Structures of autoinhibited and polymerized forms of CARD9 reveal mechanisms of CARD9 and CARD11 activation

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CARD9 and CARD11 drive immune cell activation by nucleating Bcl10 polymerization, but are held in an autoinhibited state prior to stimulation. Here, we elucidate the structural basis for this autoinhibition by determining the structure of a region of CARD9 that includes an extensive interface between its caspase recruitment domain (CARD) and coiled-coil domain. We demonstrate, for both CARD9 and CARD11, that disruption of this interface leads to hyperactivation in cells and to the formation of Bcl10-templating filaments in vitro, illuminating the mechanism of action of numerous oncogenic mutations of CARD11. These structural insights enable us to characterize two similar, yet distinct, mechanisms by which autoinhibition is relieved in the course of canonical CARD9 or CARD11 activation. We also dissect the molecular determinants of helical template assembly by solving the structure of the CARD9 filament. Taken together, these findings delineate the structural mechanisms of inhibition and activation within this protein family.
Aspase recruitment domain (CARD)-containing protein 9 (CARD9) and CARD-containing protein 11 (CARD11, aka, CARMA1) are paralogs proteins that act in a conserved manner as scaffolding proteins required to propagate signaling in immune cells. CARD9 functions in myeloid cells during innate immune responses to activate NF-κB and p38 MAPK. The most well-characterized triggers of CARD9 signaling are fungal carbohydrates, which interact with C-type lectin receptors and ultimately contribute to a Th17-type immune response critical for proper control of fungal infections. CARD11 acts in lymphoid cells downstream of T- and B-cell receptors and ultimately contributes to a Th17-type immune response critical for proper control of fungal infections.

CARD9 and CARD11 share an N-terminal domain architecture comprising an N-terminal CARD followed by ~300 residues with high coiled-coil propensity, referred to as the coiled-coil domain. CARD11 additionally contains a C-terminal membrane-associated guanylate kinase (MAGUK) domain, as well as a linker domain between the coiled-coil and MAGUK domains known as the inhibitory domain (ID). Both CARD9 and CARD11 are thought to be held in autoinhibited states prior to activation, although the specific nature of these states remained uncharacterized prior to this study. CARD11 is activated via phosphorylation of serine residues in the ID by protein kinase C, which interferes with a complex network of interactions between the ID, CARD, and coiled-coil domains required to maintain the autoinhibited state. CARD9 activation has been reported by different groups to require either phosphorylation at T231 by PKCs or ubiquitination at Lys125 by TRIM62. To our knowledge, no single study has investigated both of these modifications together, and, absent structural characterization of the system, no mechanistic explanation for CARD9 activation by either of them has been suggested.

Upon activation, both CARD9 and CARD11 recruit the same downstream binding partner, B-cell lymphoma/leukemia 10 (Bcl10), which interacts through its own CARD with the CARD of CARD9 or CARD11, a CARD–CARD interaction critical for subsequent signal propagation and NF-κB activation. The CARDs of CARD9 or CARD11 are thought to form a nucleating helical template that promotes polymerization of Bcl10 via the Bcl10 CARD, which, along with other domains of activated CARD9/11, then recruits downstream factors, including MALT1, cIAPs, TRAF6, and the LUBAC complex that mediates subsequent ubiquitination of multiple members of the complex, including both linear and K63-linked poly-ubiquitination of Bcl10. These ubiquitination events ultimately lead to activation of IKK and subsequent phosphorylation, ubiquitination, and degradation of IkB, thereby releasing NF-κB to translocate to the nucleus and induce transcription. Regulation of the pathway at CARD9/11 is thus likely achieved by modulating assembly of the helical CARD template.

A number of disease-associated mutations have been identified in CARD9 and CARD11. In CARD9, genetic deletion in mice or loss-of-function mutations in humans lead to chronic fungal infections. One intriguing CARD9 splice variant, identified as protective in inflammatory bowel disease, eliminates a TRIM62 binding site, preventing CARD9 signaling and bolstering the idea that TRIM62 ubiquitination of CARD9 is critical for its activation. In CARD11, genetic deletion in mice or loss-of-function mutations in humans lead to immunodeficiency due to defects in activation and proliferation of both T- and B-cells. A large number of mutations have also been identified in CARD11 associated with a range of lymphoproliferative disorders, including diffuse large B-cell lymphoma (DLBCL). Many of these mutations have been further characterized and shown to exhibit constitutive CARD11 signaling and aberrant cell proliferation, confirming CARD11 as an oncogene. Among these studies, the N-terminal CARD and coiled-coil domains emerged as mutational “hot-spots”, leading Chan et al. to conduct a high-throughput mutagenesis screen on the N-terminal 140 residues of CARD11, in the context of the full-length protein, in search of hyperactivating CARD11 mutants. This study identified 23 sites with hyperactivating mutations in CARD11, many of which mapped to a short stretch of residues (112–130) termed the “LATCH”, that was shown to be critical in maintaining CARD11 in an autoinhibited state.

Here, we present a structure of the N-terminal region of CARD9, which exhibits an extensive autoinhibitory interface required to prevent constitutive activation in both CARD9 and CARD11. From this structural insight, we then define the distinct structural mechanisms of activation in CARD9 and CARD11 and demonstrate that, upon activation, both proteins form helical templates that directly nucleate Bcl10 polymerization. Finally, solving a Cryo-EM structure of the CARD9 CARD helical assembly, we structurally characterize the activated form of this protein family.

Results
An extensive CARD-coiled-coil interface in CARD9. Both CARD9 and CARD11 are thought to be maintained in an autoinhibited state, and the structural basis for this inhibition had remained an open question. To probe the mechanism of inhibition, we expressed and purified 15N-labeled regions of both CARD9 and CARD11 comprising the CARD and the first predicted segment of coiled-coil that mediates homo-dimerization of each construct (CARD92-152 and CARD118-172, see Fig. 1a). Notably, CARD118-172 has been shown by Qiao et al. to be sufficient to accelerate Bcl10 polymerization in vitro. While CARD118-172 remained stable and soluble at high concentrations, the construct exhibited a “molten globule” 15N-TROSY spectrum (Fig. 1b), indicative of significant conformational heterogeneity and/or self-association. While SEC-MALS analysis of CARD118-172 indicated a molecular weight of 36.5 kDa, consistent with an expected dimeric molecular weight of 39.4 kDa, dynamic light scattering analysis at 100 μM revealed a polydispersity index of 0.40, indicating some self-association, which was corroborated by a slight improvement in the nuclear magnetic resonance (NMR) spectral quality upon twofold dilution (Supplementary Fig. 1A). In contrast to CARD118-172, the CARD11 CARD alone exhibits a well-dispersed spectrum indicative of a single dominant conformational state.

Unlike CARD118-172, the homologous CARD92-152 construct (see Supplementary Fig. 1B for sequence alignment) exhibited a well-dispersed 15N-TROSY spectrum with fewer than 150 peaks (Fig. 1c, black), consistent with a single dominant conformation and minimal self-association. We found that, at high concentrations required to collect three-dimensional (3D) NMR spectra for chemical shift assignment and structure determination efforts, CARD92-152 formed higher-order oligomers over the course of several hours, severely impacting NMR spectral quality (Supplementary Fig. 1C). We have shown previously that the CARD9 CARD is capable of forming an extensively domain-swapped dimer, and suspected that these oligomers were due to swap-dimer-mediated “daisy-chaining” of two or more coiled-coil-mediated dimers. Zn2+ binds to the CARD with subnanomolar affinity and slows interconversion of the monomer and domain-swapped CARD dimer ~50-fold, so we generated CARD92-152 complexed 1:1 with Zn2+, which slowed oligomer formation sufficiently to allow for 3D NMR data collection on the dimer (Supplementary Fig. 1C).
As shown in Fig. 1c, the CARD92–152 spectrum diverged significantly from that of the CARD9 CARD alone (CARD92–97), indicating that the CARD is not simply tethered to the coiled-coil, but instead makes extensive contact with the coiled-coil. After assigning the NMR backbone amide chemical shifts of CARD92–152, we found that large changes in amide peak positions map predominantly to one face of the CARD, containing α-helices 2, 5, and 6, suggesting these regions as the interaction interface (Supplementary Fig. 1D).

The ten C-terminal residues of CARD92–152 appear largely unstructured, and elimination of them minimally perturbed the 15N-TROSY spectrum. We thus purified and determined near-complete backbone and sidechain chemical shift assignments for CARD92–152, collected distance restraints in the form of non-specific and intermolecular-specific NOEs, collected residual dipolar coupling restraints, and determined the solution structure of the CARD92–152 dimer with a backbone RMSD among ordered residues of 1.1 Å (Fig. 1d, see Table 1 for statistics). While the overall backbone RMSD is 1.1 Å, alignment to a single CARD or the coiled-coil domain alone yields backbone RMSDs of 0.7 Å and 0.8 Å, respectively, (Supplementary Fig. 1E).

