Structural basis for specific flagellin recognition by the NLR protein NAIP5

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The nucleotide-binding domain- and leucine-rich repeat (LRR)-containing proteins (NLRs) function as intracellular immune receptors to detect the presence of pathogen- or host-derived signals. The mechanisms of how NLRs sense their ligands remain elusive. Here we report the structure of a bacterial flagellin derivative in complex with the NLR proteins NAIP5 and NLRC4 determined by cryo-electron microscopy at 4.28 Å resolution. The structure revealed that the flagellin derivative forms two parallel helices interacting with multiple domains including BIR1 and LRR of NAIP5. Binding to NAIP5 results in a nearly complete burial of the flagellin derivative, thus stabilizing the active conformation of NAIP5. The extreme C-terminal side of the flagellin is anchored to a sterically constrained binding pocket of NAIP5, which likely acts as a structural determinant for discrimination of different bacterial flagellins by NAIP5, a notion further supported by biochemical data. Taken together, our results shed light on the molecular mechanisms underlying NLR ligand perception.

Keywords: flagellin; NAIP5; NLRC4; cryo-EM

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Introduction

The nucleotide-binding domain (NBD)- and leucine-rich repeat (LRR)-containing (NLR) proteins are a family of intracellular receptors that play an important role in regulation of innate immune response [1-4]. NLR proteins share a conserved tripartite domain structure with an N-terminal protein-protein interaction domain, a central nucleotide-binding and oligomerization domain (NOD) and a variable number of C-terminal LRR [1]. Similar domain structure is also present in plant NLR-type receptors [5]. In animals, several NLRs have been shown to function as pattern recognition receptors (PRRs), detecting pathogen-associated molecules patterns (PAMPs) or host-derived danger signals in the cytosol and consequently initiating innate immune response [3, 6]. Following ligand perception, these immune NLRs oligomerize to form multiprotein complexes termed inflammasomes for activation of caspase-1 [7, 8]. Once activated, caspase-1 promotes proteolytic cleavage and secretion of IL-1β and IL-18. The activated caspase-1 can also cleave the substrate gasdermin D to induce pyroptosis, an inflammatory form of cell death [9-11]. Despite the important roles of NLRs in detecting the presence of pathogens in both animals and plants [5], the mechanisms of how NLRs sense pathogen-derived ligands still remain poorly understood.

Neuronal apoptosis inhibitory protein (NAIP)-NLR containing a caspase activating and recruitment domain
Flagellin recognition by NAIP5

NAIP5-NLRC4 complex

To determine how NAIP5 recognizes flagellin, we first purified an *Salmonella typhimurium* (*S. typhimurium*) flagellin derivative with its N- and C-terminal regions fused together (called FliC_D0) [32] in complex with wild-type NAIP5 and an NLRC4 carrying the mutations R288A-L435D-1008-1012DDYD-AAAA (called NLRC4<sup>mut</sup>) from insect cells (Supplementary information, Figures S1 and S6A) as previously described [14]. The protein purified was then used for structural analysis with cryo-EM (Figure 1; Supplementary information, Figure S2). After 3D classification, a subset of 1 663 317 particles was used for image reconstruction, generating a map with a global resolution of 4.28 Å (Supplementary information, Figures S2, S3 and Table S1), as determined with a gold standard Fourier shell correlation (FSC) (Supplementary information, Figure S3B). The resolution is anisotropic (Supplementary information, Figure S3C) with BIR1, NBD, HD2, WHD, the unannotated domain [24] (residues 921-980, call “ID” hereafter) and FliC_D0<sub>L</sub> at resolution of ~3.8-4.5 Å as supported by the visibility of larger side chains (Supplementary information, Figures S4 and S5). In contrast, LRR, BIR2 and the region N-terminal to BIR1 (residues 1-60, called “NTD” hereafter) have a lower resolution, in the range of ~5-8 Å. Compared to NAIP5, NLRC4<sup>mut</sup> is less well defined in the 3D reconstruction (Supplementary information, Figure S2). Structural comparison between NAIP5-NLRC4<sup>mut</sup> and a lateral NLRC4 dimer from NLRC4 inflammasome [14-16] showed that a conserved set of structural elements (from NBD and WHD) of NAIP5 and NLRC4 was involved in interaction with NLRC4 (Supplementary information, Figure S6). We therefore limit our discussion to NAIP5 interaction with FliC_D0<sub>L</sub>.

