Death, TIR, and RHIM: Self-assembling domains involved in innate immunity and cell-death signaling

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Abstract
The innate immune system consists of pattern recognition receptors (PRRs) that detect pathogen- and endogenous danger-associated molecular patterns (PAMPs and DAMPs), initiating signaling pathways that lead to the induction of cytokine expression, processing of pro-inflammatory cytokines, and induction of cell-death responses. An emerging concept in these pathways and associated processes is signaling by cooperative assembly formation (SCAF), which involves formation of higher order oligomeric complexes, and enables rapid and strongly amplified signaling responses to minute amounts of stimulus. Many of these signalosomes assemble through homotypic interactions of members of the death-fold (DF) superfamily, Toll/IL-1 receptor (TIR) domains, or the RIP homotypic interaction motifs (RHIM). We review the current understanding of the structure and function of these domains and their molecular interactions with a particular focus on higher order assemblies.

KEYWORDS
higher-order assembly signaling, inflammasome, NOD (nucleotide binding and oligomerization domain) and leucine rich repeat containing receptor (NLR), necrosome, signaling by cooperative assembly formation (SCAF), Toll-like receptor

1 | INTRODUCTION

The innate immune system detects and protects against microorganisms and cellular damage, and it consists of ‘pattern recognition receptors’ (PRRs) that recognize evolutionarily conserved ‘pathogen-associated molecular patterns’ (PAMPs) and endogenous ‘danger-associated molecular patterns’ (DAMPs) released by dying or damaged cells. Activation of these receptors initiates signaling pathways to induce upregulation of cytokine expression, processing of pro-inflammatory cytokines, and programmed cell-death responses. These responses control a pathogenic infection, initiate tissue repair, and stimulate the adaptive immune system.

Structural studies have provided fundamental biological insight into how these pathways are activated and an emerging theme corresponds to ‘signaling by cooperative assembly formation’ (SCAF), which involves assembly of higher order complexes—signalosomes or ‘supramolecular organizing centers’ (SMOCs).1–7 In the resting state, the PRRs are often auto-inhibited through intramolecular interactions, but conformational changes induced by PAMP/DAMP binding result in receptor oligomerization, which nucleates recruitment and oligomerization of cytosolic adaptor proteins, in turn nucleating recruitment and oligomerization of effector enzymes that are subsequently activated through proximity-induced mechanisms. SCAF enables PRRs
to respond rapidly to a low-level stimulus and although it is not an exclusive mechanism to innate immunity and cell-death pathways, it is ideally suited to signaling in these pathways.

SCAF and assembly of higher order complexes have been observed in ‘Toll-like receptor’ (TLR)5 and ‘RIG-I (retinoic acid-inducible gene 1)-like receptor’ (RLR) signaling.8–10 In inflammasome formation initiated by ‘NOD (nucleotide binding and oligomerization domain) and leucine rich repeat containing receptors’ (NLRs), and ‘pyrin and HIN (hematopoietic interferon-inducible nuclear localization 200) amino acid repeat domain containing family’ (PYHIN) receptors,11–13 in ‘receptor-interacting protein kinase’ (RIP) –1 and –3 necroosome assembly stimulated by members of the ‘tumor necrosis factor’ (TNF) and TLR superfamilies,14–18 and in the assembly of ‘CARD-CC/BCL-10/MALT1-like paracaspase’ (CBM) complexes,19,20 which are involved in signaling by some ‘C-type lectin domain receptors’ (CLRs), B- and T-cell receptors in lymphocytes, and the G protein-coupled receptor angiotensin II, and in keratinocyte immunity.21–25 In addition, SCAF has also been proposed to play an important role in effector-triggered immunity in plants.26,27

These signalosomes largely form through homotypic interactions of (i) members of the ‘death fold’ (DF) domain superfamily, (ii) ‘Toll/IL-1 receptor’ (TIR) domains, or (iii) ‘RIP homotypic interaction motifs’ (RHIMs) (Fig. 1). Here, we review the function and structure of these domains, and the molecular interactions mediated by them.

2 | DEATH-FOLD DOMAIN SUPERFAMILY

The DF superfamily is composed of 4 subfamilies, the ‘death domain’ (DD), the ‘death effector domain’ (DED), the ‘caspase recruitment domain’ (CARD), and the ‘pyrin domain’ (PYD), which are all known to form homotypic interactions leading to the formation of oligomeric signaling assemblies such as inflammasomes and the Mydosome (Fig. 2). These domains are found in many multicellular organisms including mammals, Drosophila, zebrafish and C. elegans. DED, CARD, and PYD containing proteins are also found in some viral pathogens, where they are involved in host defense system evasion (reviewed in [28]).

