The caspase recruitment domain–containing protein 9 (CARD9)–B-cell lymphoma/leukemia 10 (Bcl10) signaling axis is activated in myeloid cells during the innate immune response to a variety of diverse pathogens. This signaling pathway requires a critical caspase recruitment domain (CARD)–CARD interaction between CARD9 and Bcl10 that promotes downstream activation of factors, including NF-κB and the mitogen-activated protein kinase (MAPK) p38. Despite these insights, CARD9 remains structurally uncharacterized, and little mechanistic understanding of its regulation exists. We unexpectedly found here that the CARD in CARD9 binds to Zn$^{2+}$ with picomolar affinity—a concentration comparable with the levels of readily accessible Zn$^{2+}$ in the cytosol. NMR solution structures of the CARD9–CARD in the apo and Zn$^{2+}$-bound states revealed that Zn$^{2+}$ has little effect on the ground-state structure of the CARD; yet the stability of the domain increased considerably upon Zn$^{2+}$ binding, with a concomitant reduction in conformational flexibility. Moreover, Zn$^{2+}$ binding inhibited polymerization of the CARD9–CARD into helical assemblies. Here, we also present a 20-Å resolution negative-stain EM (NS-EM) structure of these filamentous assemblies and show that they adopt a similar helical symmetry as reported previously for filaments of the Bcl10 CARD. Using both bulk assays and direct NS-EM visualization, we further show that the CARD9–CARD assemblies can directly template and thereby nucleate Bcl10 polymerization, a capacity considered critical to propagation of the CARD9–Bcl10 signaling cascade. Our findings indicate that CARD9 is a potential target of Zn$^{2+}$-mediated signaling that affects Bcl10 polymerization in innate immune responses.

During fungal infection, fungus-specific carbohydrates bind the C-type lectin receptors in myeloid cells, including Dectin-1, Dectin-2, and Mincle (1–3). These receptors engage, via an intercellular immunoreceptor tyrosine-based activation motif (ITAM)$^4$-like motif or an adaptor protein containing an ITAM-like motif, Syk kinase. Syk, in turn, phosphorylates and activates protein kinase Cδ leading to subsequent phosphorylation and activation of the scaffolding protein CARD9 (caspase recruitment domain–containing protein 9) (4, 5). Upon activation, CARD9 recruits, via its N-terminal CARD, the CARD-containing Bcl10 (B-cell lymphoma/leukemia 10) and subsequently MALT1 to form the myeloid CARD9–Bcl10–MALT1 (CBM) signalosome. Upon complex formation, the CBM signalosome initiates the downstream activation of NF-κB required to mount an antifungal immune response (4, 6). Consistent with the critical and nonredundant role of CARD9 in this pathway, individuals deficient in CARD9 are highly susceptible to chronic fungal infections (7–10). CARD9 has also been shown to contribute to other nonfungal innate immune responses, such as cytosolic DNA sensing with Rad50 (11), intracellular bacterial infections through interaction with NOD2 (12), and viral RNA detection through association with RIG-I (13). The CARD9 signaling axis is implicated in a number of inflammatory diseases, including a genetic association with susceptibility to inflammatory bowel disease (14, 15), development of autoimmune disease of the eye (16), and progression of cardiovascular disease brought on by high-fat diet–induced obesity (17). Indeed, Wang et al. (18) observed that the activation of p38 MAPK by the CARD9–Bcl10 signaling axis induces obesity-

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$^4$ The abbreviations used are: ITAM, immunoreceptor tyrosine-based activation motif; CARD, caspase recruitment domain; ORCH, obesity-related cardiac hypertrophy; NS-EM, negative stain–electron microscopy; H5QC, heteronuclear signal quantum coherence; SOFAST-HMQC, selective optimized flip-angle short-transient heteronuclear multiple quantum coherence; CI, confidence interval; HDX, hydrogen-deuterium exchange; TEV, tobacco etch virus; IPTG, isopropyl β-D-galactopyranoside; RMSD, root mean square deviation; TCEP, tris(2-carboxyethyl)phosphine; PDB, Protein Data Bank; FP, fluorescence polarization; Ni-NTA, nickel-nitrotriacetic acid; MAPK, mitogen-activated protein kinase; MBP, maltose-binding protein; CTF, contrast transfer function.
related cardiac hypertrophy (ORCH) in mice maintained on a high-fat diet. They further found that Zn\(^{2+}\) modulates ORCH development in this context, with Zn\(^{2+}\) deficiency aggravating and Zn\(^{2+}\) supplementation mitigating disease severity in a Bcl10-dependent manner (18). These findings fit within a larger body of literature indicating that Zn\(^{2+}\) acts as a signaling molecule in a number of immune cell types, wherein transient increases in cytosolic Zn\(^{2+}\), known as Zn\(^{2+}\)-waves, serve to activate downstream signaling pathways (19–22). In general, however, the targets of the Zn\(^{2+}\)-wave and mechanisms by which Zn\(^{2+}\) impacts signaling remain poorly understood. Similarly, despite the broad importance of CARD9, CARD9 itself has remained structurally uncharacterized, and a mechanistic understanding of its regulation is lacking.

CARD9 comprises an N-terminal CARD followed by a “coiled-coil” domain of ∼450 amino acids containing multiple distinct regions predicted to have high coiled-coil propensity. The CARD is critical for CARD9’s recruitment of Bcl10, and by homology to CARD9’s closest parologue, CARD11 (CARD11, caspase recruitment domain–containing protein 11; also known as CARMA1), the CARD9–CARD is thought to act in a templating manner by forming a helical assembly able to potentiate the subsequent polymerization of Bcl10 required for signal propagation (6, 23). We therefore hypothesize that CARD9 signaling may be regulated by modulating the accessibility of its CARD and/or its propensity to generate a helical template. To better understand molecular mechanisms underlying CARD9 function, we determined the NMR solution structure of the CARD9–CARD, and we found, surprisingly, that it binds to Zn\(^{2+}\), exhibiting a dissociation constant comparable with estimates of the “free” cytosolic Zn\(^{2+}\) concentration. Although the ground-state structure of the CARD9–CARD is essentially identical in the apo and Zn\(^{2+}\)-bound states, Zn\(^{2+}\) binding strongly stabilizes the fold and reduces conformational “breathing” of the helices. Upon overexpression in Escherichia coli, the CARD9–CARD is also capable of forming an extensively domain-swapped dimer, with interconversion of the CARD monomer and dimer strongly inhibited by Zn\(^{2+}\) binding. Furthermore, Zn\(^{2+}\) binding inhibits formation of helical filaments by the CARD9–CARD monomer that otherwise spontaneously assembles in vitro. A 20-Å negative-stain EM (NS-EM) structure of these filaments reported here demonstrates that they adopt a similar symmetry as the Bcl10–CARD helical assembly. Finally, we show through both a bulk assay and direct NS-EM visualization that the CARD9–CARD helical assembly is capable of directly templating Bcl10 polymerization.