**Structure of the FliC_D0<sub>L</sub>-bound NAIP5 and its comparison with that of active NLRC4**

The NBD, HD1, WHD, HD2 and LRR domains of NAIP5 (Figure 2A) are similarly positioned to those of an active NLRC4 (Figure 2B), indicating the structure of NAIP5 represents an active state. Located between NBD and HD1, clear electron density that is not from NAIP5 likely defines an ATP molecule (Supplementary information, Figure S4A and S4B). The ATP is coordinated by residues exclusively from NBD and HD1 of NAIP5 (Supplementary information, Figure S4C). BIR1 and BIR2 are juxtaposed at one side of NAIP5 (Figure 2B, left panel), with the former making contacts with HD1 and the latter with NBD. Compared to BIR1 and BIR2, BIR3 of NAIP5 is much less well defined in the 3D reconstruction and only fuzzy density is observed within the surface groove created by BIR1, BIR2, NBD1 and

**Results**

**Cryo-EM structure of a flagellin-induced heterodimeric**

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HD1 (Supplementary information, Figure S7A and S7B). The BIR3 domain is located at the opposite side of the NAIP5 surface interacting with an active NLRC4 [14-16] and could play a role in contacting the last NLRC4 molecule for closure of the ring-like NAIP-NLRC4 inflammasome as suggested by structural comparison (Supplementary information, Figure S7C). The structures of BIR1 and BIR2 of NAIP5 resemble that of BIR3 from XIAP [33], but the peptide binding sites in these two BIR domains are completely blocked (Supplementary information, Figure S8).

Although structural domains are similarly positioned in the active NLRC4 and NAIP5, striking structure differences between them exist. In addition to NTD and the BIR domains, NAIP5 also contains an ID and extra 19 residues at its C-terminal side. The ID is mainly composed of three helices, with the middle one packing against the structural elements formed by the additional 19 residues of NAIP5 (Figure 2B, left panel). The middle helix and its surrounding regions of ID also contact one lateral side of the extreme C-terminal region of LRR, resulting in closure of the horse-shoe-like structure of the LRR domain. Another striking difference between NAIP5 and NLRC4 occurs in the regions around the conserved four-helical bundle of HD2 (Figure 2B). In the active NLRC4, a short helix is extended to establish interaction with the inner surface of its LRR (Figure 2B, right panel). By comparison, two helices pack against the conserved four-helical bundle of HD2 at a different side and interact with HD1 and WHD of NAIP5 (Figure 2B, left panel). The simultaneous interactions of these two NAIP5 helices with HD1 and WHD are expected to sta-
Figure 2 Atomic model of the FliC\_D\_0\_bound NAIP5. (A) Schematic representation of the domain structures of NAIP5. Color codes for domains are indicated. Numbers indicate the domain boundaries. (B) Cartoon representations of active NAIP5 (left panel) and active NLRC4 (right panel). The aligned NAIP5 and NLRC4 are shown in the same orientation. Red frame indicates the location where C-terminal of NAIP5 packs with middle helix of ID. (C) Structural superposition of a lateral NLRC4 trimer from NLRC4 inflammasome with NAIP5 shown in two different orientations. NAIP5 was aligned with the middle NLRC4 protomer. For clarity, NTD, BIR1 and BIR2 are shown in the same colors as in B and labeled and all the other domains of NAIP5 are shown in pink. ID, insertion domain; NTD, N terminal domain.
bilize the active conformation of NAIP5, because striking structural reorganization between these two domains has been demonstrated for NLRC4 during activation [14-16].