Members of the DF superfamily share a common α-helix bundle (usually 6 helices, H1–H6), with helices arranged anti-parallel in a Greek key type topology surrounding a hydrophobic core (Fig. 3). The arrangement of these helices results in 6 protein binding surfaces, which mediate cooperative interactions with 3 types of interfaces (type I, II, and III). These interfaces facilitate the cooperative assembly of large, oligomeric signaling complexes, and the recruitment of effector enzymes, such as caspases and kinases, which undergo concentration-dependent proximity-induced auto-activation following assembly. Proteins rarely contain the DF domain in isolation, typically this domain is found in combination with other protein–protein interaction domains or effector domains conferring enzymatic activity.

Although all members of the DF superfamily share the same overall fold, the individual subfamilies exhibit distinct structural and sequence differences that confer a degree of specificity. Members of each subfamily interact with a specific array of binding partners. Typically, these are homotypic interactions such as that of ‘apoptosis-associated speck-like protein containing a caspase recruitment domain’ (ASC) CARD recruiting procaspase-1 CARD, and the DD of the TLR adaptor protein ‘myeloid differentiation primary response gene 88’ (MyD88) interacting with the DDs of the ‘IL-1R associated kinase’ (IRAK) -2 and -4. However, the CARD of ‘apoptosis repressor with a CARD protein’ (ARC) has been reported to interact with the DDs of the ‘Fas-associating death domain-containing protein’ (FADD) in vitro,29 and the recruitment of procaspase-8 to inflammasomes via interaction between the ASC PYD and the procaspase tandem DD suggests heterotypic interactions also play a role in the formation of signaling complexes.30

2.1 | The DD subfamily

The DD serves as a homotypic protein–protein interaction domain in numerous intracellular signaling proteins involved in innate immunity and cell-death pathways (Fig. 1). Members of the death receptor subfamily of the ‘tumor necrosis factor receptor’ (TNFR) superfamily mediate cell-death pathways following ligand-induced oligomerization of extracellular receptor domains and subsequent recruitment of DD containing proteins through their own intracellular DDs. The death receptors, ‘first apoptotic signal receptor’ (Fas), ‘death receptor 4’ (DR4), and ‘death receptor 5’ (DR5) interact via a homotypic interaction with the C-terminal DD of FADD. FADD also contains an N-terminal DED and homotypically interacts with the ‘tandem DDs’ (tDDs) of procaspase-8 or -10 to form the ‘death signaling complex’ (DISC). Assembly of the complex activates the initiator caspases that in turn activate effector caspases, such as caspase-3 and -7, leading to apoptosis.31–35

Two other death receptors, TNFR1 and ‘death receptor 3’ (DR3), recruit the adaptor protein ‘TNF receptor type 1-associated protein with death domain’ (TRADD) via the N-terminal DD of TRADD. Recruitment of TRADD can facilitate either pro-cell survival and ‘nuclear factor kappa-light-chain-enhancer of activated B cells’ (NF-κB) activation through the DD:DD recruitment of RIP1 and ‘TNFR associated factor 2’ (TRAF2) termed complex I, or when RIP1 is not ubiquitylated, initiation of apoptosis via DD:DD mediated recruitment of FADD and subsequently procaspase-8, termed complex II.36–39

Outside of the TNFR superfamily, the DD-containing proteins, ‘p53-induced protein with a DD’ (PIDD) and ‘RIP-associated ICH-1 homologous protein with a death domain’ (RAIDD), form an assembly termed the PIDDosome via homotypic interactions between the DDs of PIDD and RAIDD, and between the N-terminal CARDS of RAIDD and procaspase-2. Similar to TRADD signaling, isoforms of PIDD have been reported to recruit RIP1, leading to NF-κB activation and cell survival.40,41

In addition to a C-terminal TIR domain, the TLR adaptor protein MyD88 possesses an N-terminal DD, which facilitates the assembly of an oligomeric assembly complex, termed the Mydosome, through DD:DD interactions with IRAKs, which contain an N-terminal DD and a carboxy-terminal Ser/Thr kinase or pseudokinase domain (Fig. 2). The ‘MyD88 adaptor-like protein’ (MAL) TIR domain nucleates the formation of MyD88 assemblies,5 suggesting that oligomerization of the MyD88 TIR domain serves to cluster the MyD88 DDs, followed by recruitment and proximity based auto- and cross-phosphorylation.
The DED subfamily