The CARD92–152 structure consists of the N-terminal CARD, a linker (residues 98–111) comprising a short extended strand followed by a two-turn α-helix and a sharp turn at Gly111, and then 8 turns of α-helix. This final helical element forms a canonical dimeric coiled-coil interface consisting of Val and Leu residues. α-helices 2 and 5 of each CARD are packed against both the linker α-helix and both coiled-coil helices, with additional contacts made between the linker and CARD helices 5 and 6 (Fig. 1e, f). The CARD-linker-coiled-coil interface (hereafter referred to as the CARD-coiled-coil interface) is predominately mediated through hydrophobic and aromatic packing, including Tyr33, Leu85, Tyr86, and Tyr91 of the CARD, Val102, Phe103, Met105, Ile106, and Ile107 of the linker, and Leu115, Leu118, Met120, and Met124 of the coiled-coil (Fig. 1f). There appear to be additional hydrogen bonding or electrostatic interactions mediated by Gln36, Glu81, Glu84, Arg101, Ser110, Gln117, and Glu122; however, these interactions are not sufficiently consistent among the structures determined to conclusively state which.

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Table 1 Structural statistics for the CARD92–142 dimer

| Assignments (%)a | 79 (94) |
|------------------|---------|
| 1H               | 72 (91) |
| 13C              | 90 (96) |
| 15N              | 78 (95) |
| NOE restraints   | 3917    |
| Intra-residue [i=1] | 1102    |
| Sequential [i]=| 998     |
| Medium range [i]-j| 770     |
| Long range [i]=j| 764     |
| Intermolecular   | 283     |
| Hydrogen bond constraints | 180 |
| Dihedral angle constraints | 428 |
| RDC restraints   | 166     |
| Total number of restricting constraints | 4691 |
| Restricting constraints per restrained residueb | 16.9 |
| Long range [1]=j| 3.8     |
| Total structures computed | 100 |
| Number of structures included | 20 |
| Distance violations per structure | 0.02 |
| 0.1–0.2 Å       | 47.15   |
| 0.2–0.5 Å       | 10.15   |
| > 0.5 Å         | 0       |
| R.m.s. of distance violation per constraint (Å) | 0.45 |
| Dihedral angle violations per structure | 0.07 |
| 1–10°           | 17.3    |
| > 10°           | 0.3     |
| R.m.s. of dihedral angle violation per constraint (°) | 11.2 |
| Maximum dihedral angle violation (°) | 0.1 |
| R.m.s. deviationsc |         |
| Backbone         | 1.1 (2.0) |
| Heavy atoms      | 1.4 (2.1) |
| Ramachandrand    |         |
| Most favored (%) | 90.7    |
| Additionally allowed (%) | 9.0 |
| Generously allowed (%) | 0.2 |
| Disallowed       | 0.1     |

aTotal assignment completeness, with backbone completeness reported in parentheses
bRestrices 3–142 contain conformational restraining constraints
cResidues 10–142, reported with all-residue RMSDs reported in parentheses
dResidues 10–142, calculated with Procheck

bonds are formed. Overall, among the 20 lowest energy structures calculated, on average 870 Å² are buried for each CARD, comprising ~15% of the total solvent exposed surface area of each CARD.

An interface critical for CARD9 and CARD11 autoinhibition. Amino acids 5–131 of CARD9 share 51% identity with residues 17–143 of CARD11, with no gaps in alignment (Supplementary Fig. 1B). The CARD9 residues homologous to the “LATCH” region required for CARD11 autoinhibition comprise much of the CARD-interacting residues in the linker and coiled-coil (Supplementary Fig. 1F)30. We therefore mapped the known disease-associated or mutagenesis-identified hyperactivating sites in this region of CARD11 onto the CARD92–142 structure25–29. As shown in Figs 2a, b, all oncogenic and disease-associated mutations, as well as 20 of the 23 sites identified in the random mutagenesis screen by Chan et al.30, map to the CARD-coiled-coil interface. This finding suggests that the interface is crucial for autoinhibition of CARD11.

Given the degree of homology with CARD11 and extent of the interface in CARD92–142, we suspected that the CARD-coiled-coil interface acts in an autoinhibitory manner in CARD9 as well. We generated a panel of homologous mutants in full-length CARD9 and transiently transfected them into HEK293 cells containing an NF-κB-inducible reporter gene. As shown in Fig. 2c, overexpression of WT CARD9 weakly induced NF-κB activation as compared to CARD11 lacking its inhibitory domain (CARD11-ΔID), which robustly activated NF-κB, consistent with previous studies8,32. Six of the 11 CARD-coiled-coil-disrupting mutants tested consistently and robustly activated NF-κB by ~4–12-fold, demonstrating that disruption of the CARD-coiled-coil interface is sufficient to activate both CARD9 and CARD11. Four of the mutants tested (C37Y, E81D, L85Y, and Y86F) appear to destabilize CARD9, indicated by significantly lower protein expression; increasing the concentration of transfected plasmid had a minimal effect on protein levels of these mutants. For L85Y, we observed consistently higher NF-κB activation despite the lowered protein expression, confirming that the L85Y mutation is activating. As we were unable to match WT protein expression for C37Y, E81D, and Y86F, we were unable to determine conclusively whether these mutations activate CARD9 as they do CARD11.

A weak CARD11 autoinhibitory interface reinforced by the ID. We reasoned that the increased conformational heterogeneity of CARD118–172, relative to CARD92–152 (Fig. 1b, c), may result from a more labile CARD-coiled-coil interface. Additional interactions with the ID would therefore be required to maintain the closed, autoinhibited state. To test whether the CARD11 CARD-coiled-coil interface is less functionally inhibitory than that of CARD9, we generated a chimeric expression construct in which we replaced the homologous N-terminal residues comprising the CARD, linker, and CARD-interacting stretch of coiled-coil of CARD9 with that of CARD11 (CARD11–ΔID, CARD9WT, schematic shown in Fig. 3a, sequence alignment shown in Supplementary Fig. 1B). Upon overexpression in HEK293 cells, we found that the chimeric CARD11–ΔID/CARD9 construct induced robust NF-κB activation comparable to levels of activation seen in CARD-coiled-coil-disrupting mutants (Fig. 2c) and well above that observed for CARD9WT, even at much higher CARD9WT protein levels (Fig. 3b). These data indicate that the CARD11 CARD-coiled-coil interface is less inhibitory on its own than that of CARD9.

To test whether the function of the CARD11 ID is to maintain a closed CARD-coiled-coil conformation, we generated the complementary chimera in which the N-terminal residues comprising the CARD, linker, and CARD-interacting region of the coiled-coil of CARD11-ΔID were replaced with the homologous residues in CARD9 (CARD9Δ1–131/CARD11-ΔID). CARD11-ΔID induced high levels of NF-κB activation, consistent with previous studies8,32. However, CARD9Δ1–131/CARD11-ΔID,
Disease associated
Disease associated, verified oncogenic
Activating in mutagenesis screen

**Fig. 2** The CARD-coiled-coil interface is critical for maintenance of an autoinhibited state. **a** Disease associated mutants in CARD11 (cyan) and disease associated mutants for which follow-up work confirmed that the mutants were oncogenic (red), mapped onto the homologous residues of CARD9 in the lowest energy NMR structure. **b** Hyperactivating CARD11 mutants identified by Chan et al. in a randomized mutagenesis screen, mapped onto the homologous residues of CARD9. **c** Normalized NF-κB activation in HEK-293 cells upon overexpression of WT or mutant full-length CARD9. The known activator CARD11-ΔID was used as a positive control. Three biologically independent experiments were undertaken, the outcomes of which are shown as dots. The mean value for each construct is depicted as a gray bar. Dotted line represents the value of the WT construct, to which the other values were normalized. Representative western blot is shown for CARD9, demonstrating the total protein expression levels of the constructs. Molecular weights of protein standards are indicated. Source data are provided in Source Data file. **d** The CARD9 CARD-coiled-coil interface, with residues for which mutation consistently activates NF-κB signaling depicted in magenta; sidechain heavy atom van der Walls radii are depicted as dotted surfaces.
which introduces a more robust CARD-coiled-coil interface, induced minimal NF-κB activation (Fig. 3b). To confirm that CARD91–131/CARD11-ΔID retains competency to signal, we also generated a chimeric construct that additionally contained a CARD-coiled-coil-disrupting mutation (CARD91−131/I107E/CARD11-ΔID); as shown in Fig. 3b, this construct activated NF-κB to a level comparable to CARD11-ΔID, indicating that the weak signaling induction by CARD91−131/CARD11-ΔID is due to autoinhibition by the tightly closed CARD-coil-coil interface of CARD9.