### Results

#### The CARD9–CARD binds Zn\(^{2+}\)

We purified \(^{15}\)N-labeled CARD9–CARD (residues 2–97) using a cleavable affinity tag and size-exclusion chromatography (see under “Experimental procedures”). A \(^{15}\)N HSQC NMR spectrum of the resulting monomeric CARD exhibits significant peak dispersion, suggesting that the domain adopts a well-folded structure. We determined near-complete backbone and side-chain assignments for the CARD and calculated a solution structure with a backbone RMSD of 0.7 Å (RMSD calculated for structured residues 10–97, see Table 1 for complete statistics). The CARD9–CARD adopts a canonical death-domain structure containing six antiparallel α-helices with nine N-terminal residues remaining largely unstructured. As shown in Fig. 1A, the CARD9–CARD aligns well with the crystal structure of its closest parologue, the human CARD11–CARD (1.9-Å backbone RMSD to PDB code 4LWD (23)), including the conserved kink in the α1 helix common among CARDs. The largest differences between the CARD9 and CARD11–CARDs are apparent in α-helices α3 and α4 and the flexible α3–α4 loop, a region with high B-factors in the CARD11–CARD structure and also comprising the largest sequence divergence between the CARDs. Given the role of Zn\(^{2+}\) in modulating ORCH and the central role of CARD9 in this pathology (18), we tested whether the CARD9–CARD itself interacts with Zn\(^{2+}\). The \(^{15}\)N HSQC NMR spectrum of the CARD is significantly perturbed upon addition of 1:0.5 or 1:1 concentrations of ZnCl\(_2\), with all shifted peaks in the slow-exchange limit, suggesting that the CARD binds Zn\(^{2+}\) with sub-micromolar affinity (Fig. 1B). Further increasing the Zn\(^{2+}\) concentration above a 1:1 ratio minimally affects the spectrum, indicating that the CARD9–CARD con-

### Table 1

|                     | Apo CARD | Zn\(^{2+}\)-bound CARD |
|---------------------|----------|------------------------|
| PDB code            | 6E26     | 6E25                   |
| BMRB code           | 30492    | 30491                  |
| Assignments (%)\(^a\) | 89 (97)  | 81 (77)                |
| \(^{1}\)H          | 91 (97)  | 90 (97)                |
| \(^{13}\)C         | 91 (98)  | 72 (49)                |
| \(^{15}\)N         | 76 (96)  | 75 (95)                |
| NOE restraints      | 1509     | 1592                   |
| Intra-residue (i = j) | 423     | 407                    |
| Sequential (i = j–1) | 410     | 421                    |
| Medium range (i = j <5) | 395     | 444                    |
| Long range (i = j ≥5) | 281     | 320                    |
| NOE constraints per restrained residue\(^b\) | 15.9    | 16.8                   |
| Hydrogen bond constraints | 66    | 60                     |
| Long range (i = j ≥5) | 0      | 0                      |
| Dihedral angle constraints | 140    | 140                    |
| Total no. of restricting constraints | 1715 | 1792                   |
| Restricting constraints per restrained residue\(^c\) | 18.1  | 18.9                   |
| Long range (i = j ≥5) | 3.0  | 3.4                    |
| Total structures computed | 100  | 100                    |
| No. of structures included | 20    | 20                     |
| Distance violations per structure | 0.1–0.2 Å | 9.0  | 6.4                     |
| 0.2–0.5 Å           | 2.0     | 1.25                   |
| >0.5 Å              | 0       | 0                      |
| RMS of distance violation/constraint (Å) | 0.02   | 0.02                   |
| Maximum distance violation (Å) | 0.47 | 0.46                   |
| Dihedral angle violations per structure | 1–10° | 4.1 | 2.55                   |
| >10°                | 0       | 0                      |
| RMSD\(^d\)         | 0.52    | 0.33                   |
| Backbone            | 0.7 (1.8) | 0.5 (1.5)            |
| Heavy atoms         | 1.2 (2.2) | 1.0 (1.9)            |
| Ramachandran\(^e\) | 93.6    | 94.9                   |
| Most favored (%)    | 93.6    | 94.9                   |
| Additionally allowed (%) | 6.4    | 5.0                    |
| Generously allowed (%) | 0    | 0.1                    |
| Disallowed (%)      | 0       | 0                      |

\(^a\) Total assignment completeness, with backbone completeness reported in parentheses.

\(^b\) Residues 3–97 contain conformational restraining constraints.

\(^c\) Residues 10–97 reported, with all-residue RMSDs reported in parentheses.

\(^d\) Residues 10–97, calculated with Procheck.

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**Zinc binding modulates CARD9–CARD nucleation of Bcl10**

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contains a single high-affinity metal-binding site (Fig. S1). To determine whether this binding is specific for Zn^{2+}, we incubated the CARD9–CARD with other divalent metal ions. No chemical shift changes are observed upon addition of 1:10 molar concentrations of either CaCl_{2} or MgCl_{2}, indicating that the CARD is unable to bind Ca^{2+} or Mg^{2+} (Fig. 1C). A stoichiometric concentration of NiCl_{2} likewise induces no chemical shift changes in the CARD9–CARD. However, Mn^{2+} and Cu^{2+} do interact with the CARD9–CARD at stoichiometric concentrations, with Cu^{2+} inducing chemical shift changes and the paramagnetic Mn^{2+} inducing enhanced relaxation of a number of peaks (Fig. 1C). Upon simultaneous addition of Zn^{2+} and Mn^{2+} to the CARD9–CARD in a 1:1 ratio, the peak signature indicates that all of the CARD is Zn^{2+}-bound, with remaining peak disappearances attributable to secondary Mn^{2+} interactions outside the Zn^{2+}-binding site (Fig. 1C, see arrows indicating peaks unique to the Zn^{2+}-bound CARD9–CARD). Simultaneous addition of stoichiometric concentrations of Cu^{2+} and Zn^{2+} results in populations of the CARD9–CARD bound to each of the two metals, shown by the presence of peaks corresponding to both the Cu^{2+}-bound and Zn^{2+}-bound states. The Zn^{2+}-bound peaks are ~50% less intense than when Zn^{2+} is added alone (Fig. 1C, arrowheads), indicating approximately equal affinity of the CARD for Zn^{2+} and Cu^{2+}. These findings are consistent with the relative affinities expected by the Irving–Williams series (24). The concentration of labile cytosolic copper has proven difficult to measure with high accuracy, although estimates suggest that it is in the femtomolar range or lower (25, 26). Because the CARD9–CARD dissociation constant for Zn^{2+} (and therefore for Cu^{2+}) is several orders of magnitude larger than the typical cytosolic Cu^{2+} concentration, but comparable with estimates of the “free” cytosolic Zn^{2+} concentration (see below), we suggest that Zn^{2+} is likely to be a physiological ligand for CARD9, but depending on the cellular context, Cu^{2+} binding could play a role as well. We thus proceeded to characterize Zn^{2+} binding to the CARD9–CARD.

Cysteine and histidine residues typically mediate Zn^{2+} coordination, with additional binding often provided by glutamate and aspartate side chains. To determine which residues in the CARD9–CARD are responsible for Zn^{2+} coordination, we generated alanine substitutions at either of the two cysteines (CARD^{C10A} or CARD^{C37A}) or the sole histidine (CARD^{H73A}),
the positions of which are depicted in Fig. 1D. Although CARD<sup>C37A</sup> continues to bind Zn<sup>2+</sup> in the slow-exchange limit, both CARD<sup>C10A</sup> and CARD<sup>H73A</sup> shift binding to the intermediate-to-fast exchange regime, indicating a significant reduction in affinity and demonstrating that Cys-10 and His-73 are involved in coordinating Zn<sup>2+</sup> (Fig. 1E). The only glutamates or aspartates potentially in position to provide additional Zn<sup>2+</sup> coordination are a stretch of acidic residues (Glu-5, Asp-7, Asp-8, and Glu-9) on the unstructured N-terminal region of the CARD9–CARD. Upon mutation of Glu-5 or Glu-9 to Gln, we found no substantial differences in the Zn<sup>2+</sup>-bound spectrum, indicating no contribution to coordination. In contrast, we found that the Zn<sup>2+</sup>-bound spectrum of CARD<sup>D7N</sup> differs substantially from CARD<sup>WT</sup>, despite the binding remaining in the slow-exchange regime (Fig. S2). These data suggest that Asp-7 is less critical than Cys-10 or His-73 for Zn<sup>2+</sup> affinity but that coordination by Asp-7 results in conformational changes to the CARD. CARD<sup>D8N</sup> exhibits subtler Zn<sup>2+</sup>-bound <sup>15</sup>N HSQC differences as compared with CARD<sup>WT</sup>, comprising reduced line-broadening of the binding-site–proximal amide peaks Cys-10, Trp-11 (backbone and side chain), Ser-12, and Gln-69 (side chain), suggesting a role for Asp-8 in coordination as well (Fig. S2).