The DED subfamily includes FADD, ‘phosphoprotein enriched in astrocytes 15’ (PEA-15), ‘death effector domain-containing protein’ (DEDD), and DEDD2, which all contain a single N-terminal DED, as well as procaspase-8 and -10, the ‘cellular FLICE-like inhibitory proteins’ (cFLIPs), and MC159 (viral FLIP from the Molluscum contagiosum virus), which possess tDEDs (Fig. 1). While FADD is known to interact with the tDEDs of procaspase-8 or -10 to form the DISC, the roles of PEA-15, DEDD, and DEDD2 are unknown. PEA-15 appears to have no catalytic function, acting as a modulator through protein–protein interactions. In particular, PEA-15 appears to inhibit apoptosis by binding to FADD through homotypic DD interactions and preventing the recruitment and activation of procaspases. However, PEA-15 is also implicated in ‘mitogen-activated protein kinase’ (MAPK) signaling through a non-DD interaction with ‘extracellular signal-regulated kinase’ (ERK). DEDD and DEDD2 translocate to the cell nucleus and induce limited apoptotic signaling; however, when nuclear localization is prevented, DEDD and DEDD2 bind to procaspase-8 or -10 in a DED-dependent manner and induce apoptosis. The significance of this signaling in vivo remains unknown.

The structures of the DEDs of FADD, PEA-15, procaspase-8, and MC159 reveal that DEDs display the 6-helical bundle fold representative of the DF superfamily (Fig. 3). In comparison to of IRAKs. IRAKs in turn phosphorylate TRAF6, leading to the activation of transcription factors such as NF-κB and ‘activator protein 1’ (AP-1). The structures of the DDs in the Fas/FADD DISC, PIDD/RAIDD PIDDosome, and MyD88/IRAK2/IRAK4 Myddosome complexes reveal three shared asymmetric interfaces (type I, II, and III; Fig. 3), despite the apparent differences in assembly stoichiometry and morphology.

FIGURE 1 Domain organization of selective proteins containing DF, TIR, SEFIR, and RHIM domains. ^MALT has a paracaspase domain. *cFLIP has a pseudo-caspase domain. CRD, cysteine rich domain; TM, transmembrane; LRR, leucine rich repeat; ZU5, named after the mouse tight junction protein ZO-1 and the C. elegans uncoordinated protein 5; IG, immunoglobulin; NOD, nucleotide-binding oligomerization domain; WD-40, short ~40 amino acid motifs, often terminating in a Trp-Asp (W-D) dipeptide; CC, coiled coil domain; PDZ, acronym derived from the names of the first proteins in which the domain was observed in: post-synaptic density protein 95, Drosophila disc large tumor suppressor and zona occludens 1; SH3, SRC homology 3 domain; GUK, guanylate kinase; HIN, hematopoietic IFN-inducible nuclear antigens with a 200-amino-acid repeat; NTD, N-terminal domain; ARM, armadillo repeat motif domain; SAM, sterile alpha motif domain; TIG, Ig-like domain; 3α-3α-helix structural unit; a/a, helical domain; NB-ARC, nucleotide-binding domain shared by APAF-1, certain R gene products and CED-4; FN, fibronectin; Z, Z-DNA binding domain of IRAKs. IRAKs in turn phosphorylate TRAF6, leading to the activation of transcription factors such as NF-κB and ‘activator protein 1’ (AP-1). The structures of the DDs in the Fas/FADD DISC, PIDD/RAIDD PIDDosome, and MyD88/IRAK2/IRAK4 Myddosome complexes reveal three shared asymmetric interfaces (type I, II, and III; Fig. 3), despite the apparent differences in assembly stoichiometry and morphology.

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FIGURE 2  Death-fold domain assemblies and their role as signaling platforms. (A) Activation of CARMA1 by T cell (TCR) or B cell receptor (BCR) mediated phosphorylation results in oligomerization of CARMA1, possibly due to aggregation of coiled-coil domains, and nucleates the formation of BCL-10 filaments. Recruitment of MALT1 by BCL-10 leads to activation of the MALT1 paracaspase domain, recruitment, and oligomerization of TRAF6 to MALT1, and subsequent NF-κB signaling. Note: The C-terminal MAGUK domain of CARMA1 is not shown. (B) Recognition of PAMPs or DAMPs by TLRs is thought to induce dimerization of the cytoplasmic TLR TIR domains, which then serve as a recruitment platform for the TIR containing adaptor protein MAL. MAL subsequently recruits MyD88 through TIR:TIR domain interactions causing the MyD88 DDs to form a helical assembly with IRAKs through DD:DD interactions, and the resulting proximity based auto- and cross-phosphorylation of IRAKs leads to the activation of pro-inflammatory transcription factors, such as NF-κB and AP-1. For clarity, the IRAK kinase domains are not shown. (C) Activation of the FAS receptor recruits FADD via DD:DD interactions. The resulting clustering of FADD DEDs nucleates the formation of caspase-8 tDED assemblies, this in turn causes clustering, and subsequent proximity based activation of the procaspase. Activated caspase-8 is able to induce pro-apoptotic signaling through processing of procaspase-3. (D) Detection of cytoplasmic dsDNA by AIM2 results in clustering of the AIM2 PYD. The AIM2 PYD is thought to then recruit ASC via PYD:PYD interactions, forming a filamentous assembly. Formation of the ASC PYD filament initiates clustering of the ASC CARDs and subsequent recruitment of procaspases through CARD:CARD interactions leads to caspase activation and signaling.
other DF domains, DEDs contain a surface-exposed hydrophobic patch formed predominantly by residues on helix H2, and a E/D-RxDL-motif, with the N-terminal acidic residue contributed by helix H2, and the C-terminal motif by helix H6. Also, tDEDs have an additional helix H7 at the C-terminus of DED1, linking DED1 to the second DED (DED2). The DEDs of FADD and cFLIP, and tDEDs of procaspase-8 have been shown to form filamentous assemblies in vitro. Comparison of the cryo-EM structure of procaspase-8 tDED assembly with the crystal structure suggests that the tDEDs undergo little to no conformational changes upon oligomerization. FADD DED filaments may act as a template for helical association of procaspase-8 tDED in a mechanism similar to that reported for RIG-I: mitochondrial antiviral-signaling protein (MAVS), and melanoma differentiation associated gene 5 (MDA5):MAVS assemblies (see below) (Fig. 2). The viral FLIP MC159 inhibits the concentration-dependent, proximity-based auto-activation of caspase-8 by binding to the procaspase-8 tDED and preventing further oligomerization.