Structural alignment of NAIP5 with a lateral trimer from the NLRC4 inflammasome showed that the NAIP5-bound FliC_D0L unlikely interacted with NLRC4 directly (Figure 2C), suggesting that the flagellin derivative allosterically activates NLRC4 to assemble NAIP5-NLRC4 inflammasome. This structural comparison also revealed that BIR1 and BIR2 are located far from the NAIP5-NLRC4 dimeric interface, indicating that these two domains are less likely to make a direct contribution to NAIP5-NLRC4 oligomerization.

**Overall structure of the FliC_D0L-NAIP5 complex**

The N- and C-terminal sides of FliC_D0L (termed FliC-N and FliC-C, respectively) form two parallel α helices with few interactions formed between them (Figure 3A), as observed in the structure of full-length flagellin [34]. Both of the two helices interact with NAIP5, resulting in a nearly complete burial of FliC_D0L (Figure 3B). FliC forms extensive interactions with NAIP5 via packing against NTD, BIR1, HD1, HD2, ID and LRR of NAIP5 (Figure 3B), supporting the observation that the C-terminal 35 residues of flagellin are necessary and sufficient for NAIP5 activation [21]. Participation of the non-conserved NTD, BIR1 and ID domains in interaction with FliC-C explains specific recognition of flagellin by NAIP5 but not by NLRC4. Compared to FliC-C, FliC-N forms much less dense contacts with NAIP5 (Figure 3B). The helical portion of FliC-N is sandwiched between HD2 and the inner surface of LRR of NAIP5 (Figure 3B), although identities of the residues from LRR cannot be unambiguously determined in the density map. In addition, the N-terminal side of FliC-N binds to a surface groove formed by HD1 and HD2 (Figure 3B), which likely also contributes to stabilization of the active conformation of NAIP5. Interestingly, NBD and WHD that are important for oligomerization of the NAIP5-NLRC4 inflammasome [14-16] are not involved in NAIP5 recognition of FliC_D0L.

**Recognition mechanism of FliC_D0L by NAIP5**

The extreme C-terminal side of FliC-C binds to a deep hydrophobic pocket formed by NTD, BIR1 and HD1 (Figures 3A and 4A). Four hydrophobic residues, L491, L493, L494 and V490 from this region of FliC-C contacts their neighboring residues from these three domains of NAIP5 (Figure 4B). Supporting this structural observation, simultaneous substitutions of L491, L493, L494 with alanines significantly compromised the activity of the C-terminal 35-residue peptide [21] or full-length flagellin [12, 13] in activating NLRC4-mediated immune response. The central region of FliC-C is sandwiched between ID and one helix from HD2 of NAIP5 mainly via van der Waals and hydrophobic contacts (Figure 4A, 4C and 4D). Similar types of interactions are also important for mediating interaction of the helix portion of FliC-N with HD2 (Figure 4C) and binding of the N-terminal side of FliC-N to the surface groove formed by HD1 and HD2. The extensive contacts of HD2 with FliC_D0L are consistent with previous data suggesting that the central NBD-associated domains of NAIP6 are crucial structural determinants for recognition of flagellin [31]. Structure-based sequence alignment indicate that the FliC_D0L-interacting residues of NAIP5 are highly conserved in NAIP6 but not in NAIP2 (Supplementary information, Figure S9), explaining the specific recognition of flagellin by NAIP5 or NAIP6 [12, 13].