CARD9 ubiquitination disrupts the autoinhibitory interface. CARD9 has no known region analogous to the ID, and based on the weak NF-κB activation induced by the CARD91−131/CARD11-ΔID chimeric construct (Fig. 3b), the CARD9 CARD-coiled-coil interface is sufficiently tight alone to maintain the CARD in an inhibited state. These observations suggest that signaling-induced disruption of the autoinhibitory CARD-coiled-coil interface must proceed through a different mechanism for CARD9 than for CARD11. Cao et al.12 reported that CARD9 is specifically ubiquitinated at Lys125 by TRIM62 and that this ubiquitination is required for signal propagation through CARD9. The dimeric CARD92−142 structure shows that Lys125 is not directly involved in the CARD-coiled-coil interface; however, Lys125 is in proximity to the CARD, suggesting that ubiquitin conjugation at Lys125 could sterically disrupt the CARD-coiled-coil interface.

To generate sufficient quantities of ubiquitinated CARD92−152, we utilized a chemical ligation approach. A CARD92−152 construct was generated with both cysteines (C10 and C37) mutated to serine, and Lys125 mutated to cysteine (CARD92−152/SSC). In parallel, a ubiquitin variant was generated for which the C-terminal glycine was mutated to cysteine (UbG76C). After individually purifying each component, the two proteins were conjugated using the short bi-functional maleimide crosslinker bismaleimidoethane (BMOE) and purified by size exclusion chromatography, yielding CARD92−152/SSC dimers with ~95% ubiquitin conjugation (Supplementary Fig. 2A). As CARD92−152/SSC and UbG76C were purified separately, we generated ligated preparations for which only CARD92−152/SSC or UbG76C was 15N-labeled, allowing selective investigation of the impact of ligation on each of the proteins by NMR.

Fig. 3 The CARD11 CARD-coiled-coil interface is weaker than that of CARD9, but is reinforced by the ID. a Cartoon schematic of the domain architecture of CARD9, CARD11-ΔID, and CARD11 along with the chimeric constructs CARD91–131/CARD11-ΔID and CARD11–143/CARD9, wherein the N-terminal residues comprising the CARD, linker, and CARD-interacting stretch of the coil-coil are swapped between the two proteins (See Supplementary Fig. 1B). b Normalized NF-κB activation in HEK-293 cells upon overexpression of the indicated constructs. Three biologically independent experiments were undertaken, the outcomes of which are shown as dots. The mean value for each construct is depicted as a gray bar. Dotted line represents the value of the CARD9WT construct, to which the other values were normalized. Representative western blot is shown for CARD9 and CARD11 constructs, using antibodies specific for regions outside of the N-terminal swapped domain or ID. Source data are provided in Source Data file.
The overall peak dispersion and linewidths remain comparable between CARD9\(^{2-152/SSC}\) and CARD9\(^{2-152}\) (Supplementary Fig. 2B), indicating that the mutations have not disrupted the CARD-coiled-coil interface, confirming that Lys125 is not crucial for maintenance of the interface, and allowing us to assign 79% of the backbone amide chemical shifts in CARD9\(^{2-152/SSC}\). Mixing 14\(N\)-Ub\(^76\)C and 15\(N\)-CARD9\(^{2-152/SSC}\) in a 1:1 ratio, we observed no changes in the CARD9\(^{2-152/SSC}\) spectrum, indicating that, in the absence of conjugation, the two proteins do not appreciably interact (Supplementary Fig. 2C). However, as shown in Fig. 4a, the BMOE-conjugated 15\(N\)-CARD9\(^{2-152/SSC}\)-14\(N\)-Ub\(^76\)C spectrum exhibited both a global line broadening, attributable to the larger molecular weight of the complex (52.1 kDa vs. 34.6 kDa), as well as site-specific, selective line broadening indicative of intermediate time-scale dynamics for certain regions of CARD9\(^{2-152/SSC}\) (Fig. 4b). Mapping all residues with a >75% reduction in peak intensity onto the CARD9\(^{2-152/SSC}\) apo structure (PDBID 1UBQ [http://dx.doi.org/10.2210/pdb1UBQ/pdb], right). Selectively line-broadened residues (<25% reduction in peak intensity for Ub\(^76\)C and >10% reduction in peak intensity for ubiquitin) are colored blue. Residues Lys125 in CARD9\(^{2-152/SSC}\) and Gly76 in ubiquitin are colored cyan. All source data are provided in Source Data file.

Fig. 4 Ubiquitination of CARD9 at Lys125 disrupts the CARD-coiled-coil interface. a 15\(N\)-TROSYs of CARD9\(^{2-152/SSC}\) apo (black) or conjugated to 14\(N\)-Ub\(^76\)C (red). b Peak intensities of spectra in a for all assigned, non-overlapped peaks. c Lowest energy CARD9\(^{2-152}\) dimer structure (left) and ubiquitin crystal structure (PDBID 1UBQ [http://dx.doi.org/10.2210/pdb1UBQ/pdb], right). Selectively line-broadened residues (<25% apo intensity for CARD9\(^{2-152}\) and <5% apo intensity for ubiquitin) are colored red. Minimally line-broadened residues (>50% apo intensity for CARD9\(^{2-152}\) and >10% apo intensity for ubiquitin) are colored blue. Residues Lys125 in CARD9\(^{2-152}\) and Gly76 in ubiquitin are colored cyan. d 15\(N\)-TROSYs of CARD9\(^{2-152/WT}\) (black) and CARD9\(^{2-152/I107E}\) (red). e 15\(N\)-TROSYs of Ub\(^76\)C apo (black) or conjugated to 14\(N\)-CARD9\(^{2-152/SSC}\) (red). f Peak intensities of spectra in panel e for all non-overlapped peaks. g Deuterium exchange as a function of labeling time for representative peptides from CARD9\(^{2-152/SSC}\) and Ub\(^76\)C un conjugated (black) or conjugated (red). h Global protection factors for CARD9\(^{2-152/SSC}\) and Ub\(^76\)C after conjugation as compared to apo. Each dot represents the mid-point of a single identified peptide. Dotted line indicates a protection factor of 1, indicating no change upon conjugation. Values less than one indicate de-protection, i.e., faster deuterium exchange. i CARD9\(^{2-152}\) dimer structure with regions de-protected upon ubiquitination plotted in red and unaffected regions plotted in blue. Lys125 is indicated in cyan. All source data are provided in Source Data file.
are distal from the interface. We compared the $^{15}$N-CARD9$^{2-152}$/SSC–$^{14}$N-UbG76C spectrum to those of the constructs containing the activating mutations I107E (CARD9$^{2-152}$/I107E) or L115I (CARD9$^{2-152}$/L115I) and found that the activating mutants induce selective line broadening to a similar extent and to nearly identical residues (Fig. 4d and Supplementary Fig. 2D), suggesting that this pattern of line broadening represents a signature of functionally significant CARD-coiled-coil disruption.

The inversely labeled $^{15}$N-CARD9$^{2-152}$/SSC–$^{15}$N-UbG76C complex likewise exhibited both global and selective line broadening as compared to $^{15}$N-UbG76C– alone (Fig. 4e, f). The degree of global line broadening is more severe in this case due to the much larger molecular weight difference (52.1 kDa vs. 8.8 kDa). Highlighting those residues with the greatest peak intensity reduction (>95%, Fig. 4c, red) onto a ubiquitin crystal structure, the line broadened residues map predominantly to a well characterized hydrophobic patch that includes L8, I44, and V70—the most common protein–protein interaction interface on ubiquitin18.