2.3 | The CARD subfamily

The CARD subfamily (Fig. 1) was first identified in the apoptotic signaling proteins RAIDD, caspase-2, and ‘cell death protein 3’ (CED-3), an orthologue of caspase-9, due to sequence similarity and their known involvement in apoptotic signaling. Originally described
as a motif that facilitated the recruitment of caspases to an apoptotic signaling complex, CARD-containing proteins are now known to have a broad range of functions in multiple immune system signaling pathways. The roles of CARDs can be broadly classified into 3 groups: the CARDs found as pro-domains of caspases, CARDs that act as receptors or adaptors in the assembly of signaling complexes, and CARDs that inhibit or otherwise modulate signaling complexes. The CARD-containing proteins 'apoptotic protease activating factor-1' (Apaf-1), RIG-I, and 'NLR family CARD domain-containing protein 4' (NLRC4), serve as recruitment domains for downstream adaptor proteins or caspases following activation of the receptors' pattern recognition domain (e.g. a leucine-rich repeat domain or WD repeat domain) by PAMP or DAMP ligands such as mitochondrial cytochrome c, viral genomic material, or bacterial flagellin. Adaptor proteins such as ASC, which contains both an N-terminal PYD and a C-terminal CARD, act to link the PYD-containing receptor proteins to caspase recruitment and amplify receptor signaling due to homotypic interactions, with subsequent recruitment, oligomerization, and activation of CARD containing procaspases through CARD:CARD interactions (Fig. 2).

The 'CARD-coiled coil' (CARD-CC) containing proteins, including 'caspase-recruitment domain membrane-associated guanylate kinase (MAGUK) protein' (CARMA) 1–3 and CARD9, nucleate the formation of the CBM complexes, typically upon T- or B-cell receptor, natural killer cell receptor, or dectin-1-induced phosphorylation of the CARD-CC by protein kinase C. In the prototypical CBM, phosphorylation of CARMA1, which is thought to reside in an auto-inhibited state, activates CARMA1, which in turn recruits 'B-cell lymphoma 10' (BCL-10) and nucleates the formation of filamentous BCL-10 assemblies. The CARD-CC proteins CARMA 1–3 contain an N-terminal CARD, central CC domain, and C-terminal MAGUK domain, while CARD9 lacks the MAGUK domain, containing only an N-terminal CARD and C-terminal CC domain. Recruitment of BCL-10, which contains an N-terminal CARD and C-terminal Ser/Thr-rich domain, occurs via CARD:CARMA interactions with CARMA1. The paracaspase, 'mucosa-associated lymphoid tissue lymphoma translocation protein 1' (MALT1), is recruited to the CARMA1/BCL-10 complex through interactions between the CARD and the Ser/Thr-rich domain of BCL-10 and the N-terminal DD and central 'immunoglobulin' (Ig) domains (Ig1 and Ig2) of MALT1 (Fig. 2). The recruited MALT1 rapidly dimerizes, leading to paracaspase activation, recruitment, and oligomerization of TRAF6, induction of TRAF6 ubiquitin ligase activity, and subsequent NF-κB activation.