**Mutagenesis analysis of NAIP5 responsiveness to flagellin**

We used the cell-based assay as previously described [13] to further verify our structure. Amino acids from NAIP5 that are important for the interaction with FliC_D0L were mutated and effects of these mutation on the activity of NAIP5 were monitored. As positive controls, FliC and FliC_D0L strongly promoted the production of IL-1β in 293T cells when co-expressed with wild-type NAIP5, NLRC4 and procaspase-1 (Figures 5 and 6). In contrast, deletion of the N-terminal 40 residues of NAIP5 that cap FliC-C (Figure 3A) resulted in no detectable production of mature IL-1β induced by FliC. Furthermore, substitution of the three residues (106-108) from BIR1 that recognize the C-terminal side of FliC_D0L together with other two non-conserved residues with their equivalents in NAIP2 greatly reduced FliC-induced maturation of IL-1β. These results agree with the observation that deletion of the BIR domains from NAIP5 resulted in loss of flagellin-induced activation of NAIP5-NLRC4 inflammasome [13] and support an important role of BIR1 in dictating ligand specificity of NAIP5. Consistently, swapping of BIR1 and BIR3 of NAIP5 led to no detectable FliC-induced production of IL-1β. In further support of our structure, mutations of Leu840 and Gly847 from the helix in HD2 that simultaneously packs against the two helices of FliC_D0L (Figure 4C) to their equivalents in NAIP1 also significantly reduced responsiveness of NAIP5 to FliC (Figure 5A). Similar observation was also made for the mutants of NAIP5 with Phe844 substituted by its corresponding Cys887 in NAIP2 and deletion of the C-terminal 14 residues, which contact the C-terminal side of FliC-C and stabilize the ID. In contrast, little effect on IL-1β maturation was observed for mutating
S857, a residue that is solvent-exposed and does not interact with FliC_D0L. Strikingly, the NAIP5 mutant F844C was partially responsive to PrgJ in mediating the production of IL-1β (Figure 5B), further supporting an important role of this domain in mediating specific NAIP5 recognition of flagellins. However, further replacement of residues surrounding Phe844 with those in NAIP2 did not enhance responsiveness of the resulting NAIP5 mutants to PrgJ. Collectively, these results support our cryo-EM structure.

**Mechanism of differential flagellin recognition by NAIP5**

Structure-based sequence alignment showed that the NAIP5-interacting residues of FliC_C are highly conserved among bacterial flagellin (Figure 6A). However, flagellins from different bacteria vary significantly in their NAIP5-interacting and NLRC4-inducing activities [13]. It is of interest to note that flagellins with higher activities have the arginine residue at their C-termini, whereas this residue is substituted with Gln or Gln-Gly in those with lower activities (Figure 6A). In the structure, the arginine residue of FliC_D0L binds to a pocket of NAIP5 with a limited size (Figure 3B) and tightly packs against residues 106-109 from BIR1 (Figure 4B). These data suggest that the last arginine residue of flagellin may be important for its NAIP-NLRC4 inflammasome-inducing activity. In support of this possibility, mutation of the last residue Arg495 of FliC_D0L to Gln significantly reduced its activity of inducing NAIP5 activation (Figure 6B). The activity was further decreased by addition of glycine to this FliC_D0L mutant, likely because of steric effect caused by the limited size of the pocket recognizing the extreme C-terminal side of FliC_D0L. Consistently, introduction of Gly at the C-terminus of FliC_D0L substantially compromised the peptide activity in inducing NAIP5 activation. However, the *Legionella pneumophila* flagellin with a glycine inserted before the last arginine residue has a similar activity to the *S. typhimurium* flagel-
Figure 4 Recognition mechanism of Flic_D0L by NAIP5. (A) A close-up view of the interaction of the C-terminal Flic_D0L (cartoon) with NAIP5 (transparent surface). Detail interactions between Flic_D0L and NAIP5 within the highlighted regions are shown in B-D. (B) Detailed interactions of the C-terminal side of Flic_D0L with NAIP5 highlighted within the red square in A. (C) Detailed interactions of the central region of Flic_D0L with NAIP5 highlighted within the blue square in A. (D) Detailed interactions of the C-terminal side of Flic_D0L with NAIP5 ID highlighted within the purple square in A.
Flagellin recognition by NAIP5

The reason for this might be that the C-terminal side of FliC\(_{D0}\) can be slightly kinked by introduction of the achiral glycine, thus allowing the C-terminal side to be accommodated by the NAIP5 pocket. Taken together, these results show that the last residue of flagellin is an important structural epitope recognized by NAIP5, although contributions from other positions to differential recognition of flagellins by NAIP5 are fully possible.