The selective line broadening in both CARD9 and ubiquitin suggests that the hydrophobic patch of ubiquitin competes with the CARD to interact with the hydrophobic surface of the coiled-coil and/or linker region and thus displaces the CARD. To test whether the CARD is physically displaced by ubiquitin conjugation, we monitored conjugated and unconjugated CARD9$^{2-152}$/SSC and UbG76C by time-resolved mass spectrometry-based hydrogen-deuterium exchange (HDX). By comparing deuterium incorporation over time between a conjugated and unconjugated sample, a protection factor can be assigned to each peptide identified in both CARD9$^{2-152}$/SSC and UbG76C (representative examples shown in Fig. 4g, see Source Data for all identified peptides); protection factors of less than one correspond to de-protection (i.e., faster exchange, structural destabilization) upon ubiquitination, while protection factors of greater than one correspond to protection. Globally, we identified several regions of CARD9$^{2-152}$/SSC that exhibit de-protection upon ubiquitination (Fig. 4h), consistent with the displacement of the CARD and elimination of structurally stabilizing interactions. By comparison of many overlapping peptides, we found that residues 24–38, 71–78, 83–85, 93–102, 106–108, and 116–120 were de-protected upon ubiquitination, while residues 4–23, 39–66, and 129–135 remained unaffected. Mapping these residues onto the structure, we found increases in exchange rates along the CARD-coiled-coil interface upon ubiquitination, while residues distal from the interface remained unaffected (Fig. 4i), corroborating our interpretation of the NMR data that Lys125 ubiquitination physically displaced the CARD. UbG76C exhibited a modest global protection upon conjugation, consistent with a stabilizing interaction between the conjugated ubiquitin and CARD9$^{2-152}$/SSC. Finally, we compared HDX rates between CARD9$^{2-152}$/WT and a construct containing an activating mutation, CARD9$^{2-152}$/I107E. At both the individual peptide level (Supplementary Fig. 2E) and globally (Supplementary Fig. 2F), ubiquitination of CARD9$^{2-152}$/SSC induced a similar pattern and extent of de-protection as the I107E mutation, indicating that ubiquitination at Lys125 displaces the CARD to a functionally significant degree.

CARD9 CARD-coiled-coil disruption promotes helical assembly. Canonically, death domains like CARDs propagate signaling cascades via homo- or heterotypic interactions with other death domains, forming helical assemblies34. For CARD9 and CARD11, the CARD–CARD interaction with Bcl10 is critical for signal propagation, and CARD11$^{8-172}$ has been shown to accelerate polymerization of Bcl10 filaments in vitro15. We hypothesized that the CARD-coiled-coil interaction sequesters the CARD9 and CARD11 CARDs in the absence of activation, thus preventing homotypic interactions required to generate a Bcl10-nucleating template.

To test this hypothesis, we purified recombinantly expressed CARD9$^{2-152}$ with mutations identified as NF-kB activating (Fig. 2b). We found that CARD9$^{2-152}$/I107E spontaneously formed filaments after a short incubation at elevated temperatures, which are readily visible by negative-stain electron microscopy (NS-EM, Fig. 5a). As we have shown previously for the CARD9 CARD alone, filament formation was hindered in the presence of zinc. Upon addition of excess EDTA to 0.5 mM CARD9$^{2-152}$/I107E that had been purified with a 1:1 concentration of zinc, filaments were readily observable by NS-EM after 10 min at 25 °C, while none were observed without adding EDTA (Supplementary Fig. 3A). In the presence of zinc, filaments could be induced by elevating the temperature to 37 °C, suggesting that zinc inhibits but does not completely block filament formation. A second mutant, CARD9$^{2-152}$/L115I, was additionally identified to form filaments in vitro that are similarly modulated by Zn$^{2+}$. The I115L mutation appears to be somewhat less permissive of filament formation, however, as filaments formed within 10 min upon Zn$^{2+}$ removal only at 37 °C, and not at 25 °C. Under none of these conditions were filaments of CARD9$^{2-152}$/WT ever found, even after removal of Zn$^{2+}$ from 1 mM CARD9$^{2-152}$/WT followed by incubation for 16 h at 37 °C (Fig. 5a).

To test whether the CARD-coiled-coil interaction blocks CARD9 CARD assembly in a cellular context, we utilized Distributed Amphiphilic FRET (DAmFRET)35. Briefly, in this method, a protein of interest is tagged with the photoconvertible fluorescent protein mEos3.1 and expressed over a range of concentrations in cells. By exposing the cells to a limiting dose of 405 nm light, a fraction of the fluorophore molecules are photoconverted, allowing them to act as FRET acceptors to the fluorophore molecules that have not converted. This system thus allows for monitoring protein self-assembly, which increases FRET signal, as a function of the protein’s concentration in cells. The DAmFRET assay has been established in the budding yeast, S. cerevisiae, which has the advantage of not containing native death domain proteins or associated regulatory mechanisms that could potentially obscure intrinsic CARD assembly properties. For the CARD9 CARD alone (CARD1$^{97}$), DAmFRET revealed a sharp increase in polymerization at concentrations above ~100 μM (Fig. 5b and Supplementary Fig. 3B, C), consistent with previous in vitro measurements31. Inclusion of the coiled-coil (CARD9$^{1-142}$) blocked this polymerization while simultaneously increasing the FRET level of soluble protein (Fig. 5b, inset), consistent with the expected formation of dimers. Disrupting the CARD-coiled-coil interface within this construct via mutation (CARD9$^{1-142}$/I107E) restored polymerization at high concentrations (Fig. 5b and Supplementary Fig. 3B, C). Intriguingly, the mutation also increased FRET levels for the soluble protein (Fig. 5b, inset), suggesting an increase in oligomerization even at subsaturating concentrations. The CARD9 CARD-coiled-coil interaction thus inhibits CARD9 CARD polymerization in living cells, as well as in vitro.

CARD9 templates Bcl10 nucleation in vitro and in cells. We anticipated that the observed in vitro filaments (Fig. 5a) likely represent the nucleating template that seeds Bcl10-filament formation. To test this hypothesis, we utilized a fluorescence polarization (FP)-based Bcl10 polymerization assay described previously31, based on the assay developed by Qiao et al.13. Bcl10 was purified with an N-terminal MBP tag linked by a TEV protease cleavage site and sparsely labeled with a fluorophore. The MBP tag blocks Bcl10 polymerization; however, the MBP tag
is rapidly removed upon addition of TEV protease, with >50% cleaved in 2 min (Supplementary Fig. 3D). Upon removal of MBP, Bcl10 spontaneously forms filaments, leading to an increase in FP. Factors that nucleate Bcl10 induce a more rapid increase in FP upon addition of the protease.

Utilizing this FP assay, we found that CARD92–152/I107E and CARD92–152/L115I filaments accelerate Bcl10 polymerization, while CARD92–152/WT, or mutant constructs under conditions that do not generate filaments, fail to accelerate Bcl10 polymerization. Specifically, we added excess EDTA to 1:1 Zn$^{2+}$-complexed CARD92–152/I107E and incubated it at either a concentration that allows (0.5 mM) or does not allow (4 μM) filament formation. Adding an equimolar concentration of the filamentous preparation to 2 μM MBP-Bcl10, but not the non-filamentous preparation, accelerated Bcl10 polymerization (Fig. 5c). As shown in Supplementary Fig. 3E, Zn$^{2+}$-bound CARD92–152/I107E is also capable of promoting Bcl10 polymerization, but only when first incubated under conditions permissive of filaments. Under no conditions was the CARD92–152/WT construct able to accelerate Bcl10 polymerization. CARD92–152/L115I is likewise able to promote Bcl10 polymerization only under conditions for which it forms...
CARD-coiled-coil disruption promotes Bcl10-nucleating CARD9 helical assembly. a NS-EM micrographs of CARD9 

![CARD9-2-152/WT, CARD9-2-152/L110I, and CARD9-2-152/I115I](image), after removal of Zn²⁺ by EDTA and incubation at the indicated concentration, time, and temperature. Incubation of CARD9-2-152/WT for shorter times, at lower temperature (25 °C) or lower concentration (0.5 mM) also failed to produce filaments by NS-EM. Scale bars of all panels are 200 nm. b DAmFRET assay performed on CARD9-1-97, CARD9-1-142/WT, or CARD9-1-142/L110I. Inset depicts AmFRET binned at 50-60 μM for each construct, with 50 μM indicated by vertical dashed lines. See Supplementary Fig. 4D for replicate data, statistical analysis, and data binned at 100-200 μM. c Bcl10 FP nucleation assay. MBP-Bcl10 was mixed with CARD9-2-152 preparations as indicated. EDTA addition and incubation of CARD9-2-152 constructs occurred prior to initiation of nucleation assay; 11 and 5:1 refers to the molar ratio during the FP assay, with the final concentration of Bcl10 always set to 2 μM. TEV protease was added at t = 0 to indicated samples. Vertical bars represent the s.d. of three technical replicates. The data shown in the two plots were collected in the same experiment and were plotted separately to aid in visualization; hence, the "CARD9-1-97" and "CARD9-1-142/WT" data are identical in both plots. Source data are provided in Source Data file. d Bcl10, CARD9-2-152/L110I, or CARD9-2-152/I115I filaments, 2 min after addition of TEV protease and with addition of anti-HA antibody during NS-EM grid preparation. e, f Direct templating of Bcl10 by CARD9-2-152/L110I (e) or CARD9-2-152/I115I (f) filaments. Grids were prepared 2 min after addition of TEV protease and with anti-HA antibody added during NS-EM grid preparation. White arrows indicate CARD9-2-152-to-Bcl10-filament transitions. For panels d-f, samples were prepared under identical conditions and concentrations as in e with all grids prepared 2 min after addition of TEV protease. g Genetic correlation of Bcl10-mEos3.1 alone (left), showing that it partitions cells into overlapping populations that contain either monomer alone or polymer; or for Bcl10-mEos3.1 with co-expressed with μNS-Bcl10 (right), which shifts all cells to the polymer-containing population. h DAmFRET assay for Bcl10-mEos3.1 with co-expressed CARD9-1-97, CARD9-1-142/WT, or CARD9-1-142/L110I either alone or tethered to μNS.