The structures of several CARDs have been determined in isolation by various methods, and display the 6-helical bundle fold characteristic of the DF superfamily. A feature that appears unique for the DF of CARDs is the morphology of helix α1, which is often kinked or separated into 2 separate helices. As with all members of the DF superfamily, the orientations and lengths of several helices often differ among the individual CARDs, and presumably confer some degree of specificity for the binding partners of CARD-containing proteins. Notably, RIG-1, MDA5, and NOD2 contain 'tandem CARDs' (tCARDs) located sequentially at the N-terminus of the protein.
conditions used for NMR spectroscopy appear to have abolished self-association, and thus provide little insight into the interfaces of the PYD assemblies. However, the cryo-EM structure of the ASC PYD filament at near-atomic resolution demonstrates that few conformational changes are apparent between monomeric PYD and those of the filament. In the filament, the a3 helix and a2-a3 loops appear to be stabilized by the type I, II, and III interfaces. The ASC PYD filaments display a 3-fold helical symmetry, when looking down the helical axis, featuring a hollow core with an inner diameter of ∼20 Å and outer diameter of ∼90 Å. The crystal structure of the AIM2 PYD displays little difference to that of the ASC and NLRP3 PYDs, with the exception of the α1 and α6 helices, which are elongated in the AIM2 PYD.12,69,72 PYDs typically display highly positively or negatively charged surface regions, which are either involved in self-association or conferring specificity for a particular binding partner.12,69

2.5 | Death-fold family oligomerization interfaces

The DF domains display 3 shared asymmetric non-overlapping interfaces that can potentially interact with up to 6 different binding partners (Fig. 3). The interfaces between layers or strands of the DF assemblies may differ depending on the symmetry or oligomerization pattern used to describe the assembly, for instance the Myddosome42 and procaspase-1 CARD assemblies65 have been described as left-handed helices with the type III interface composed of intrastrand interactions, while the ASC PYD filament has been described as a 3-stranded right-handed assembly with intrastrand interactions mediated by the type I interface.12 However the interfaces remain relatively conserved among DF family members.12,42,45

The type I interfaces form between helices H1 and H4 (interface la) of one DF domain and H2 and H3 (interface Ib) of an adjacent domain, and are largely composed of electrostatic interactions. The type II interaction is formed between residues of helices H4 (IIa) of one DF domain and residues of the loop region between helices H5 and H6 of another. Finally, the type III interaction is formed between helix H3 (IIla) of one DF domain and the 2 loop regions connecting helices H1 and H2, and helices H3 and H4, respectively (patch IIlb) of the other DF domain.12,42,45 The procaspase-8 tEDDs differ slightly from this arrangement as within each tDED, the type Ib surface of the first DED (DED1) interacts with the type Ia surface of the second DED (DED2). Furthermore, the type I interface of the tDED filament is predominantly mediated by hydrophobic interactions, whereas the type II and III interactions are complementary electrostatic interactions.49

In mammals TIR domains are found (i) on the cytosolic side of TLRs, which recognize a wide variety of PAMPs and DAMPs either at the cell surface or in the endosomes, and trigger activation of NF-κB and IFN regulatory factor (IRF) transcription factors to induce cytokine production75; (ii) on the cytosolic side of Interleukin-1 receptors (IL-1Rs), which regulate the activities of the IL-1 family of proinflammatory cytokines76; (iii) in the cytoplasmic signaling adaptor molecules MyD88, MAL, ‘TIR domain-containing adaptor protein inducing IFN-β’ (TRIF), and ‘TRIF-related adaptor molecule’ (TRAM), which are recruited to TLRs and IL-1Rs (only MyD88) via TIR:TIR domain interactions upon activation by PAMP/DAMP or cytokine recognition77; (iv) in ‘sterile-α and TIR motif-containing protein’ (SARM), which was first described as an adaptor protein involved in regulation of TLR signaling,78 but has more recently been established as an executor of neuronal cell-death following insults such as oxygen and glucose deprivation, axonal degeneration following nerve transection and vincristine treatment79,80; and (v) in ‘B-cell adaptor for phosphoinositide 3-kinase’ (BCAP), which has recently been proposed to be the sixth TIR domain-containing TLR adaptor, linking phosphoinositide metabolism with negative regulation of TLR pathways.81,82

In plants, TIR domains are the signaling component of a major subclass of disease resistance proteins that recognize pathogen effector proteins introduced into the plant cell during the invasion of the plant. These proteins are typically referred to as plant NLRs, based on their similarity to mammalian NLRs, and they trigger defense responses that often include localized cell death at the site of infection.26 Self-association of these receptors via TIR:TIR interactions is critical for activation of defense responses.