The CARD9–CARD exhibits a picomolar dissociation constant for Zn<sup>2+</sup>

Zn<sup>2+</sup> binding to the CARD9–CARD is sufficiently tight to preclude direct affinity determination by NMR titration. We instead utilized competition against the fluorescent Zn<sup>2+</sup>-binding dye mag-fura-2, which has a dissociation constant of 20 nM for Zn<sup>2+</sup> (27). As shown in Fig. 2A, the CARD9–CARD competes effectively against mag-fura-2, although less effectively than EDTA (K<sub>D</sub> of ~10<sup>-16</sup> for Zn<sup>2+</sup>). Fitting these data to the exact competitive binding equation described by Wang (28), CARD<sup>WT</sup> binds Zn<sup>2+</sup> with a dissociation constant of 0.73 nM (95% confidence interval (CI) 0.45–1.07 nM). Consistent with the NMR data, CARD<sup>C10A</sup> and CARD<sup>H73A</sup> are unable to compete with mag-fura-2, whereas CARD<sup>C37A</sup> binds with comparable affinity to CARD<sup>WT</sup>. The mutations D7N and D8N each decrease binding affinity ~2–3-fold, whereas the double mutant CARD<sup>D7N/D8N</sup> exhibits an ~25-fold decrease, suggesting that the two acidic residues may trade-off responsibility for coordinating the Zn<sup>2+</sup> ion (Fig. 2C). Because the measured picomolar affinity of the CARD9–CARD is approaching the lower limit accessible in competition with the 20 nM mag-fura-2, we additionally assessed affinity among the tightly binding constructs in competition with the more tightly binding indo-1 dye (K<sub>D</sub> for Zn<sup>2+</sup> of 0.16 nM (29)). As shown in Fig. 2, B and C, affinities as measured in competition with indo-1 agree with those determined with mag-fura-2, confirming that the CARD9–CARD binds Zn<sup>2+</sup> with a picomolar dissociation constant, comparable with estimates of the free cytosolic Zn<sup>2+</sup> concentration.

Zn<sup>2+</sup> binding does not significantly alter the CARD9–CARD structure

Given the high affinity and specificity that the CARD9–CARD exhibits for Zn<sup>2+</sup>, we were curious as to the impact of Zn<sup>2+</sup> binding on the CARD structure. We thus determined near-complete backbone and side-chain chemical shift assignments for the Zn<sup>2+</sup>-bound CARD9–CARD and calculated the NMR solution structure to a backbone RMSD value of 0.5 Å (RMSD calculated for structured residues 10–97, see Table 1 for complete statistics).

In the Zn<sup>2+</sup>-bound structure, the Zn<sup>2+</sup> ion forms a bridge between Cys-10 at the beginning of α1 and His-73 at the beginning of α5 (Fig. 3A). Because of the ambiguity in coordination by Asp-7 and Asp-8, we only imposed constraints to maintain coordination by Cys-10 and His-73 during structure calculations. Although full quantum mechanical calculations were not performed, Asp-7 interacts with the Zn<sup>2+</sup> ion with one or both carboxyl oxygens in all of the 20 lowest energy structures, consistent with its prominent role in coordination (representative structures shown in Fig. 3A). The apo and Zn<sup>2+</sup>-bound struc-
tatures are remarkably similar, with a backbone RMSD (residues 10–97) of 0.95 Å, which is only slightly higher than the RMSD of the apo state itself (Fig. 3B). We further manually compared the NOESY spectra of the CARD in the apo and Zn\(^{2+}\)-bound states in search of more subtle differences in the structures. Although the specific NOE cross-peaks used in the two calculations vary somewhat due to differential peak overlap and line-broadening between the apo and Zn\(^{2+}\)-bound states, we were unable to conclusively identify any instances in which an NOE cross-peak was present in one state and not the other. We therefore conclude that Zn\(^{2+}\) binding does not significantly alter the ground-state solution structure of the CARD9–CARD.

**Zn\(^{2+}\) binding stabilizes the CARD9–CARD and inhibits α-helical unraveling**

Although binding of a Zn\(^{2+}\) ion to the CARD9–CARD does not substantially alter its ground-state structure, we were curious as to the potential impact of Zn\(^{2+}\) binding on the CARD stability and conformational dynamics. We monitored denaturation of the CARD9–CARD monomer via differential scanning fluorimetry and found that addition of Zn\(^{2+}\) increases the thermostability of the CARD by nearly 14 °C, reflecting a substantial stabilization of the domain upon Zn\(^{2+}\) binding (Fig. 4).

To monitor conformational stability, we performed an NMR-based hydrogen-deuterium exchange (HDX) experiment to monitor the solvent accessibility of backbone amides in the apo and Zn\(^{2+}\)-bound states. Aqueous \(^{15}\)N-labeled CARD9–CARD was lyophilized and then resuspended in 99.99% D\(_2\)O, followed by a collection of a series of SOFAST-

**Figure 3. Solution structure of the CARD9–CARD in the Zn\(^{2+}\)-bound state.** A, two lowest energy Zn\(^{2+}\)-bound CARD9–CARD structures, demonstrating Zn\(^{2+}\) coordination by Cys-10, His-73, and Asp-7. B, alignment of the 20 lowest energy structures of the CARD9–CARD in the apo (black) and Zn\(^{2+}\)-bound (red) states.

**Figure 4. Zn\(^{2+}\) binding stabilizes the CARD9–CARD and slows HDX.** A, differential scanning fluorimetry melt curve for apo (black) and Zn\(^{2+}\)-bound (red) CARD9–CARD. Inflection points are indicated by dashed vertical lines. Two technical replicates are shown for each condition, which agreed to within 0.1 °C of the mean values shown on the graph. B, representative NMR HDX peak-height decay curve for Thr-31 in the absence (black) and presence (red) of equimolar Zn\(^{2+}\). Circles represent \(^{15}\)N-SOFAST-HMQC peak intensities, and dotted lines are best-fit single exponential decay curves. C, CARD9–CARD, lowest energy Zn\(^{2+}\)-bound solution structure. Residues for which Zn\(^{2+}\) binding increased the HDX lifetime, which includes all observed peaks, are colored cyan. The indole amide of Trp-11 also exhibits enhanced protection and is colored cyan. Those residues for which lifetimes could be calculated in both the apo and Zn\(^{2+}\)-bound states and were increased greater than 5-fold by Zn\(^{2+}\) binding are colored red. D, global CARD9–CARD backbone amide HDX exchange lifetimes in the absence (black) and presence (red) of equimolar ZnCl\(_2\). Error bars represent profile likelihood 95% confidence intervals. Residues are excluded for which no signal remained at the first time point or for which overlap precluded accurate peak height determination.

HMQC experiments, which allow for rapid data collection. Approximately 35% of residues remain at least partially protonated by the first 1.5-min time point. For all residues that we were able to monitor, the HDX lifetime was significantly increased in the context of Zn\(^{2+}\) binding, with half-lives increasing by 1.5–14–fold over the apo state (Fig. 4, B and D). The residues most strongly protected by Zn\(^{2+}\) binding map predominantly to helices α4 and α5, which lie on either side of His-73 (Fig. 4C). These data demonstrate that Zn\(^{2+}\) binding locks the CARD in a more stable compact conformation, with less conformational breathing in the helices than in the apo state.