**Discussion**

The data presented here showed that optimal recognition of FliC\(_{D0}\) involves multiple structural domains of NAIP5, although their contributions to interaction with the flagellin can vary. Binding of FliC\(_{D0}\) functions to stabilize the active conformation of NAIP5, indicating that NAIP5 as a seeding NLR needs a stabilized active conformation to ensure NLRC4 activation. A similar function can also be expected for the ligands of NAIP1 and its human homolog hNAIP; and NAIP2, which have been shown to play similar roles in ligand-induced assembly of NAIP-NLRC4 inflammasomes. Future structural studies are needed to investigate how these NAIPs specifically recognize their respective ligands. The observation that mutations of critical residues in HD2 of NAIP5 to their equivalents in NAIP1 significantly compromised FliC-induced activation of the NAIP5-NLRC4 inflammasome (Figure 5A) suggests that this structural domain may be also important for NAIP1 and hNAIP recognition of their ligands. In contrast with flagellin, cytochrome c is not involved in stabilizing the active conformation of APAF-1 in the APAF1 apoptosome [35, 36]. It is noteworthy to mention that, however, assemblies of NAIP5-NLRC4 inflammasome and APAF-1 apoptosomes follow different mechanisms [14, 15]. An active conformation-stabilizing role of FliC\(_{D0}\) in NAIP5

![Figure 5 Mutagenesis analysis of NAIP5 responsiveness to flagellin.](image-url)

(A) Mutagenesis analyses of NAIP5 responsiveness to FliC. 293T cells were transfected with plasmids as indicated. 24 h after the transfection, the culture medium was supplemented with PA and LFn-FliC proteins with final concentration of 3 \(\mu\)g/mL. The cells were harvested and lysed, and the cleaved IL-1\(\beta\) was detected by anti-IL-1\(\beta\) immunoblotting analysis after 12 h. GAPDH was used as a loading control. Trans837-845: 837-845 residues of NAIP5 replaced with 880-888 residues of NAIP2. BIR321: exchange of BIR1(61-129) and BIR3(278-345) without changing linker region. (B) Mutagenesis analyses of NAIP5 responsiveness to PrgJ. The assays were performed as described in A.
activation is in line with the observation that the constitutively active plant NLR L6 displayed a higher ligand binding affinity than wild-type protein [37]. In contrast with those in the NAIP5-NLRC4 inflammasome [13-16], NLRC4 from the dimeric NAIP5-NLRC4 complex is much less defined, indicating that oligomerization of NLRC4 is important for stabilization of its active conformation. These results appear to suggest that stabilization of the active conformation of an NLR protein can play an important role in its activation. But more studies are needed to determine whether ligand recognition or perception by other NLRs has a similar function to flagellin binding to NAIP5.