A wide range of both pathogenic and nonpathogenic bacterial species also have TIR domain-containing proteins.83,84 Some of these proteins such as TcpB from Brucella melitensis and TcpC from uropathogenic Escherichia coli CFT073 have been shown to supress TLR signaling in mammals by interfering with receptor–adaptor TIR domain interactions.85,86

‘Similar expression to fibroblast growth factor genes and IL-17R’ (SEFIR) domains, which in mammals are present in both the cytosolic segment of IL-17Rs and in the cytosolic adaptor protein ‘NF-κB activator 1’ (Act1), share similarity with TIR domains, and these 2 domain families have been classified as the STIR (SEFIR/TIR) domain superfam.87 Analogous to the role of TIR domains in TLR and IL-1R signaling, Act1 is recruited to IL-17R complexes via SEFIR:SEFIR interactions upon ligand stimulation.88 SEFIR domains are also found in bacterial proteins, but their functions are not known.89

3 | TIR DOMAINS

The TIR domain is a small globular domain consisting of 125–150 residues, and in animals and plants it is usually found in multidomain proteins involved in innate immunity pathways (Fig. 1). Many bacterial proteins also contain TIR domains, and at least some of them are associated with virulence by suppressing host innate immunity signaling pathways.

3.1 | TIR and SEFIR domain structure

TIR and SEFIR domains have a flavodoxin-like fold, consisting of a central 5-stranded parallel β-sheet surrounded by 4–7 α-helices on both sides of the sheet. The secondary elements and loops are labeled sequentially; for example, the AB loop connects helix αA and strand βB, while the BB loop connects strand βB and helix αB. TIR domain structures are available for the human TLRs 1, 2, 6, and 1090,92; the human TLR adaptors MyD88, MAL, TRAM, TRIF, and BCAP81,86,93–95,97–99; human IL-1R accessory protein-like 1 (IL-1RAPL)100; the Toll-related
receptor TRR-2 from the lower metazoan Hydra magnipapillata (PDB ID 4W8G and 4W8H); the flax NLR protein L6; the Arabidopsis NLR proteins RPS4, RRS1, SNC1, and RPP1; the Arabidopsis protein AtTIR; the grape NLR protein RPV1; the Paracoccus denitrificans protein PdTIR; and the Brucella melitensis virulence factor TcpB.

In addition, SEFIR domain structures are available for the human IL-17RA and IL-17RB receptors and the Bacillus cereus protein BcSEFIR. The structural core of all TIR domains is conserved but there are significant structural differences in the surrounding loops and α-helices (size, number, and orientation), in particular between the different sub-types of TIR domains (TLRs, TLR-adaptors, IL-1Rs, plant, and bacteria) (reviewed in [109]). For example, in plant TIR domains the region between the αD and βE strands contains 3 well-defined helices (αD1, αD2, and αD3) and is significantly different compared to both mammalian and bacterial TIR domains, which only contain 1 or 2 short helices (Fig. 4A). The central 5-stranded β sheet of SEFIR domains as well as the 3 helices αA, αB, and αE superimpose well with the TLR receptor TIR domains, but there are significant structural differences in the CC region, the αC and αD helices, and the DD region (Fig. 4A).

3.2 | TIR domain assemblies

PAMP or DAMP recognition by TLRs drives the intracellular TIR domains to self-associate and subsequently recruit TIR domain-containing adaptor proteins via homotypic TIR-domain interactions to initiate signaling. Although structures are available for many receptor and adaptor TIR domains, the molecular mechanisms underpinning adaptor recruitment to TLRs via homotypic TIR-domain interactions have remained controversial, due to the difficulty of reconstituting stable TIR domain oligomers (reviewed in [3]). Our studies of MAL and MyD88-dependent TLR4 signaling using electron microscopy and in vitro reconstitution assays revealed that the TIR domain of MAL can (i) self-assemble or assemble with the TLR4 TIR domain into filaments, and (ii) nucleate large crystalline assemblies of the MyD88 TIR domain. Both the MAL filament and the MyD88 assembly consist of proto-filaments containing 2 parallel strands of TIR-domain subunits in a head-to-tail arrangement, mediated by the highly conserved BB loop (links the βB-strand to the αB-helix), which has previously been shown to be critical for TLR signaling (Fig. 4B). Although similar-scale TIR-domain assemblies are unlikely to form in the cell during a normal signaling event, the MAL and MyD88 TIR-domain proto-filament structures reflect the molecular mechanisms of TIR:TIR interactions during this process: binding of lipopolysaccharide (LPS) to TLR4 causes the dimerization of its TIRs, which provides a platform for MAL to start an open-ended assembly, which in turn nucleates an open-ended assembly of MyD88 TIRs. Consistent with this model, Latty et al. recently demonstrated, using live cell imaging of macrophages, that TLR4 assemblies into dimers upon LPS stimulation, which rapidly nucleates formation of MyD88 assemblies consisting of 6 or 12 subunits. Several of the residues in the TIR:TIR interfaces in the MAL and MyD88 proto-filaments are highly conserved in mammalian TIR domains, suggesting that this may be a general mechanism of assembly formation in both TLR and IL-1R signaling.