**The CARD9–CARD can adopt a domain-swapped dimeric structure**

Upon recombinant overexpression of the CARD9–CARD in E. coli and subsequent purification, two distinct species can be
isolated: the monomeric CARD for which NMR structures are shown in Fig. 4, and a kinetically stable dimeric state that accounts for \( \frac{20\%}{H} \) of the purified protein at the final gel filtration step (Fig. 5A). We determined the 1.36-Å resolution X-ray crystal structure of this CARD9–CARD dimer (see Table 2 for complete statistics) using molecular replacement with the CARD11–CARD structure (PDB code 4LWD). The dimer is composed of two six-helix bundles, each of which aligns well to the CARD9–CARD monomer (Fig. S3A). Each bundle, however, contains three helices from each of the two polypeptide chains, forming a domain-swapped dimer with a short linker crossing over between helices \( H3 \) and \( H4 \) (Figs. 5B and Fig. S3B).

The \( Zn^{2+} \)-binding site in the CARD9–CARD is distal from the strand swap between \( H3 \) and \( H4 \), such that the domain swap would not be expected to alter \( Zn^{2+} \) binding. Consistent with this expectation, NMR chemical shift changes upon addition of \( Zn^{2+} \) to the dimer are nearly identical to those seen for the monomeric state. We thus soaked \( Zn^{2+} \) into domain-swapped dimer crystals, identified a single condition where \( Zn^{2+} \) occupies one of the two binding sites, and solved the crystal structure at a resolution of 1.81 Å (see Table 2 for complete statistics). The \( Zn^{2+} \) ion binds where expected based on our NMR structure of the \( Zn^{2+} \)-bound monomer, with clear electron density demonstrating its coordination by both Cys-10 and His-73 (Fig. 5C and Fig. S3B). Additional electron density is present, which suggests \( Zn^{2+} \) coordination by a third residue in the N-terminal tail; however, the residues N-terminal of Asp-9

| Table 2 |
| Structural statistics for the CARD9 CARD apo and \( Zn^{2+} \)-bound domain-swapped dimer crystal structures |
| --- |
| Apo CARD dimer | \( Zn^{2+} \)-bound CARD dimer |
| PDB code | 6E28 | 6E27 |
| Wavelength | 1.000 | 1.000 |
| Space group | \( P_{121} \) | \( 1 \) |
| \( a, b, c (\AA) \) | 43.98, 37.43, 56.88 | 44.01, 37.46, 56.96 |
| \( \alpha, \beta, \gamma (\degree) \) | 90.101, 47.90 | 90.101, 79.90 |
| Resolution range (Å) | 37.94–1.36 (1.41–1.36) | 43.08–1.81 (1.88–1.81) |
| \( R_{merge} \) (%) | 3.905 (84.11) | 7.536 (120) |
| \( I/I_{max} \) | 21.67 (1.71) | 14.06 (1.37) |
| Completeness | 95.72 (73.23) | 98.24 (95.65) |
| Redundancy | 6.3 (5.2) | 6.6 (6.6) |
| Resolution (Å) | 1.36 | 1.81 |
| Unique reflections | 37,560 (2849) | 16,495 (1582) |
| \( R_{work/R_{free}} \) (%) | 0.1797 / 0.1937 | 0.2066 / 0.2577 |

Non-hydrogen atoms

| Protein | 1524 | 1498 |
| Ligands | 0 | 1 |
| Water | 153 | 89 |

Average \( B \)-factor (Å²)

| Protein | 36.48 | 44.78 |
| Ion | 63.81 |
| Water | 41.34 | 45.46 |

RMSD

| Bond length (Å) | 0.009 | 0.011 |
| Bond angles (°) | 1.01 | 1.05 |

Ramachandran

| Favored (%) | 99.45 | 98.89 |
| Allowed (%) | 0.55 | 1.11 |
| Outliers (%) | 0 | 0 |
are too poorly resolved to conclusively identify which (likely Asp-7 or Asp-8) is participating in the coordination, likely reflecting conformational heterogeneity in the coordination throughout the crystal. In agreement with the minimal ground-state differences observed between the apo and Zn²⁺-bound monomeric structures, no notable structural changes are observed in the domain-swapped dimer upon Zn²⁺ binding (Fig. 5C).

The domain-swapped dimer exhibits a similar ¹⁵N HSQC spectrum as the monomer, but with distinct differences, especially for those residues near the strand swap. By monitoring a single amide peak (Ser-28, Fig. 5D) with resolvable monomeric and dimeric chemical shifts, we were able to track the kinetics of interconversion of the two states. At 25 °C, we found that the dimer and monomer interconvert with a half-life of 4.1 h (95% CI 3.8–5.4 h) in the absence of Zn²⁺. This interconversion is dramatically slowed, however, in the presence of Zn²⁺, where the dimer exhibits a half-life of 179 h (95% CI 155–206 h), representing a nearly 50-fold decrease in interconversion rate (Fig. 5E). There additionally appears to be a shift in the monomer–dimer equilibrium, with ~3-fold more dimer present at equilibrium in the presence of Zn²⁺ relative to the apo state. These findings are consistent with the HDX data, suggesting that the presence of Zn²⁺ locks the CARD9–CARD in a more stable conformation, preventing helical unraveling that must be required for the monomeric and domain-swapped dimeric conformations to interconvert.

We observed that the homologous CARD11–CARD (50% identity to the CARD9–CARD) is also capable of adopting a relatively long-lived dimeric conformation upon overexpression in E. coli, with an in vitro half-life of 34 min (95% CI 28–48 min) at 25 °C (Fig. S4, A–C). We were unable to conclusively demonstrate that the CARD11–CARD adopts a homologous domain-swapped dimeric structure as the CARD9–CARD, but the long half-life and a comparable extent of chemical shift changes intimate a similar structure. Upon addition of stoichiometric ZnCl₂ to the ¹⁵N-labeled CARD11–CARD monomer, we observed no significant NMR chemical shift perturbations, indicating that Zn²⁺ binding is not conserved within the protein family (Fig. S4D).

The CARD9–CARD forms in vitro filaments in a Zn²⁺-regulated manner

In the absence of Zn²⁺, we found that upon concentrating monomeric CARD9–CARD above ~150 μM, the solution became cloudy. We visualized this opaque solution by NS-EM and observed that the CARD9–CARD monomer assembles into long filaments with a diameter of ~90 Å (Fig. 6A). To monitor the effect of Zn²⁺ binding on these filaments, we purified the CARD9–CARD monomer bound to a stoichiometric amount of Zn²⁺ by adding a saturating concentration of ZnCl₂ prior to a final gel-filtration column. In contrast to the apo CARD, we found that at 200 μM the Zn²⁺-bound CARD remained clear by eye and filament-free as monitored by NS-EM (Fig. 6B). Addition of EDTA to chelate the Zn²⁺ away from the CARDs leads to formation of filaments within ~10 min at 25 °C. We monitored filament formation though UV absorbance at 350 nm and found that they form readily at 200 and 150 μM, but minimally at 100 μM (Fig. 6C). Doubling the salt concentration to 300 mM also effectively blocks CARD9–CARD polymerization at these protein concentrations, a property that permitted NMR data collection and structure determination of the apo CARD9–CARD monomer described above (Fig. 1A). These assemblies are readily reversible upon re-binding of Zn²⁺, as addition of Zn²⁺ stoichiometrically equal to the EDTA concentration induces disassembly within ~5 min. Unlike the monomeric CARD9–CARD, the domain-swapped CARD9–CARD dimer solution remains clear at concentrations of >2 mM, irrespective of the presence of Zn²⁺, indicating that the dimeric state of the CARD is unable to form filaments.