CARD9 helical assembly, and not the dimeric form of the protein, is capable of accelerating Bcl10 polymerization and that disruption of the CARD-coiled-coil interface promotes this activity. We have previously shown that CARD9-2-97 helical assemblies are capable of directly templating Bcl10 polymerization. To demonstrate that the CARD9-2-152/L110I and CARD9-2-152/I115I helical assemblies are likewise accelerating Bcl10 polymerization through direct templating, we visualized, by NS-EM, CARD9-2-152 filament-nucleated Bcl10 filaments shortly after addition of TEV. To distinguish CARD9-2-152 and Bcl10 filaments, we utilized a Bcl10 construct with a C-terminal HA tag and added anti-HA antibody after incubation with CARD9-2-152 preparations as indicated. EDTA addition and incubation of CARD9-2-152 constructs occurred prior to initiation of nucleation assay; 11 and 5:1 refers to the molar ratio during the FP assay, with the final concentration of Bcl10 always set to 2 μM. TEV protease was added at t = 0 to indicated samples. Vertical bars represent the s.d. of three technical replicates. The data shown in the two plots were collected in the same experiment and were plotted separately to aid in visualization; hence, the "CARD9-1-97" and "CARD9-1-142/WT" data are identical in both plots. Source data are provided in Source Data file. d Bcl10, CARD9-2-152/L110I, or CARD9-2-152/I115I filaments, 2 min after addition of TEV protease and with addition of anti-HA antibody during NS-EM grid preparation. e, f Direct templating of Bcl10 by CARD9-2-152/L110I (e) or CARD9-2-152/I115I (f) filaments. Grids were prepared 2 min after addition of TEV protease and with anti-HA antibody added during NS-EM grid preparation. White arrows indicate CARD9-2-152-to-Bcl10-filament transitions. For panels d-f, samples were prepared under identical conditions and concentrations as in e with all grids prepared 2 min after addition of TEV protease. g Genetic correlation of Bcl10-mEos3.1 alone (left), showing that it partitions cells into overlapping populations that contain either monomer alone or polymer; or for Bcl10-mEos3.1 with co-expressed with μNS-Bcl10 (right), which shifts all cells to the polymer-containing population. h DAmFRET assay for Bcl10-mEos3.1 with co-expressed CARD9-1-97, CARD9-1-142/WT, or CARD9-1-142/L110I either alone or tethered to μNS.

CARD11-18-172 also forms Bcl10-nucleating helical templates. Consistent with a more labile CARD-coiled-coil interface in CARD11 (Figs. 1b and 3), CARD11-8-172 was able to polymerize in vitro without introduction of a mutation. When incubated at 37 °C, solutions of 1 mM CARD11-8-172, but not 4 μM CARD11-8-172, were observed to contain filaments visualized by NS-EM (Supplementary Fig. 4A). The filament-containing sample of CARD11-8-172 strongly accelerated Bcl10 polymerization, whereas the sample that lacked filaments had no effect (Supplementary Fig. 4B). Although CARD9-2-152 filament-containing samples formed intractable gels after extended incubation, CARD11-8-172 filament-containing samples instead remained clear. We thus utilized SEC to fractionate the filament-containing sample, yielding three distinct species: dimers, filaments, and fairly uniform oligomers of ~300 kDa (Supplementary Fig. 4C, D). Separation was incomplete, however, as small numbers of large filaments could be identified by NS-EM in the dimer and oligomer fractions (Supplementary Fig. 4E); presumably these filaments were retained by the in-line pre-column filter and subsequently dislodged over the course of the SEC run. At a 1:5 molar ratio, the filament fraction was capable of robust Bcl10 nucleation, while the dimer and oligomeric fractions inhibited nucleation only slightly above background (Supplementary Fig. 4F), indicating that, as with CARD9, the CARD11 helical assembly is required for Bcl10 nucleation. Given their inability to nucleate and their uniform size, we suspect the observed CARD11-8-172 oligomers represent an "off-pathway", non-helical conformational state, perhaps mediated by domain-swapping as was observed for CARD9-2-152 (Supplementary Fig. 1C). Utilizing an anti-HA antibody to decorate the Bcl10 filaments, we were also able to identify continuous CARD11-8-172-Bcl10 filaments (Supplementary Fig. 4G), demonstrating direct templating as the mechanism of enhanced Bcl10 polymerization.
The structural basis for polymerization of CARD9. As previously reported\textsuperscript{31}, CARD9\textsuperscript{2–97} filaments present almost exclusively as large bundles when frozen for Cryo-EM. With the CARD9\textsuperscript{2–152/1107E} construct, however, we found that the filaments now remained predominantly singular when frozen shortly after inducing filament formation. With this sample, we were able to generate a 4.0-Å reconstruction of the CARD9 CARD filament (Fig. 6a and Supplementary Fig. 5). The coiled-coil domain is disordered relative to the CARD core, such that the reconstruction contains only CARD residues 8–95. The overall resolution was estimated at 4.0-Å by Fourier shell correlation (FSC, Supplementary Fig. 5B), with regions towards the center of each CARD estimated at ~3.5 Å and the N-terminal region and flexible α3-α4 loop estimated at ~4.5 Å (Table 2 and Supplementary Fig. 5C). The map was of sufficient quality to resolve most sidechains (Supplementary Fig. 5E), allowing us to model the CARD and define the interfaces involved in the assembly. The CARD adopts the canonical helical architecture of a death domain, with helical symmetry of a 5.11 Å rise and 101.6° left-handed rotation, consistent with a growing body of literature demonstrating a narrow range of parameters adopted by helical assemblies of CARDs (4.85–5.13 Å rise and 100.2°–101.6° left-handed rotation) including that of Bcl10\textsuperscript{36,38–41}. As is typical of this architecture, the CARD utilizes three CARD-CARD interfaces, known as the Type I, Type II, and Type III interfaces, corresponding to the 3-, 4-, and 1-start helical symmetries, respectively (Fig. 6a, b). As detailed in Fig. 6c, interactions within the interfaces are predominantly electrostatic in nature, consistent with our previous observation that increasing salt concentration can inhibit assembly\textsuperscript{31}.

To confirm the importance of the identified interfaces in CARD helical assembly, we generated a series of mutants in each interface and tested their impact on the CARD filament formation. As we have previously demonstrated, upon removal of Zn\textsuperscript{2+} from 150 μM 1:1 Zn\textsuperscript{2+}-bound CARD9\textsuperscript{2–97/WT} by addition of EDTA, the CARD assembles into filaments. Density is only present for the CARD core of the assembly. Seven CARDs of the modeled structure are shown, depicting the six CARDs contacted by any one CARD, colored as in Supplementary Fig. 6A, B). We also tested a clinically identified mutant (R35Q), which was found in a patient with invasive fungal infections in the central nervous system and digestive tract\textsuperscript{18}. As with the other interface mutants, R35Q completely
blocked CARD filament formation, providing a molecular explanation for the clinically observed deficiency in CARD9 signaling. Within the Type III interface, D43 and K58 form a relatively isolated salt bridge. By swapping the charges on these two residues (CARD9_{2-97/D43K58}), the CARD regains filament-forming capacity, albeit at a higher protein concentration, confirming the importance of this salt-bridging interaction in the formation of the CARD assembly.

Comparing the Cryo-EM structure of CARD9_{2-152/1107E} to a previously determined NMR structure of CARD9_{2-97} (PDB ID 6E26), most of the CARD remains minimally changed between the monomeric and polymerized form (Fig. 6f). The exception is the flexible a3-a4 loop, which adopts a "closed" conformation in the NMR structure, but an "open" conformation in the filament, stabilized by interactions across the Type III interface. The Zn\(^{2+}\)-binding site, comprising critical residues Cys10 and His73, is also minimally perturbed upon filament formation (Supplementary Fig. 6C), failing to resolve the outstanding question as to how Zn\(^{2+}\) binding modulates CARD9-CARD filament assembly.