Structures of plant TIR domains involved in effector-triggered immunity have revealed 2 functionally relevant TIR:TIR domain interactions that involve highly conserved residues: the DE interface, which involves residues from the αD, βE, and αE regions, found important for signaling by the L6 NLR from flax; and the AE interface, which involves residues from αA, αE, and the AA and EE loops, found critical for RPS4-mediated signaling in Arabidopsis and RPV1-dependent signaling in the wild grapevine Muscadina rotundifolia. These 2 interfaces are symmetric and different to the asymmetric interactions observed in the MAL and MyD88 proto-filaments, suggesting that the TIR-domain fold may have evolved to mediate different types of homotypic interactions (Fig. 4B). Recent structure-function studies of TIR domains from the SNC1 and RPP1 R proteins demonstrated that both AE and DE interactions control self-association and are required for triggering an immune response. The SNC1 crystal structure also revealed an extended TIR domain superhelix propagated through the AE and DE interfaces, which may facilitate SCAF in plant cell-death signaling (Fig. 4B).

The available crystal structures of bacterial TIR domains, PdTIR from the nonpathogenic Paracoccus denitrificans and TcpB from the pathogenic Brucella melitensis, reveal a common symmetric self-association interface distinct from the plant TIR domains, which involves residues in the DD and EE loops. The BB loops from the 2 interacting molecules are exposed to the solvent for possible interaction with host molecules.

Common interaction interfaces have not yet been reported for SEFIR domains, but functional analyses have identified helix αC as a critical structural motif for heterotypic SEFIR:SEFIR interactions between Act1 and IL-17RA/IL-17RB while the αB helix of Act1 is important for homotypic interactions between Act1 SEFIR domains.

3.3 | TIR domain enzyme activity

SARM has been shown to have TLR-independent roles in neurons and the local cell death program induced by SARM after axonal injury involves rapid breakdown of ‘nicotinamide adenine dinucleotide’ (NAD+) into nicotinamide and ADP-ribose. Enforced dimerization of the C-terminal TIR domain of SARM is sufficient to cause depletion of axonal NAD+ and induce axonal degeneration in the absence of injury and Essuman et al. recently demonstrated that the SARM TIR domain has intrinsic NADase activity and suggested it shares similarities to nucleotide hydrolases and nucleoside transferases. The TLR4 and MyD88 TIR domains did not show NADase activity, suggesting this activity is a unique feature of SARM. Essuman et al. also showed that a glutamate residue, E642, in the predicted αC helix of SARM TIR is critical for the NADase activity. An equivalent glutamate residue is not found in TLR and IL-1R receptor and adaptor TIR domains in mammals, but it is conserved in many bacterial and plant TIR domains, and in a second paper by Essuman et al. it was confirmed that several of the bacterial TIR domains also have NADase activity. For both SARM and the bacterial proteins, clustering of the TIR domains on beads was required for NADase activity, suggesting that TIR domain self-association may be a requirement for enzymatic
FIGURE 4 Structural features of TIR and SEFIR domains, and RHIM regions and their assemblies. (A) Structures of human TLR1 (PDB ID 1FYV), human MAL (PDB ID 5UZB), Brucella melitensis TcpB (PDB ID 4C7M) and flax L6 (PDB ID 3OZI) TIR domains and the human IL-17RA (PDB ID 4NUX) SEFIR domain. The structures are shown in analogous orientations and unique structural elements are highlighted. (B) Left panel: Cartoon representation of the MAL TIR domain proto-filament (PDB ID 5UZB). The assembly consists of 2 parallel strands of TIR-domain subunits in a BB-loop-mediated head-to-tail arrangement. The schematic diagram highlights the assembly interfaces: BB surface, BB loop; EE surface, \( \alpha \)D, \( \alpha \)E strands, and \( \alpha \)E helix; BC surface, \( \alpha \)B and \( \alpha \)C helices; CD surface, \( \alpha \)D helix and CD loop. Right panel: Cartoon representation of a hypothetical assembly of plant TIR domains utilizing the symmetric AE and DE interfaces observed in the SNC1 crystal structure (PDB ID 5TEC). The schematic diagram highlights the assembly interfaces: AE surface, \( \alpha \)A and \( \alpha \)E helices; DE surface, \( \alpha \)D1 and \( \alpha \)E helices and connecting loops. (C) The structure of the RIP1:RIP3 RHIM domain hetero-amyloid (PDB ID 5V7Z). Strands of the amyloid are formed by alternating RHIM domains aligned forming 2 parallel \( \beta \)-sheets; the 2 \( \beta \)-sheets are arranged in an antiparallel fashion and form a hydrophobic core. The hydrophobic core is clearly visible in the crystal structure of the RIP3 homo-amyloid (PDB ID 5ZCK).

activity. Interestingly, the Act1 SEFIR domain has recently been shown to directly bind and stabilize mRNAs encoding key inflammatory proteins, suggesting that nucleotide binding may be a common feature among TIR and SEFIR domains.