We found that Zn²⁺ binding inhibits filament assembly of the CARD9–CARD monomer but does not block it entirely. Upon concentrating the Zn²⁺-bound CARD9–CARD solution to ~800 μM, it also becomes cloudy, and filaments can be observed by NS-EM (Fig. S5A). These filaments are ~180 Å in diameter and appear to be tandem assemblies of two filaments, as they often end ~90 Å wide, off-center “tails.” After addition of Zn²⁺ to the single-width filaments induced by Zn²⁺ depletion (as depicted in Fig. 6, A and C), we found that all single-width filaments had disassembled when visualized by NS-EM, whereas a small population (undetectable by UV absorbance, Fig. 6C) of tandem filaments had formed, presumably from a subset of filaments that had been able to bind Zn²⁺ and adopt a stabilized tandem conformation prior to disassembly (Fig. S5A). In the context of the full CARD9 protein, the coiled-coil domain would necessarily protrude from any helical CARD assembly, likely blocking side-mediated interactions of the filaments. We therefore anticipate that these observed tandem CARD9–CARD filaments are likely an in vitro artifact. Nonetheless, the vast majority of filaments rapidly disassemble in the presence of Zn²⁺, demonstrating that Zn²⁺ binding regulates the stability of CARD9–CARD helical assemblies.

CARD9–CARD filaments comprise a similar helical assembly as Bcl10 and are able to template Bcl10 nucleation

Because CARD9 is thought to propagate signaling via nucleation of Bcl10 helical assemblies, we wondered whether these in vitro filaments are representative of the helical template that seeds Bcl10 polymerization. To determine whether the filaments adopt a conformation consistent with this nucleating capacity, we determined an ~20-Å resolution NS-EM structure of the CARD9–CARD filaments (Fig. 6D, left). The CARD9–CARD filaments are 90 Å in diameter and form a hollow helical assembly with a 5-Å rise and 102° rotation, which are nearly identical to the 5.0-Å rise and 100.8° rotation determined previously for the Bcl10–CARD helical assembly (30). Direct comparison of low-resolution NS-EM structures of the CARD9–CARD and Bcl10–CARD assemblies (Fig. 6D, right) (EMD-5729 (23)) demonstrates that although the slight differences in helical symmetry lead to discernable differences over several turns of the helix, the two CARDs adopt highly similar filamentous structures. Although the Bcl10 CARD contains an extended C-terminal helix that enables unambiguous fitting of the CARD monomers into the filament structure, the CARD9–CARD contains no such large asymmetry, preventing us from independently placing our CARD monomeric structure con-
clusively into the EM density. However, given the similarity of the helical symmetry between the assemblies, we predict that a higher resolution CARD9–CARD structure would reveal an orientation comparable with the Bcl10–CARD filament. The CARD9–CARD filaments are thus consistent in structure with what we would expect in a templating assembly.

To directly monitor the capacity for the CARD9–CARD filaments to nucleate Bcl10, we adapted a fluorescence polarization (FP)-based Bcl10 nucleation assay described by Qiao et al. (23). Briefly, a Bcl10 construct was generated linked N-terminally to MBP via a TEV protease-cleavable linker; this MBP tag was shown to block in vitro Bcl10 polymerization, which otherwise occurs rapidly for either the full-length Bcl10 or the Bcl10 CARD alone. Bcl10 was also sparsely labeled with Alexa Fluor 488 dye prior to a final gel-filtration column. Bcl10 polymerization is induced by addition of TEV protease, which removes >50% of the MBP in under 2 min and nearly all MBP within 10 min, irrespective of the presence of CARD9–CARD (Fig. S5B). Subsequent Bcl10 polymerization is then monitored by the change in FP corresponding to the increased molecular weight of the filament.

As shown in Fig. 6F, addition of CARD9–CARD filaments to MBP–Bcl10 at a 5:1 molar ratio accelerates the formation of Bcl10 filaments relative to Bcl10 alone, although the addi-
tion of monomeric CARD9–CARD slows Bcl10 polymerization significantly, presumably by competing with Bcl10 homotypic binding sites. We additionally performed a replicate of the experiment utilizing an independent preparation of the CARD9–CARD filaments and purification of MBP–Bcl10. As shown in Fig. 5SC, we observed nearly identical results with an identical replicate (2 μM MBP–Bcl10) and additionally found comparable CARD9–CARD-induced acceleration utilizing a lower concentration of 1 μM MBP–Bcl10.

The capacity for the CARD9–CARD filaments to accelerate bulk Bcl10 polymerization could stem either from direct CARD–CARD templating wherein the Bcl10–CARD helical assembly extends continuously from the CARD9–CARD assembly or from indirect effects, e.g. increasing local Bcl10 concentration. To distinguish these possibilities, we visualized CARD9–CARD filament-nucleated Bcl10 filaments shortly (2 min) after addition of TEV protease by NS-EM. Because Bcl10 contains a C-terminal Ser/Thr-rich domain in addition to its CARD, its filaments are appreciably wider than the CARD9–CARD filaments (Fig. 6, F and G). As shown in Fig. 6H for the nucleated sample, we were able to readily identify continuous filaments comprising regions of both CARD9–CARD and Bcl10, demonstrating direct CARD9–CARD–CARD-templated nucleation of the Bcl10 helical assembly. We were unable to find any instance of more than a single CARD9–Bcl10 transition within a given filament, suggesting that nucleation, like Bcl10 filament extension (30), is unidirectional.

Discussion

Although total eukaryotic cellular Zn2+ concentrations are typically hundreds of micromolar, the vast majority of it is sequestered by tight interactions with proteins (31). Indeed, ~10% of the human proteome has been estimated to bind Zn2+ in structural, catalytic, or regulatory capacities (32). Considerable effort has been made to address the challenging question of what concentration of free Zn2+ is readily available in the cytosol of eukaryotic cells, with estimates ranging from 5 pM to 1.67 nM (33, 34); however, over diverse cell types and detection methods, most studies have measured concentrations in the range of 0.1–1 nM (35–40). Structural binding sites (e.g., zinc finger domains) typically bind Zn2+ with dissociation constants much lower than the free or readily available cytosolic Zn2+ concentration, and therefore, they remain saturated under all conditions. In contrast, regulatory binding sites typically bind with dissociation constants comparable with the picomolar cytosolic Zn2+ concentration, such that fluctuations in local Zn2+ concentration can modulate function via changes to the Zn2+-bound status of a given protein (31).

The immune system modulates Zn2+ concentrations from the organismal to the subcellular levels over time scales of seconds to hours in response to diverse stimuli (21, 22, 41). Within immune cells, several stimuli have been shown to induce a Zn2+-wave wherein cellular stores of Zn2+ are released on the order of minutes after signal initiation, acting as a second messenger; these include stimulation of IgE in mast cells that activate MAPK, extracellular signal-regulated kinase (ERK), and c-Jun N-terminal kinase (JNK) signaling (20) and activation of monocytes by a range of stimuli, including lipopolysaccharide, inducing downstream NF-κB and MAPK signaling (19). Generally, the mechanism of Zn2+ release and the targets of increased cytosolic Zn2+ remain unknown. Specific to CARD9 signaling, Wang et al. (18) demonstrated that Zn2+ deficiency exacerbates cardiac hypertrophy in response to diet-induced obesity and that the onset of disease and an observed Zn2+-mediated rescue depend on the activation of p38 MAPK by the CARD9–Bcl10 signaling axis.

Here, we have demonstrated that the CARD9–CARD specifically binds to Zn2+ with picomolar affinity, akin to proteins that utilize Zn2+ in a regulatory role; the in vitro affinity suggests a regulatory role for Zn2+ binding to CARD9; however, further characterization will be required to determine to what extent the cellular context influences the binding of Zn2+ to the CARD9–CARD. To our knowledge, this is the first observation of metal binding in the CARD family or the larger death-domain family. Among the close paralogues to CARD9 (CARD11, CARD10, and CARD14), the coordinating cysteine and histidine are not conserved (Fig. S7A), consistent with the lack of Zn2+ binding observed for the CARD11–CARD (Fig. S4D). The histidine and cysteine are conserved within CARD9 orthologues among mammals but not in reptiles or more divergent species (Fig. S7B).

