDB: 4BTF). (b) Structure of AtZAR1 (PDB: 6J5W) in the resting state. The four helices of the CC domain are coloured in purple, orange, green, and blue. The loop region connecting $\alpha$4A and NB is shown as a curved dashed line in red, which becomes a helix termed $\alpha$4B in the active state. (c) Structure of active AtZAR1 (PDB: 6J5T) shown as a monomer from the pentameric resistosome highlighting the $\alpha$1 helix being flipped and penetrating the plasma membrane (PM). The $\alpha$4A in the resting state becomes a potentially flexible loop region not resolved in the electron density map and is indicated as a curved dashed line in blue. The newly formed helix $\alpha$4B is shown in red. (d) Structure of AtZAR1 resistosome (PDB: 6J5T) highlighting the channel pore with a diameter of about 1 nm.

11 | UNDEFINED FUNCTIONS OF LIPASE-LIKE PROTEINS AND RNLS DOWNSTREAM OF TNL

All tested TNLs require lipase-like proteins, including Enhanced Disease Susceptibility1 (EDS1), Senescence-Associated Gene101 (SAG101), and Phytoalexin Deficient4 (PAD4), for function (Gantner et al., 2019; Lapin et al., 2019; Wagner et al., 2013). In Arabidopsis, AtEDS1 and AtSAG101 form a stable heterodimer and cooperate with the AtNRG1s to control cell death, while AtEDS1 and AtPAD4 physically associate and function together with the AtADR1s to mediate bacterial growth restriction and resistance. EDS1, SAG101, and PAD4 contain a lipase-like domain, whose enzymatic activity is dispensable for function, and a plant-unique EDS1-PAD4 (EP) domain. In EDS1/SAG101 and EDS1/PAD4 heterodimers, the pockets between the two EP domains are critical for TNL function and are proposed to be the potential binding sites of TIR-generated small molecules, but are not validated (Sun et al., 2021). More interestingly, activation of the TIR signalling pathway has been shown to trigger the association between EDS1/SAG101 and NRG1s as well as the interaction between EDS1/PAD4 and ADR1s (Sun et al., 2021; Wu et al., 2021). The activation mechanisms of EDS1/SAG101/NRG1 and EDS1/PAD4/ADR1 to initiate cell death and resistance responses, respectively, are currently unknown.

12 | AUTOACTIVE NRG1 AND ADR1 FUNCTION AS A Ca$^{2+}$ CHANNEL

Similar to lipase-like proteins, RNLS including NRG1s and ADR1s are genetic requirements for TNL function (Castel et al., 2019; Qi
et al., 2018; Salle et al., 2020). In the context of ETI, RNLs function cooperatively with lipase-like proteins. Autoactive RNLs or their CC\(\text{R}\) domains trigger EDS1-independent cell death in *N. benthamiana* (Collier et al., 2011; Jacob et al., 2021). The AtNRG1.1 CC\(\text{R}\) domain adopts a 4H structure fold reminiscent of the cell death domain of MLKL that has been shown to function as cation channel. Further investigations demonstrated that autoactive NRG1 mutant and ADR1 function as a Ca\(^{2+}\) channel when expressed in planta and in human cells to trigger cell death (Jacob et al., 2021) (Figure S2a).

13 | NLR SIGNALLING NETWORK AND UNRESOLVED LINKS

Based on current knowledge, a plant NLR functioning network includes three distinct signalling modes. First, some CNL singletons like ZAR1 are both sensors of pathogen effectors and immunity effectors as a Ca\(^{2+}\) channel (Figure S1a). Whether all CNL singletons and pairs function as Ca\(^{2+}\) channels or alternative signalling mechanisms exist remains to be demonstrated. The structural bases of CNL pairs autoinhibition and activation require future investigations. Second, in solanaceous species, phylogenetically related sensor CNLs on effector recognition activate CNL-type helper NRCs to confer resistance and cell death. Interestingly, the pore-forming helix α1 in the CC domain is interchangeable between ZAR1 and NRCs. It is reasonable to speculate that helper NRCs function as Ca\(^{2+}\) channel-like ZAR1 and RNL helpers, which is not confirmed experimentally. The biochemical functions of the sensor CNLs and how effector-activated sensor CNLs further activate helper NRCs are not clear. Whether sensor CNLs are also Ca\(^{2+}\) channels or use different biochemical function to activate NRCs remains to be determined (Figure S1b). Third, effector-activated TNLs function as NADase and nuclease to produce undefined signalling molecules whose receptors remain to be determined. Whether full-length TNLs can form filaments and function as nuclease remains to be demonstrated. The activation mechanisms of EDS1/SAG101/NRG1 and EDS1/PAD4/ADR1 by different TIR enzymatic activities are not clear. The biochemical function of lipase-like family proteins EDS1/SAG101/PAD4 in the TNL signalling pathway is not clear either. In the biologically relevant context of the effector-activated TNL signalling pathway involving EDS1/SAG101 and EDS1/PAD4, whether RNLs indeed function as a Ca\(^{2+}\) channel requires further investigation (Figure S2b). The structural bases of helper RNLs functioning as a Ca\(^{2+}\) channel remain unresolved. The structures of TNL pairs in the resting and active states are not known either. The Ca\(^{2+}\)-responsive factors perceiving the channel activities of CNLs, RNLs, and potential NRCs and their links to immune signalling require further investigation.

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DATA AVAILABILITY STATEMENT

Data sharing is not applicable to this article as no new data were created or analysed.

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**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of the article at the publisher’s website.

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