Modeling studies showed that the inactive NAIP5 has a similar conformation to that of the inactive NLRC4 [31, 38]. Then how is flagellin initially recognized by an inactive NAIP5? Structural comparison between the active NAIP5 and a modeled inactive NAIP5 (Supplementary
information, Figure S10) suggested that the FliC_D0_L binding pocket formed by BIR1 and HD1 is an attractive site for initial recognition of the extreme C-terminal side of FliC_D0_L because other FliC-D0 binding sites are largely occluded by the positioning of LRR in the inactive NAIP5. This can afford an explanation for the ability of NAIP5 to differentiate bacterial flagellins with subtle differences in their C-terminal sides. However, we cannot exclude the possibility that inactive NAIP5 assumes a structure strikingly different from the predicted one. Nonetheless, nearly complete burial of FliC_D0_L indicates that the conformation of inactive NAIP5 should be different from its active one. Thus, FliC_D0_L binding is expected to trigger structural re-organization to the active conformation of NAIP5 for full interaction with flagellin. Conformational selection, proposed for self-activation conformation of NAIP5 for full interaction with flagellin. In this case, NAIP5 may adopt a metastable active state that exists in an equilibrium with its more stable inactive state. Stabilization by flagellin binding can shift the equilibrium toward the active state of NAIP5.

Structural studies showed that dATP/ATP acts to stabilize the active conformation of APAF-1 via the γ-phosphate of the bound dATP/ATP, supporting an essential role of dATP/ATP in assembly of the APAF-1 apoptosis [35, 36]. However, the γ-phosphate of the NAIP5-bound ATP does not interact with other domains than the NBD and HD1, indicating that ATP makes no direct contribution to the stabilization of the active conformation of NAIP5. Mutations [32] of P-loop showed that nucleotide binding of NAIP5 is not essential for formation of NAIP5-NLRC4 inflammasome. Our previous studies [14, 38] suggested that ATP binding may be a consequence of ligand-induced conformation changes in NLRC4. Collectively, these data suggest that a role of the bound ATP in NLR activation might be associated with whether it acts to stabilize the ligand-induced active conformation of the NLR protein [16].

Our structural and biochemical data showed that the last arginine residue from the tested flagellins with higher NLRC4 inflammasome-inducing activity [13] is an important epitope recognized by NAIP5. Mutation of this residue in S. typhimurium flagellin has been shown to impair bacterial mobility [39], suggesting a correlation between the functional significance of the arginine residue as a critical moiety of PAMP for bacteria and its recognition by NAIP5. A similar correlation has been demonstrated for flagellin recognition by TLR5 [39, 40]. Some bacterial flagellins possess different residues at this position and display much lower NLRC4 inflammasome-inducing activity [13], suggesting that these alterations might be a bacterial strategy to evade host immune recognition by NAIP5. On the other hand, at least for EPEC and Burkholderia thailandensis carrying such altered flagellins, flagellin-independent caspase-1 activation has been demonstrated during infection [13], probably reflecting an evolutionary arms race between bacteria and their hosts.

### Materials and Methods

**Plasmids and antibodies**

Full-length NAIP5 and NAIP2 genes were obtained by DNA synthesis. Mouse NLRC4, pro IL-1β and pro caspase-1 were amplified from reverse-transcribed mouse cDNAs. Full-length PrgJ and FliC genes were amplified from S. typhimurium genomic DNA. NLRC4-R288A-L435D-1 008-1 012<sup>DDYD</sup>AAAA(NLRC4<sup>44</sup>), all NAIP5 and flagellin mutations were generated by standard molecular biology procedures. FliC_D0_L was designed as described before [32]. All constructs were verified by sequencing.

Antibody used: anti-Myc (cw0299, CWBIO), anti-HA (cw0092, CWBIO), anti-IL-1β (GTX74034, GeneTex) and anti-GAPDH (cw0100, CWBIO).

**Recombinant protein expression and purification**

A heterodimeric complex between NAIP5 and NLRC4<sup>44</sup> induced by FliC_D0_L was purified from insect cells as described previously [14]). Briefly, the related genes were cloned into modified pFastBac vectors containing cleavable N-terminal GST or N-terminal 6× His-SUMO tag and co-expressed in sf21 insect cells (Invitrogen). SF21 cells were grown in SF900 (GIBICO) medium by shaking at 120 rpm at 28 °C until the density reached 2.0 × 10<sup>6</sup>/mL. One liter of cells (2.0 × 10<sup>9</sup>/mL) were infected with 22 mL of recombinant baculovirus. Cells were harvested after 48 h of infection, re-suspended in the buffer containing 25 mM Tris (pH 8.0), 150 mM NaCl and 1 mM PMSF, and lysed by sonication before centrifugation. The supernatant was flowed through glutathione sepharose 4B beads (Invitrogen). The bound proteins were digested with PreScission protease (GE Healthcare) to remove the GST tag and further purified by size exclusion chromatography (Hiload 16/60 Superdex 200 prep grade, GE Healthcare) in a buffer containing 10 mM Tris (pH 8.0) and 150 mM NaCl.