Although Zn2+ binding does not substantially alter the CARD9–CARD structure, it does increase its stability, reducing the conformational flexibility of the domain. Moreover, Zn2+ dramatically affects the polymerization propensity of the CARD9–CARD. Indeed, the most striking in vitro impact of Zn2+ binding that we observed was the modulation of filament formation, shown in Fig. 6, A–C. Although it has been assumed that CARD9, like CARD11, propagates signaling by forming a nucleating seed, Fig. 6 represents the first direct evidence that the CARD of CARD9 is capable of forming a helical assembly, that this assembly closely mirrors the symmetry of the Bcl10 helical assembly, and that the CARD is capable of directly templating Bcl10 polymerization. The micrometer length filaments shown in Fig. 6A were generated with high concentrations of a CARD9 construct lacking the coiled-coil region and are thus unlikely to reflect the size of CARD9 assemblies that would form in cells. Rather, we suggest a model wherein CARD9–CARDs are brought to a high local concentration by coiled-coil domain-mediated oligomerization, driving formation of a CARD9–CARD helical “seed” that acts to template and thereby nucleate assembly of Bcl10 filaments. Given the role of Zn2+ in modulating this template formation, we predict that the primary role of Zn2+ binding in CARD9 will prove to be in modulating the propensity to form this template and hence propagate signaling. Within dendritic cells where CARD9 functions, cytosolic zinc concentrations have been shown to decrease during maturation, potentially serving to prime the CARD9–CARD for subsequent signaling events (42). Alternatively, a Zn2+-wave type of transient Zn2+ increase could serve to blunt excessive signaling though CARD9 by promoting helical disassembly after stimuli. A detailed temporal study of zinc levels during maturation and stimulation will be required to tease out the specific mechanisms by which Zn2+ binding may modulate the CARD9 signaling axis.
Zinc binding modulates CARD9–CARD nucleation of Bcl10

The similarity between the NMR solution structures of the apo and Zn$^{2+}$-bound CARD9–CARD (Fig. 3) along with the dramatic differences in stability and conformational flexibility (Fig. 4) suggest that the CARD within the helical assembly may require conformational rearrangement as compared with the monomer in solution. This phenomenon was recently demonstrated for the Bcl10 CARD, for which significant structural rearrangement was observed in a 4-Å resolution cryo-EM structure of the helix as compared with the monomeric NMR solution structure (30). Indeed, the largest differences between the Bcl10–CARD structures are in the orientation of helix α1, which in the CARD9–CARD contains Cys-10 and would thus be conformationally restricted by Zn$^{2+}$ binding. Unfortunately, as was reported for filaments of the Bcl10 CARD alone, CARD9–CARD filaments present primarily as single filaments on NS-EM grids but almost exclusively as massive bundles of filaments (approximately micrometer diameter) when imaging by cryo-EM is attempted. The presence of these large bundles may also help to explain the relatively weak bulk solution nucleating capacity of the CARD9–CARD filaments (Fig. 6E), as they necessarily sequester large numbers of unproductive CARDs. David et al. (30) were ultimately able to determine a cryo-EM structure of the Bcl10–CARD filament using a construct containing the Ser/Thr-rich domain that is disordered relative to the CARD core and therefore absent in the reconstruction. Nevertheless, the Ser/Thr-rich domains serve to block side-to-side filament associations and are observable en masse on each individual filament, allowing us to distinguish the Bcl10 filaments from the thinner CARD9–CARD filaments (30). A similar strategy in CARD9 may allow for future high-resolution structure determination, which would provide insight into both homotypic and heterotypic CARD–CARD interactions, as well as into the specific mechanism by which Zn$^{2+}$ binding modulates CARD9–CARD helical assembly.

As is common in the death-domain family, the CARD9–CARD engages in interactions with other CARDs (Fig. 6, D and H), generating helical assemblies in which the individual domains share a common orientation relative to the helical axis (43). The symmetric nature of the domain-swapped dimer would interfere with this assembly, explaining why the dimer is unable to incorporate into filaments. We thus speculate that the domain-swapped CARD9–CARD dimer (Fig. 5) may act as a negative regulator of CARD9 signaling that could be modulated by Zn$^{2+}$ binding. The domain-swapped structure, however, is formed between CARD9–CARDs under the high concentrations of E. coli overexpression and outside of the context of the full proteins. Further characterization of the full-length protein under physiological conditions, perhaps by utilizing conformationally specific antibodies or an engineered protein deficient in domain swapping, will be required to determine whether the domain-swapped conformation is indeed biologically relevant.

In conclusion, we have identified and structurally characterized multiple conformations accessible to the CARD9–CARD, including a monomer, a domain-swapped dimer, and a filamentous helical assembly (graphically summarized in Fig. 7). CARD9 binds to Zn$^{2+}$ with picomolar affinity, which modulates interconversion between these states, stabilizing the CARD9–CARD ground-state conformation and restricting its capacity to form Bcl10-nucleating filaments. We have thus identified CARD9 as a potential target of Zn$^{2+}$-mediated signaling during innate immune responses.

Experimental procedures

Protein purification

CARD9 and CARD11–CARDs (residues 2–97 for CARD9 and 8–109 for CARD11) were expressed in BL21(DE3) cells with an N-terminal, TEV protease-cleavable His$_6$ tag. Protein production was achieved by growth for 48–72 h at 16 °C in TB autoinduction media or $^{15}$N autoinduction media for unlabeled and $^{15}$N-labeled protein, respectively (44). $^{13}$C,$^{15}$N-Labeled protein was generated by induction in $^{13}$C,$^{15}$N minimal media with 0.5 mM IPTG for 6 h at 37 °C. Proteins were purified by Ni-NTA (Qiagen), overnight cleavage by TEV protease, removal of imidazole by dialysis, and removal of TEV and any remaining uncleaved protein by Ni-NTA. Final purification and separation of monomeric and dimeric species were achieved via a Superdex 75 gel-filtration column (GE Healthcare). For all proteins used in metal-binding experiments, 5 mM EDTA was added prior to the final gel-filtration column. For proteins used in CARD9–CARD filament assays, 500 μM ZnCl$_2$ was added after the second Ni-NTA column, prior to concentration before the gel-filtration step.

For the Bcl10 FP assay, an E. coli expression construct was generated comprising an N-terminal His$_6$ tag followed by MBP, a TEV cleavage site, Bcl10(C29A/C10A), a short linker, and an HA peptide (MBP–Bcl10). The cysteines were mutated to allow for labeling exclusively on the Ser/Thr-rich domain and not the CARD. The ORF was designed to exactly replicate the construct generated by Qiao et al. (23), with the addition of GSGSYPYDVPDYA at the C terminus. MBP–Bcl10 was expressed in BL21(DE3) cells in LB media, which were induced by addition of 0.2 mM IPTG at $A_{600}$ 0.7 for 1 h at 37 °C. MBP–Bcl10 was purified by Ni-NTA and eluted in ~3 ml, to which 2 μM Alexa Fluor 488 C$_5$ maleimide (ThermoFisher Scientific)
was added, followed by incubation for 10 min at room temperature. The labeled protein was then loaded directly with no concentration onto a Superdex 200 gel-filtration column (GE Healthcare) in 20 mM Tris, 150 mM NaCl, 0.5 mM TCEP, pH 7.5. Both the FP assay and preparation of grids for NS-EM were initiated within 2 h of the protein eluting from the gel-filtration column.