Given the structural similarity of the monomeric CARD9 and CARD11 CARDs\(^{31}\), we anticipated that the CARD11 CARD (CARD11_{8-109}) forms a helical assembly in a similar manner. To probe the CARD11 CARD helical assembly, we used Rosetta to generate a model of the CARD11_{8-109} filament based on our EM structure of the CARD9 CARD filament. As shown in Supplementary Fig. 7A, our model predicts utilization of the same three canonical interfaces; while the residues involved in the interactions are not strictly conserved with those of CARD9, the interaction remains predominantly electrostatic. To experimentally validate our model, we generated mutations within each of the interfaces at residues involved in putative CARD-CARD interactions. Unlike the WT CARD11 CARD, which forms filaments in vitro at 250 \(\mu\)M, each of the mutants were incapable of filament formation at double that concentration (Supplementary Fig. 7B), confirming their relevance in mediating the CARD11 CARD helical assembly.

A number of CARD domains, including those of Caspase-1 and Bcl10, have been demonstrated to polymerize unidirectionally\(^{36,39}\), implying that only one end of a growing filament is structurally compatible with recruiting additional monomers. To test whether CARD9 CARDs likewise exhibit unidirectional growth, we induced CARD9_{2-97/WT} filament formation in the presence of an equimolar concentration of interface mutants, demonstrated in Fig. 6d to be unable to form filaments. If the "b" interface tip (top as oriented in Fig. 6a) of the filament can recruit additional monomers, mutants in the "a" interfaces would be unable to effectively bind growing CARD9_{2-97/WT} filaments, while mutants in the "b" interface would be able to bind and "cap" growing CARD9_{2-97/WT} filaments, inhibiting further growth (See Supplementary Fig. 6D for schematic cartoon). We found that the three "a"-interface mutants tested minimally impacted CARD9_{2-97/WT} filament growth, while the three "b"-interface mutants significantly impaired CARD9_{2-97/WT} filament growth, demonstrating that the CARD9 CARD polymerizes unidirectionally, with the "b" interface at the leading end. The directionality of this growth is consistent with the directionality previously suggested for Bcl10.

### Discussion

The existence of an autoinhibited state of CARD11 was first indicated by Sommer et al.\(^8\), who demonstrated that phosphorylation of the ID relieves apparent autoinhibition of the protein. The biological and pathological importance of autoinhibition was reinforced in a series of studies demonstrating that oncogenic, constitutively activating mutations functionally disrupt this repression\(^{30,32,42}\). However, the specific mechanism by which CARD11 and the rest of the protein family are held in an autoinhibited state prior to activation has remained an open question.

Here, we have determined the structure of an autoinhibited CARD9 construct, demonstrating that the CARD forms an extensive, predominantly hydrophobic, interaction with the adjacent linker, and coiled-coil elements (Fig. 1). The residues involved in this interaction are largely conserved, both between CARD9 and CARD11 and among the four-member family that also includes CARD10 (CARMA3) and CARD14 (CARMA2) (Supplementary Fig. 1G), suggesting that the nature of the autoinhibited state is conserved throughout the family. Further in support of this notion, nearly all of the activating mutations previously identified in CARD11 map to this interface, and in particular to "inward-facing" residues; for example, in the linker, the only residues not identified as activating are the solution-exposed residues in the short helical segment (Fig. 2b).

With insight provided by this structural understanding of the autoinhibited state, we demonstrated that the CARD11 interface is more labile than that of CARD9, but is reinforced by the ID. Replacing the CARD and CARD-interacting region of CARD11 with the more stable region of CARD9 (Fig. 3b) revealed that the ID is functionally redundant in maintaining autoinhibition in the context of a robust CARD-coiled-coil interaction. Jattani et al.\(^9\) recently demonstrated that the CARD11 ID comprises numerous redundant sections that interact in a multivalent manner with the CARD and the larger coiled-coil domain, explaining why activating mutations within the ID itself have not been identified\(^{9,43}\). Following similar logic, we anticipate that the multivalent nature of the ID interaction precludes single mutations in the CARD or coiled-coil domain from functionally disrupting all interactions.

### Table 2 Structural statistics for the CARD9 CARD helical assembly

| Magnification | 130,000 |
| Voltage (kV) | 300 |
| Electron exposure (e-/Å\(^2\)) | 50.9 |
| Defocus range (µm) | 1.25-2.5 |
| Pixel size (Å) | 1.085 |
| Symmetry imposed | Helical: 5.11 Å, −101.6° |
| Initial particle images (no.) | 141,592 |
| Final particle images (no.) | 31,908 |
| Map resolution (Å) | 4.0 |
| FSC threshold | 0.143 |
| Map resolution range (Å) | 3.5-4.5 |
| Initial model used (PDB code) | 6E26 |
| Model resolution (Å) | 4.0 |
| FSC threshold | 0.143 |
| Map sharpening B factor (Å\(^2\)) | 90 |
| Validation MolProbity score | 2.03 |
| Clashscore | 7.4 |
| Poor rotamers (%) | 0.0 |
| Ramachandran plot
| Favored (%) | 87.2 |
| Allowed (%) | 13.8 |
| Disallowed (%) | 0.0 |
with the ID; yet, disruptions to the CARD-coiled-coil interface that promote helical assembly of the CARD may block interactions with the ID, thus explaining the reduced ID-CARD co-immunoprecipitation in the presence of CARD-coiled-coil disrupting mutants previously reported. Further investigation of the CARD11 ID, CARD, and coiled-coil will be required to detail the specific structural determinants of this complex, multivalent interaction and how upstream signaling events disrupt this repressed state to fully activate CARD11.

For CARD9, activation is mediated by ubiquitination at Lys125, which displaces the CARD from the coiled-coil to a comparable extent as an activating mutation (Fig. 4). Cao et al. showed that the ubiquitin ligase TRIM62 activates CARD9 via a specific ubiquitination at Lys125 and that, upon TRIM62 overexpression, CARD9 is heavily modified via K27-linked ubiquitin, but did not investigate whether poly-ubiquitination is required for CARD9 activation. Our studies (Fig. 4 and Supplementary Fig. 2) suggest that conjugation of a single ubiquitin to Lys125 is sufficient to disrupt the CARD-coiled-coil interface in a similar manner and to a similar degree as mutations that are activating in cells. We anticipate that poly-ubiquitination could function to promote more robust signaling through CARD9 by providing additional hydrophobic ubiquitin surfaces to compete with the CARD for coiled-coil binding and/or slowing inactivation by deubiquitinases.

Another activating post translational modification described for CARD9, phosphorylation at Thr231, occurs outside the region of the protein structurally characterized here, further into the larger coiled-coil domain. Given that disruption of the CARD-coiled-coil interface is sufficient alone to activate CARD9 upon overexpression (Fig. 2), we suggest that the T231 site may play a role in modulating coiled-coil interactions. In particular, we note that the polypeptide nature of the multiple coiled-coil-forming segments in CARD9 (Fig. 1a) may allow the protein to partition into phase-separated condensates that would be expected to promote the formation of nucleating seeds by increasing the local concentration of the CARD. In support of this theory, Qiao et al. found that a CARD11 construct comprising this region (CARD11–302) purifies as a large multimer capable of nucleating Bcl10 polymerization. Furthermore, a second “hot-spot” of CARD11 oncogenic mutations exists near this site, including the homologous residue, Ser250, suggesting it as a conserved structural determinant of this complex, multivalent CARD9 activation. We do, however, note a discrepancy in our findings as a large multimer capable of nucleating Bcl10 polymerization. We do, however, note a discrepancy in our findings as a large multimer capable of nucleating Bcl10 polymerization. However, addition of TEV protease overnight to remove the 6xHis tag did not require higher-order multimerization to nucleate Bcl10 polymerization, while our observations for both CARD9–152 and CARD11–182 suggest that the proteins are capable of seeding Bcl10 nucleation only after assembling into a helical template (Fig. 5 and Supplementary Fig. 4). In our studies, the CARD11–182 dimer is capable of forming Bcl10-nucleating filaments upon concentration and incubation, suggesting that Qiao et al. may have handled the CARD11–182 protein in such a way as to inadvertently allow some nucleating multimers to form subsequent to SEC purification.

The structures of the CARD91–142 dimer and the CARD9 CARD helical assembly we present here provide insight into both the autoinhibited and activated forms of this family of Bcl10-nucleating proteins. In addition to providing a structural explanation for pathogenic mutations in both CARD9 and CARD11, these structures allowed us to define the mechanisms of activation for both proteins in the course of canonical signaling. While significant questions remain, in particular in understanding the regulatory role of the larger coiled-coil domain, our study provides a number of critical steps towards a full structural description of regulation within the protein family.