PrgJ, FliC, and FliC_D0_L mutants used in 293 assays were fused with N-terminal 1-263 aa of lethal factor (LF<sub>L</sub>) and cloned to pET15b vector. *Escherichia coli* BL21(DE3) strains transformed with the expression plasmids were grown in LB medium and induced overnight at 18 °C with 0.6 mM IPTG after OD<sub>600</sub> reached 0.7. Bacteria were harvested, re-suspended and lysed in the buffer containing 25 mM Tris (pH 8.0) and 150 mM NaCl. The soluble fraction was purified by Ni-NTA (Novagen) and further purified by size exclusion chromatography (Superdex 200, GE Healthcare) in a buffer containing 10 mM Tris (pH 8.0) and 150 mM NaCl.

**Analysis of NLRC4 inflammasome in 293T cells**

mNAIP5, mNAIP2, mouse pro IL-1β, pro caspase1 and NAIP5 mutations were cloned to pcDNA3.1 vector. 293T cells were seeded to six-well plate 12 h before transfection. The complex of plasmids containing 1 μg pro IL-1β, 50 ng pro caspase 1, 50 ng NLRC4, NAIP5 and 2 μl vigofect reagent (Vigorous) per well were
added to the culture supernatant. After 24 h of transfection, PA and LF, fused proteins were added to the culture medium with a final concentration of 3 μg/mL. After another 12 h, the transfected cells were harvested in the lysis buffer containing 25 mM Tris 7.4, 150 mM NaCl, 1% Triton X-100 and 1× protease inhibitor cocktail (Thermo Scientific). The cleaved IL-1β was detected by anti-IL-1β immunoblotting analysis. All the reconstitution experiments were performed at least three times.

Cryo-EM sample preparation and data collection

The purified FliC_D0,NAIP5-NLRC4 complex was concentrated to ~2 mg/mL. Then aliquots of 4 μl of this sample were applied to the glow-discharged Quantifoil R1.2/1.3 holey carbon grids (Quantifoil Micro Tools GmbH), blotted for 2 s and plunge-frozen by FEI Vitrobot Mark IV. Grids were examined using an FEI Titan Krios operated at 300 kV, and images were recorded using a K2 Summit direct electron detector (Gatan) in super-resolution mode, at a nominal magnification of 22,500, and with the defocus ranging from ~1.5 to ~3.0 μm. Images were collected under low-dose condition in a semiautomatic manner using UCSF-Image4 [41]. The dose rate on the camera was set to be 8.2 electrons per pixel per second. For each micrograph stack, a total of 32 frames were collected with an exposure time of 8 s, leading to a total accumulated dose of 50 electrons per Å² on the specimen.