**NMR assignments and solution structure determination**

For apo assignments and structure determination, monomeric $^{13}$C,$^{15}$N-labeled CARD9–CARD was purified in 50 mM Tris, 300 mM NaCl, 0.5 mM TCEP, pH 7.0, and concentrated to 400 $\mu$L for all NMR experiments. All experiments were collected at 37 °C on an 800-MHz Bruker spectrometer with a cryogenically cooled probe. Backbone assignments were collected at 37 °C on an 800-MHz Bruker spectrometer HCCH-TOCSY, and $^{13}$C NOESY-HSQC (aliphatic and aromatic, 150-ms mixing times) and $^{13}$C NOESY-HSQC (150-ms mixing time) experiments. Additional $^{15}$N NOESY-HSQC (150-ms mixing time) and $^{13}$C NOESY-HSQC (aliphatic and aromatic, 150 ms mixing times) experiments, with additional $^{15}$N HSQC (aliphatic and aromatic), HCCH-TOCSY, and $^{13}$C NOESY-HSQC (aliphatic and aromatic, 150 ms mixing times) experiments. Additional $^{15}$N HSQC (aliphatic and aromatic) experiments, were transferred from the apo form, utilizing $^{15}$N HSQC, CBCA(CO)NH, HNCA, HN(CA)CO, and (H)CC(CC)NH experiments. Side-chain assignments were determined using $^{13}$C HSQC (aliphatic and aromatic), HCCH-TOCSY, and $^{13}$C NOESY-HSQC (aliphatic and aromatic, 150 ms mixing times) experiments. Additional $^{15}$N NOESY-HSQC (150-ms mixing time) and $^{13}$C NOESY-HSQC (aliphatic and aromatic, 150 ms mixing times, dissolved in 99.99% D$_2$O) experiments were collected to assist in structure determination.

For the Zn$^{2+}$-bound CARD9–CARD, monomeric $^{13}$C,$^{15}$N-labeled CARD was purified up to 400 $\mu$L followed by addition of 480 $\mu$L ZnCl$_2$. Backbone and side-chain assignments were transferred from the apo form, utilizing $^{15}$N HSQC, HNCA,CB, HNCA, $^{13}$C HSQC (aliphatic and aromatic), HCCH-TOCSY, and $^{13}$C NOESY-HSQC (aliphatic and aromatic, 150 ms mixing times) experiments, with additional $^{15}$N NOESY-HSQC (150-ms mixing time) and $^{13}$C NOESY-HSQC (aliphatic and aromatic, 150 ms mixing times, dissolved in 99.99% D$_2$O) experiments were collected to assist in structure determination. Stereoscopic assignments for valine and leucine methyl groups were determined for the Zn$^{2+}$-bound CARD9–CARD by expressing the protein in M9 media with a 1:913C/12C glucose ratio and collection of $^{13}$C HSQC spectra as described by Senn et al. (45); these stereoscopic assignments were subsequently transferred to the spectra of the apo protein. All spectra were referenced directly ($^1$H) or indirectly ($^{13}$C and $^{15}$N) to an internal 2,2-dimethyl-2-silapentane-5-sulfonic acid standard. All spectra were processed using Bruker TopSpin version 3.5 and subsequently analyzed in CcpNMR Analysis version 2.4 (46).

For both apo and Zn$^{2+}$-bound samples, a small proportion of the monomeric protein converted to the domain-swapped dimer over the course of extended NMR data collection; however, the concentration remained sufficiently low as to not register in the 3D experiments and was therefore ignored for both sequential assignments and structure calculation.

For structure determination of both the apo and Zn$^{2+}$-bound CARD9–CARD, dihedral angles were estimated using TALOS+ (47). For the Zn$^{2+}$-bound CARD, restraints were enforced to maintain coordination by Cys-10 5y and His-73 N81. NOE peaks were assigned and initial structure determination was achieved using the CYANA version 3.97 NOE assignment and structure determination package (48, 49). Sum of $r^{-6}$ averaging was used for all NOEs. For each round of CYANA NOE assignment and structure determination, 100 structures were generated, with the 20 lowest target function structures proceeding to the next round. After the final round of NOE assignments, 100 structures were calculated and subsequently refined in explicit water using the PARAM19 force field in CNS version 1.2 (50, 51) and the WaterRefCNS package developed by Dr. Robert Tejero. The 20 lowest energy structures for each of the apo and Zn$^{2+}$-bound refinements in water are presented here. Structures were evaluated using PROCHECK-NMR, with statistics presented in Table 1. All structural depictions were generated in PyMOL.

**Metal binding by NMR**

$^{15}$N HSQC spectra were collected on 100 $\mu$L $^{15}$N-labeled WT and mutant CARD9–CARDs in 50 mM HEPES, 300 mM NaCl, 0.5 mM TCEP, pH 7.0, on a 600-MHz Bruker spectrometer at 37 °C, unless otherwise noted. $^{15}$N-SOFAST HMQC spectra of 150 $\mu$L $^{15}$N-labeled CARD11–CARD with or without 150 $\mu$L ZnCl$_2$ were collected in 50 mM HEPES, 300 mM NaCl, 0.5 mM TCEP, pH 7.0, on an 800-MHz Bruker spectrometer at 25 °C. Stock concentrations of ZnCl$_2$, MnCl$_2$, CuCl$_2$, and NiCl$_2$ were determined via inductively coupled plasma MS.

**Metal competition assays**

Competition assays were performed in 20-µL volumes in 10 mM HEPES, 150 mM NaCl, pH 7.5. Wildtype (WT) or mutant CARD9–CARDs were mixed 1:1 with either mag-fura-2 or indo-1 dyes, with final concentrations of 10 $\mu$L each. Stock CARD9–CARD concentrations were determined by measuring absorbance at 280 nm; mag-fura-2 and indo-1 stock concentrations were determined by measuring absorbance at 369 and 346 nm, respectively, in the presence of EDTA. CARD9–CARD and dye were incubated at 25 °C with the indicated concentrations of Zn$^{2+}$ for 15 and 45 min for mag-fura-2 and indo-1, respectively, to ensure that measurements were made under equilibrium conditions. For mag-fura-2 samples, Zn$^{2+}$ binding was monitored by measuring emission at 497 nm upon excitation at 325 nm. For indo-1 samples, Zn$^{2+}$ binding was monitored by measuring emission at 460 nm upon excitation at 320 nm. All measurements were made on a Molecular Devices SpectraMax M5e plate reader. Data were fit to the exact competitive binding equation described by Wang (28) using GraphPad Prism 7.

**Differential scanning fluorimetry**

Protein denaturation was monitored by differential scanning fluorimetry using the NanoTemper Prometheus NT.48 for both data collection and analysis. 100 $\mu$L CARD9–CARD was prepared with or without addition of 100 $\mu$L ZnCl$_2$ in 50 mM HEPES, 150 mM NaCl, 0.5 mM TCEP, pH 7.5. Temperature was increased at 1 °C/min, and the protein-folding state was monitored via the ratio of tryptophan fluorescence at 350 and 330 nm. Melting temperature was determined as the inflection point of the 350/330 ratio.
Zinc binding modulates CARD9–CARD nucleation of Bcl10

Hydrogen-deuterium exchange

250 μM 15N-labeled monomeric CARD9–CARD samples were generated in 50 mM HEPES, 300 mM NaCl, 250 μM ZnCl2, 0.5 mM TCEP, pH 7.0, with or without 1 mM EDTA. Samples were lyophilized and resuspended in 99.99% D2O, followed by immediate collection of 15N-labeled SOFAST-HMQC experiments at 25 °C on a Bruker 600-MHz spectrometer for time points shown. The mid-point of the SOFAST-HMQC experiments was used as the time points for both plotting and fitting. For all resolved peaks, peak height decay curves were fit to a single-phase exponential using GraphPad Prism 7.