**Methods**

**Protein purification.** All purified CARD9, CARD11, and ubiquitin constructs comprised the human sequences and were expressed in BL21(DE3) or with an N-terminal TEV-cleavable 6xHis tag. Protein production was achieved by growth for 48–72 h at 16 °C in Terrific Broth autoinduction media or 15N autoinduction media for unlabeled and 15N-labeled protein, respectively. Uniform 15N-labeled and 15N-13C-labeled protein was generated by induction in 13C15N or 13C15N minimal media with 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) for 6 h at 37 °C. U176G was purified by lysing cell pellets in nickel buffer (50 mM HEPES, 500 mM NaCl, 20 mM imidazole, pH 7.5) via sonication. After a high-speed spin, the soluble fraction was passed over a Ni-NTA (Qiagen) affinity column, washed with nickel buffer, and eluted with nickel buffer supplemented with 400 mM imidazole. TEV protease was added to this solution, which was then dialyzed overnight to into nickel buffer at 30 °C to cleave the 6xHis tag. A majority of the uncleaved U176G was removed by an additional Ni-NTA column, however a small amount of uncleaved U176G nonetheless remained (see Supplementary Fig. 2A). U176G was finally purified over a Superdex 75 gel filtration column (GE Healthcare).

A significant percentage of CARD11–172, CARD9–152 (WT and mutant constructs), and CARD9–142 remained in the insoluble fraction upon gentle lysis; these constructs were thus purified under denaturing conditions. Cell pellets were lysed in 5 M guanidine hydrochloride, 100 mM HEPES, 100 mM NaCl, pH 7.5. After high-speed spin, the soluble fraction was passed over a Ni-NTA affinity column, washed with urea wash buffer (6.5 M Urea, 50 mM HEPES, 50 mM NaCl, 20 mM imidazole, pH 7.5), and eluted with urea wash buffer supplemented with 400 mM imidazole. Five millimolar TCEP was added to eluate, which was then dialyzed twice against nickel buffer with 0.25 mM TCEP to allow for refolding, followed by addition of TEV protease overnight to remove the 6xHis tag. Uncleaved protein and TEV protease were subsequently removed by Ni-NTA. Finally, all proteins were purified via a Superdex 200 gel filtration column (GE Healthcare). For formulations containing zinc, a saturating concentration of ZnCl2 was added prior to the final gel filtration column. WT and most mutants of CARD9–142 and CARD11–182 were purified as U176G, except that TEV protease cleavage was achieved through overnight incubation at 4 °C and saturating ZnCl2 concentrations were added to CARD9–97 prior to the final Superdex 75 gel filtration column. CARD9–97 mutants that were predominantly insoluble upon gentle lysis were purified under denaturing conditions, as described for CARD9–142 except that the final gel filtration column was a Superdex 75.

The MBP-Bcl10 construct comprises an N-terminal MBP, followed by a TEV cleavage site, human Bcl10, with mutations C10A and C29A to prevent fluorophore labeling of the Bcl10 CARD, and a C-terminal HA-tag (YDYDDPPYDYA). The protein was expressed in BL21(DE3) cells grown in LB media at 37 °C to an optical density at 600 nm of 0.7. 0.2 mM IPTG was added to induce protein production for one hour at 37 °C. MBP-Bcl10 pellets were lysed in nickel buffer via sonication and spun down. The soluble fraction was bound to Ni-NTA (Qiagen) affinity column, washed with nickel buffer, and eluted in 5 ml 150 mM NaCl, 20 mM imidazole supplemented with 400 mM imidazole. Two micromolar Alexa Fluor 488 C5 maleimide (ThermoFisher) was added to the eluate, which was incubated for 10 min at room temperature. The eluate was then purified over a Superdex 200 gel filtration column. The monomeric MBP-Bcl10 peak was collected, stored at 4 °C, and utilized for FP assays or microscopy within 2 h of elution from the gel filtration column with no concentration during any step of the purification.

For all purifications, the subsequent downstream application dictated the final gel filtration buffers, which are described for each application in the following sections.

**NMR data collection.** All NMR spectra shown were collected in 50 mM HEPES, 300 mM NaCl, 0.5 mM TCEP, pH 7.0 at 37 °C on a Bruker 800 MHz Bruker Spectrometer with a cryogenically cooled probe. Backbone assignments and structure determination utilized a construct comprising residues 1–152 of human CARD9. Assignments were performed on 1 mM samples with 1:1 15N–13C double labeling, with the exception of the CARD9–97 spectrum in Fig. 1c, which was a 15N-HQCC. The CARD11–172 spectrum in Fig. 1b was collected at 150 μM, while the CARD9–152 and CARD9–97 spectra depicted in Fig. 1c were collected at 200 and 400 μM, respectively. The spectra depicted in Fig. 4 and Supplementary Fig. 2 were all collected at 200 μM using identical experimental parameters.

**NMR assignments and solution structure determination.** CARD9–142 assignments and structure determination utilized a construct comprising residues 2–142 of human CARD9. Assignments were performed on 1 mM samples with 1:1 ZnCl2; in 50 mM HEPES, 300 mM NaCl, 0.5 mM TCEP, pH 7.0, 10% D2O, at 37 °C, with data collected on a 600 or 800 MHz Bruker Spectrometer with a cryogenically cooled probe. Backbone assignments were determined through sequential assign-
Side chain assignments were determined with 13C/15N CARD9−142 using 13C- and 15N-HSQC (aliphatic and aromatic), (1H/15N)-CO/NH, HCCH-TOSY, HCCH-COSY, and 13C- and 15N-HSQC (aliphatic and aromatic, 120 ms mixing times) experiments, all collected in 99.9% D2O buffer, as well as a 13C-NOESY-TROSY experiment collected in 10% D2O. Stereoscopic assignments of valine and leucine methyl groups were determined by generating protein with a 1:9 13C/12C unlabelled glucose ratio and collection of 13C-HSQC spectra as described by Senn et al.13. To determine structure, intermolecular NOE assignments were collected using an intermolecular 13C-NOESY-HSQC experiment that allows for selective detection of NOEs between 13C- and 13C-bound protons.47 The sample for this experiment was prepared by mixing equimolar concentrations of denatured 13C15N and unlabeled CARD9−142 after elution from the Ni-NTA column and continuing with an otherwise identical purification protocol. The sample was concentrated to 2 mM for data collection. As only 25% of the sample as prepared is detectable by this experiment (i.e., only the 13C15N protein in a labeled/unlabeled dimer), and ~1% of the carbons in the sample are differentially labeled (i.e., 13C in an "unlabeled" protein due to natural abundance or 15N in a "labeled" protein due to lack of complete isotopic purity in 13C-glucose), intermolecular NOEs were included conservatively, only when peaks were relatively intense as compared to a standard 13C-NOESY-HSQC collected on a fully labeled sample. Concentration to 2 mM led to substantial line-broadening due to protein−protein interaction; this experiment was therefore collected with 900 mM NaCl rather than 300 mM to mitigate these interactions. Only cross-peaks for which minimal chemical shift perturbations were observed between 300 and 900 mM NaCl were included in the structure calculation.

N-H residual dipolar coupling (RDC) restraints were also collected on the CARD9−142 dimer. Samples were generated with 800 mM H13C15N−CARD9−142, with and without 14 mM Pt(bipy)2Cl2 (Asla BioTech). TROSY and anti-TROSY peaks were collected in an IPAP manner. RDC values were determined as the difference in δN between the sample with and without added bacteriophage. The alignment tensor was determined iteratively during structure determination using the CYANA v3.97 FitTensor protocol36. An additional 13C-NOESY-HSQC (mix times ranging from 3 to 10 s, 13C in 90% H2O/10% D2O buffer) experiment was collected to assist in structure determination. All spectra were referenced directly (proton) or indirectly (nitrogen and carbon) to an internal DSS standard. All spectra were processed using Bruker TopSpin v3.5 and subsequently analyzed in CcpNMR Analysis v2.49.

For structure determination, dihedral angles were estimated using TALOS−v1.252,53 and the WaterRefCNS package developed by Robert Tejero. The 20 lowest energy structures were evaluated using PROCHECK-NMR, with statistics presented in Table 1. Backbone amide assignments were transferred to CARD92−152/SSC in 50 mM HEPES, 150 mM NaCl, pH 7.0, and 0.5 mM TCEP, with 0.7 molar equivalents of residual dipolar coupling restraints collected on CARD92−152/SSC and UbG76C. This mix was passed over a HiTrap desalting column (GE Healthcare) equilibrated in 50 mM HEPES, 150 mM NaCl, pH 7.0. BMOE was solubilized at 20 mM in DMSO and added incrementally to the solution, with 0.1 molar equivalents (as compared to the total protein concentration) were added at a time, followed by mixing and 30 s of incubation at room temperature. BMOE was added in this manner past saturation, to a final concentration of 0.7 molar equivalents, followed by neutralization by excess β-mercaptoethanol. The desired CARD9−152/SSC−UbG76C conjugate was purified from other off-target products with a Superdex 200 gel filtration column (GE Healthcare) in 50 mM HEPES, 300 mM NaCl, 0.5 mM TCEP, pH 7.0 (See Supplementary Fig. 2A).