4 | RHIM REGIONS

The ‘RIP homotypic interaction motif’ (RHIM) is a short sequence of \(~15–20\) amino acids, first identified as a homotypic interaction motif that facilitates association between RIP1 and RIP3. Located at the C-terminus of RIP3 and adjacent to the C-terminal DD in RIP1, the RHIM domain forms the core of the amyloid signaling complex known as the necosome, which coordinates necroptosis, a regulated form of necrosis that occurs upon inhibition of caspase-8 mediated apoptosis. The RHIM region contains a core motif of (V/I)Q-(V/I/L/C)-G, and is also found in ‘DNA-dependent activator of IFN regulatory factors’ (DAI), TRIF, mouse cytomegalovirus ‘viral inhibitor of RIP activation’ (vIRA), ‘herpes simplex virus’ (HSV)-1 ICP6 and HSV-2 ICP10, and shares similarities with the amyloid-forming domain of the fungal protein HET-s. Necroptosis appears to be activated when recruitment of RIP1 by either TNFR1 or TRIF fails to result in caspase-8 activation, thus sparing the cell from apoptosis. In the absence of apoptosis, RIP1 and RIP3 associate via their RHIM domains, forming an amyloid fibril; oligomerization and subsequent activation of RIP3 leads to the recruitment and phosphorylation of ‘mixed lineage kinase domain-like protein’ (MLKL) via the N-terminal kinase domain of RIP3, triggering MLKL oligomerization,
recruitment to the plasma membrane and cell death, presumably by permeabilization of the membrane or disruption of membrane homeostasis.\textsuperscript{16,18,116,117,124–127} In addition to RIP1-mediated necroptosis, TRIF and DAI have been reported to recruit RIP3, resulting in MLKL activation and necroptosis independent of RIP1.\textsuperscript{118–120} While ICP6 and ICP10 appear to interact with RIP1 and RIP3 and disrupt necroptosis in a RHIM-dependent manner.\textsuperscript{122}

Structures of the RHIM regions are limited to the crystal structure of a homo-amyloid formed by the RIP3 tetrad VQVG and the solid-state NMR structure of the RIP1/RIP3 hetero-amyloid complex (Fig. 4C).\textsuperscript{18} Both the RIP3 homo-amyloid and RIP1/RIP3 hetero-amyloid fibril feature two parallel $\beta$-sheets arranged together in an anti-parallel fashion, with the hydrophobic residues of the (V/I/L/C)-G tetrad forming a hydrophobic core. The overall structure of the RIP1/RIP3 hetero-amyloid complex further reveals that each strand in the parallel $\beta$-sheet is broken into 4 short $\beta$-strand segments separated by short turns. In the RIP1/RIP3 hetero-amyloid fibril, the parallel strands of the $\beta$-sheets are formed from alternating RIP1/RIP3 RHIM domains. In addition to the hydrophobic core, stacking of the parallel RIP1:RIP3 RHIM domains is supported by Asn and Gln ladders formed between the side chains of Asn353 in RIP1 and Asn454 in RIP3 and the side chains of RIP1 residue Gln540 and RIP3 residue Gln459. Additional interactions including tyrosine stacking interactions formed between Tyr534 (RIP1) and Tyr453 (RIP3), and a Ser536 (RIP1)/Cys455 (RIP3) ladder further stabilize the assembly.\textsuperscript{18}

5 | CONCLUDING REMARKS

In innate immunity and cell-death pathways, SCAF is emerging as a key signaling mechanism. The first characterized examples of SCAF involved members of the DF superfamily, but more recently RHIM regions and TIR domains have also been shown to form open-ended assemblies and to engage in SCAF. The higher order assemblies formed by these domains have distinct properties. The RHIM assemblies resemble amyloids and prions, in which adjacent interacting elements are tightly packed, resulting in cooperative contacts and a high barrier of dissociation. The DF signalosomes consist of folded domains, but cooperative interactions among 6 weakly associating adjacent surfaces lead to assemblies that can be as stable as amyloids.\textsuperscript{6,12} By contrast, the 2-stranded head-to-tail assemblies of TIR domains have lower valency and are less stable. The MAL filament, for example, disassembles at low temperatures in solution.\textsuperscript{5} The static and dynamic properties associated with the different assemblies could be directly related to the function of the assemblies and the biological outcome of the signaling pathways they are involved in. For example, in TLR signaling, it is likely that the low valency and observed dynamic properties of TIR domain assemblies are important for the observed rapid assembly and disassembly of TLR4:MAL:MyD88 complexes in macrophages.\textsuperscript{110} On the other hand, DF and RHIM assemblies may persist even after cell lysis.

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DISCLOSURE

The authors declare no conflicts of interest.

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