Crystallography and structure determination

Dimeric WT CARD9–CARD was purified in 20 mM Tris, 150 mM NaCl, 0.5 mM TCEP, pH 7.0, and concentrated for crystallization. Apo crystals were generated by vapor diffusion at 19 °C in 0.4-μl sitting drops by mixing 20 mg/ml CARD9–CARD dimer and 1.1 M ammonium tartrate dibasic, pH 7.0 (Hampton Research), at a 1:1 ratio. Crystals were transferred into a cryo-protectant solution of the crystallization solution supplemented with 20% glycerol and frozen in liquid nitrogen. Crystals into which Zn2+ was soaked were generated by vapor diffusion at 19 °C in 0.4-μl sitting drops comprising a 1:1 mix of 20 mg/ml CARD9–CARD dimer and 5% v/v tacsimate, 0.1 M HEPES, pH 7.0, 10% w/v PEG monomethyl ether 5,000 (Hampton Research). These crystals were transferred into the crystallization buffer supplemented with 3 mM ZnCl2 and incubated ~16 h at 19 °C. Crystals were then transferred into a cryo-protectant solution of the crystallization solution supplemented with both 1 mM ZnCl2 and 20% glycerol and subsequently frozen in liquid nitrogen.

Diffraction images were collected at the Advanced Light Source beamline 5.0.2. Data were indexed, integrated, and scaled using XDS and XSscale (53). Both structures were solved by molecular replacement with Phaser-MR within the Phenix package (54, 55), using the monomeric CARD11–CARD structure (PDB code 4LWD) as a search model for the apo structure and the apo domain-swapped dimer structure as a search model for the Zn2+ bound structure. For both structures, iterative cycles of model building in COOT (56) and refinement in Phenix were used to generate final models. All structural depictions were generated in PyMOL.

Monomer–dimer interconversion kinetics

300 μM 15N-labeled dimeric CARD9–CARD samples were generated in 50 mM HEPES, 300 mM NaCl, 500 μM ZnCl2, 0.5 mM TCEP, pH 7.0, with or without 1 mM EDTA. Samples were transferred to 25 °C followed by immediate collection of SOFAST-HMQC (52) experiments at 25 °C on a Bruker 600-MHz spectrometer. Between time points, samples were incubated at 25 °C. For CARD11, 180 μM 15N-labeled dimeric CARD11–CARD was prepared in 50 mM HEPES, 300 mM NaCl, 0.5 mM TCEP, pH 7.0; the sample was generated by concentrating fractions from the dimer peak in Fig. S4A at 4 °C for ~1 h, followed by immediate data collection at 25 °C. The mid-point of the SOFAST-HMQC experiments was used as the time points for both plotting and fitting. Monomeric and dimeric peak heights were fit simultaneously to a single-phase exponential for each sample using GraphPad Prism 7 using amide peak of Ser-28 (CARD9) or the peak boxed out in Fig. S4B (CARD11).

CARD9–CARD filament formation

Zn2+–bound CARD9–CARD was prepared by saturating with Zn2+ prior to a final size-exclusion purification step. 50 μl of Zn2+–bound CARD9–CARD was prepared in 50 mM Tris, 150 or 300 mM NaCl, 0.5 mM TCEP at the indicated concentrations in a 384-well clear-bottom plate. At t = 0, Zn2+ was removed by addition of 250 μM EDTA. Filament formation was monitored by measuring absorbance at 350 nm on a Molecular Devices SpectraMax M5e plate reader while shaking at 25 °C. At the indicated time point, an additional 250 μM ZnCl2 was added to each well, and monitoring was continued. Samples for EM were taken just prior to EDTA addition, just prior to Zn2+ addition, and at the end of the assay.

Bcl10 fluorescence polarization assay

The assay was performed in a 20-μl volume in 20 mM Tris, 150 mM NaCl, 0.5 mM TCEP, pH 7.5, with 1 or 2 μM final concentration of MBP–Bcl10 as indicated. CARD9–CARD filaments were prepared as in Fig. 6C by addition of 250 μM EDTA to 200 μM Zn2+–saturated CARD9–CARD, followed by incubation with shaking at 25 °C for 90 min. The CARD9–CARD monomer control was treated identically to the filament sample, but without addition of EDTA. At the initiation of the experiment, TEV protease was added to 0.05 mg/ml along with CARD9–CARD monomer or filaments as indicated. Fluorescence polarization was measured by exciting at 495 nm and monitoring at 519 nm on a Molecular Devices SpectraMax M5e plate reader at 25 °C. We note that by the 90-min end point of the FP assay at 25 °C, small numbers of filaments form in the MBP–Bcl10 sample in the absence of TEV, indicating that the MBP is not absolute in its ability to block polymerization (Fig. S5D). These filaments are sufficiently sparse as to not register on the FP assay (Fig. 6E) and much thicker than the Bcl10 filaments that form upon TEV cleavage, ensuring that we are not observing them in Fig. 6, F and H.

EM and helical reconstruction

Negative stain samples were generated by incubating on glow-discharged carbon on 400-mesh copper grids (Electron Microscopy Sciences) for 30 s, followed by staining with 2% uranyl acetate. For Fig. 6, A and B, and Fig. S5, A and D, and micrographs used for helical reconstruction of the CARD9–CARD filaments, 4-μl samples were applied at 200 μl with no dilution. For Fig. 6, F and H, samples were prepared by mixing MBP–Bcl10 (2 μM), CARD9–CARD filaments (10 μM), and/or TEV (0.05 mg/ml) as indicated and incubating at room temperature for 2 min followed by direct application of 4 μl onto the grid, with no dilution.

Grids were imaged using a Talos F200C microscope operated at 200 kV and Ceta camera (ThermoFisher Scientific). Images for helical reconstruction were collected at 2.006 Å/pixel; all other images were collected at 4.097 Å/pixel. For helical reconstruction of the CARD9–CARD filament, filaments were manually picked using the EMAN2 program e2helixboxer (57), and...
all subsequent processing steps were performed utilizing routines in Spring (58). CTF parameters were determined using Miccftdetermine, which utilizes CTFFIND (59). Micrographs were CTF corrected and segmented using Segment, yielding 16,754 segments, which were classified into 50 classes using Segmentclass. Six of these classes (chosen by visual inspection) were analyzed using Segclassreconstruct, which computes a 3D reconstruction based on a single class average over a set of incremented helical symmetries (i.e. rise and rotation); the projections of these reconstructions are quantitatively compared against the original class average to identify helical symmetries compatible with the class average. Of those helical parameters that returned a high-correlation coefficient, the 3D reconstructions were visually inspected to identify structures that the resolution is compatible with the globular CARD9–CARD. These helical parameters were then used as an input to Segmentrefine3D to iteratively refine the structure from the segment stack, beginning from a cylinder of radius 100 Å. Finally, the 2D projections and power spectra of the reconstructions were compared against the class averages; only the segment stack, beginning from a cylinder of radius 100 Å. Finally, the 2D projections and power spectra of the reconstructions were compared against the class averages; only the reported parameters of a 5-Å rise and 102° rotation yielded a projection and power spectrum that matched the class averages (Fig. S6, A–C). The final reconstruction utilized 11,226 segments. Using six independent class averages, we ran Segclassreconstruct with fine spacing (0.1 Å and 0.1°) around 5 Å and 102°. The highest correlation symmetry parameters in all cases were within ±0.1 Å and ±0.5° of the reported symmetry, providing an estimate of the uncertainty in these values. Fourier shell correlation analysis (Fig. S6D) of the final structure indicates a resolution of 13.0 Å; however, we were unable to discern significant structural details beyond the location of the individual CARD9–CARDs, and so we suggest by comparison with other comparable structures that the resolution is ~20 Å.

**Sequence alignment**

Multiple sequence alignments were performed using Clustal Omega (60).

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