CARD9−152 and CARD11−172 filament formation. CARD9−152 constructs were purified bound 1:1 to Zn2+ in 50 mM HEPES, 150 mM NaCl, 0.5 mM TCEP, pH 7.0. A twofold excess of EDTA was added to the indicated concentrations of protein, followed by incubation at the indicated temperature for the indicated time. CARD9−172 was purified in 20 mM Tris, 150 mM NaCl, 0.5 mM TCEP, pH 7.5, followed by incubation at either 4 μM or 1 μM as indicated for 16 h at 4°C.

Bcl10 fluorescence polarization assay. The assay was conducted as previously described31 in 20 mM Tris, 150 mM NaCl, 0.5 mM TCEP, pH 7.5 in a 20 μl final volume. Purified MBP-Bcl10 sparsely labeled with AF488 C8 maleimide (see above) was incubated alone or with addition of CARD9 or CARD11 constructs prepared as described in the text. For the indicated sample, TEV protease was added to 0.05 mg/ml at the initiation of the assay. Bcl10 polymerization was monitored by measuring fluorescence polarization at 519 nm while exciting at 495 nm on a Molecular Devices SpectraMax M5e plate reader at 25°C.

NS-EM sample preparation and imaging. All NS-EM samples were incubated on glow-discharged carbon on 400-mesh copper grids (Electron Microscopy Sciences), followed by staining with 2% uranyl acetate. Images were collected on a Talos F200C microscope operated at 200 kV equipped with a Ceta camera (Thermo- Fisher) at 2.006 Å/pixel (Supplementary Fig. 4D, right), 4.097 Å/pixel (Fig. 5 and Supplementary Figs. 3, 4A, 4D left/center, 4E, and 7), or 10.6 Å/pixel (Fig. 6 and Supplementary Fig. 6). CARD9−152 samples in Fig. 4a were diluted to 50 μM just before grid preparation. CARD11−172 samples in Supplementary Fig. 4a were diluted to 4 μM just before grid preparation. CARD11−189 samples depicted in Supplementary Fig. 7 were diluted to 100 μM just before grid preparation. Grids in Supplementary Fig. 4D, E were prepared directly from SEC fractions (Supplementary Fig. 4C, green) without dilution. Samples in Fig. 5b and Supplementary Fig. 4G were prepared identically as the corresponding FP assay, with 2 μM MBP- Bcl10, 2 μM CARD9−152/152T107S filaments, 10 μM CARD9−152/152L131L33 filaments, and/or 2 μM CARD11−172 filaments and incubated with 0.05 mg/ml TEV protease. After a 2-min incubation at room temperature, 4 μl of sample was transferred to the EM grid. For samples including antibody, monoclonal anti-HA antibody (1:1000 dilution, Cell Signaling 4440S), or a monoclonal anti-β-actin antibody (1:1000 dilution, Licor 926−2212), blots were visualized using an HRP- conjugated antibody (1:5000 dilution, Sigma NA934V, for CARD9 and CARD11), SuperSignal West Pico Chemiluminescent Substrate (ThermoScientific), and a PXi6 imager (Syngene) or film.

Hydrogen-deuterium exchange. Samples, purified in 50 mM HEPES, 300 mM NaCl, 0.5 mM TCEP, pH 7.0, were diluted ~15-fold into 10 mM phosphate buffer, 0.5 mM DCl pH 7.0 with 150 μl of 100% D2O and incubated at 20°C for times ranging from 30 s to 1 day. Exchange was arrested by 1:1 dilution with a pH 2.3 buffer containing 400 mM Glycine and 4 M
Guanidinium chloride. For measurement of carried deuterium, 100 μl of this mixture was immediately passed through an immobilized pepsin column (21.3 × 30 mm, Applied Biosystems) and peptides produced were bound to an online trap column for desalting (Acquity Vanguard C8), separated by a reverse-phase chromatography (Acquity UPLC BEH C18, 1.7 μm particle size, 1.0 × 50 mm) all at 0 °C, and then introduced into the mass spectrometer by electrospray ionization (Thermo-Fisher Orbitrap XL, 60k resolution at m/z 200). A LEAP Pal X7 robotics platform was used to automate sample preparation, quenching, and handling. Measurements generally followed protocols described previously. Extracted ion chromatograms were produced by the ExMS program and analyzed by custom python scripts to extract the amount of carried deuterium by each peptide at each time, product. Pairwise parameters determined in RELION were used to generate nine symmetry-mates according to the determined helical parameters. The map was iteratively refit in map refinement scheme.

Cryo–EM data collection and analysis. In all, 0.5 mM CARD92–152/I107E purified in 50 mM HEPES, 150 mM NaCl, 0.5 mM TCEP, pH 7.0 bound 1:1 to Zn2+ was incubated for 10 min at 25 °C after addition of 1 mM EDTA. This solution was diluted to 0.1 mM in the same buffer, 3 μl of which was added to glow-discharged Protocips C-Flat 2/1 200 mesh holey carbon grids. After a 45 s incubation, the grid was washed and blotched six times with buffer, followed by a final 3.5 μl buffer addition and plunge freezing in liquid ethane using a Vitrobot Mark IV (ThermoFisher).

The sample was imaged on a Titan Krios Cryo Transmission Electron Microscope (ThermoFisher) operated at 300 kV and equipped with a K2 Summit direct electron detector camera, and a Bioquantum energy filter (Gatan). Movies were recorded at a magnification of 80k, corresponding to 1.2 Å per pixel. Forty frames were collected over 10 s, with an exposure rate of 5.093 e/Å2/s at between 1.25 and 2.5 μm defocus with an energy slit width of 20 eV.

See Supplementary Fig. 5A for a graphical depiction of the refinement scheme. Four-thousand four-hundred and three movies were collected and aligned using cisTEM. Twelve-thousand three-hundred and eighty-six individual micrographs were produced by the ExMS program and analyzed by custom python scripts to extract the amount of carried deuterium by each peptide at each time, product. The helical parameters were optimized in RELION by searching between a 4.8 to 2.5 Å defocus range, yielding a final resolution limit of 5.11 Å. Using a 5.0 Å high-resolution limit for particle alignment, refinement in Relion yielded a 4.0 Å reconstruction and −101.6° rotation, along with a −60.0 Å reconstruction. This volume was imported into cisTEM, in which further refinement in C1 (no helical symmetry applied) was carried out, leading to a −4.2 Å reconstruction. This volume and particle stack were finally exported into helical Frealign 9.1 using the refined parameters determined in RELION. Using a 5.0 Å high-resolution limit for particle alignment, refinement in Frealign yielded a 4.0 Å reconstruction (see Supplementary Fig. 5B for FSC curve). A local resolution estimate was determined using local resolution, a re-implementation (Rohub, manuscript in preparation) of blocres.20 The reconstruction was sharpened in cisTEM using a pre-cutoff B-factor of −90 Å2.

“Fit in map” in UCSF Chimera was used to fit the previously determined, lowest energy NMR solution structure (PDB ID 6E26) into the sharpened density and to generate nine symmetry-mates according to the determined helical parameters. The map was iteratively refitted in Phenix and COOT, maintaining strict non-crystallographic symmetry among the CARDS. Structural deformations were generated in UCSF Chimera or Pymol (pymol.org).

CARD92–97 filament formation. The in vitro CARD92–97 polymerization assay (Fig. 6d) was performed in 50 mM Tris, 150 mM NaCl, 0.5 mM TCEP, pH 7.0 as previously described.25 All CARD92–97 constructs utilized were purified bound to equimolar Zn2+ by addition of ZnCl2 prior to the final gel filtration column. Fifty microliters of CARD92–97WT or mutants were prepared at the indicated concentrations in a 384-well clear-bottom plate. The assay was initiated by addition of a super-stoichiometric concentration of EDTA as indicated. Polymerization was monitored using either monitoring optical density at 350 nm on a Molecular Devices SpectraMax M5e plate reader while shaking at 25 °C. At the 90 min endpoint of the assay, NS-EM grids were generated.

CARD18–109 filament formation. WT or mutant CARD18–109 constructs were purified in 50 mM HEPES, 50 mM NaCl, 0.5 mM TCEP, pH 7.0, concentrated to either 250 μM or 500 μM, and incubated at 25 °C for 2 h prior to grid preparation.
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