Cryo-EM image processing and refinement

The collected original micrographs stacks were aligned and summed using whole image-motion correction [41], and binned two-fold, resulting in a pixel size of 1.30654 Å per pixel. The defocus values of micrographs were estimated by CTFIND3 [42]. The collected particles were picked by EMAN2 [43] and RELION 1.4 [44, 45]. 2D and 3D classifications and refinements were performed using RELION 1.4 [44, 45]. A total of 123,876 particles were first boxed from 525 micrographs of the FliC_D0,NAIP5-NLRC4 complex using e2boxer.py in EMAN2 [43], respectively. Then, the boxed particles were extracted and reference-free 2D class averaging was performed. The generated 2D class averages were used as the templates for the subsequent autopicking of particles of Naip5-Nlrc4 complex using Relion 1.4 [44, 45]. A total of 2,192,107 particles of the NAIP5-NLRC4 complex were automatically picked out from 2,864 micrographs, respectively, and 2D classified using Relion 1.4 [44, 45]. About 1,663,317 particles were empirically selected from 2D classifications using and auto-refined against model from RELION 1.4 [44, 45]. About 1,663,317 particles were divided into 10 classes. The two most homogeneous classes of 245,315 particles and 381,293 particles were subjected to auto-refinement without any symmetry imposed and resulted in reconstructions at an overall resolution of 6.53 and 7.90 Å, respectively, based on the gold-standard FSC 0.143 criterion [46]. The original image stacks were then aligned and summed using whole image-motion correction2 [47], and binned two-fold. All the particles above had been replaced by particles from new summed micrographs from image-motion correction2, with keeping the alignment parameters already obtained from previous auto-refinement using RELION 1.4 [44, 45]. All of the new 245,315 and 381,293 particles were subjected to auto-refinement with local search methods using RELION 1.4 [44, 45], and at this stage the resolution of these two classes of particles were refined to 4.93 and 4.58 Å, respectively. A soft mask was applied around the rigid part of NAIP5, which further improved resolution to, 4.42 and 4.51 Å, respectively. All the particles of the two classes were then merged together for auto-refinement with the soft mask imposed around the rigid part of NAIP5 and resulted in a reconstruction at a resolution of 4.28 Å. Additional cycles of 3D classifications and refinements did not further improve the overall resolution of the maps. Local resolution was estimated using ResMap [48].

Model building and structure refinement

To build the NAIP5 model, the previously reported activated NLRC4 structure and baculoviral IAP repeat-containing protein 7 (PDB 1TW6, chain A) were used as the initial models. The initial NLRC4 model, in which the conserved residues in NLRC4 were retained and the non-conserved residues were substituted with alanine residues using CHAINSAW [49]. We individually separated the NBD, HD1, WHD and LRR domains of NLRC4 and rigid body fitted these domains into the EM map in COOT [50]. The chain A of baculoviral IAP repeat-containing protein 7 was fitted into the electron density from the BIR1 and BIR2 domains. Residues 921-980 from the ID of NAIP5 were manually built into the EM density in COOT. De novo model building was performed for FliC_D0, by manually docking two alpha helices into the electron density. Residue assignment of FliC_D0 was guided by its last arginine and other bulky residues.

The molecular dynamics flexible fitting (MDFF) [51] method was used to flexibly fit the atomic structure into the density map in VMD after model building in COOT. The model was finally refined against the map at overall 4.28 Å using phenix.real_space_refine application in PHENIX [52] in real space with secondary structure and geometry restraints to prevent structure over-fitting. Residues with Ramachandran outliers were further manually adjusted in COOT. The coordinates of the final model (FliC_D0,NAIP5-NLRC4 complex) were randomly displaced by 0.5 Å using the Phenix (PDB Tools) to remove potential model bias. The displaced model was then refined against one of the half maps (produced from a half set of all particles during refinement by RELION). FSC curves were calculated between the resulting model and half1 map (model versus half1 map, FSCwork, that is, used for refinement), the resulting model and half2 map (model versus half2 map, FSCfree, that is, not used for refinement) and the resulting model and the final density map (model versus summer map) from all particles. The lack of significant separation between work and free FSC curves suggested that the models were not over-fitted.

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Author Contributions

JC and S-FS conceived, designed the project and wrote the manuscript. JC and S-FS supervised the project. ZH made initial contribution to the project. XY and GL purified the proteins for EM and performed 293 assays. XY and FY performed cryo-EM sample preparation and data collection. FY calculated the cryo-EM map and WW built the atomic model. All authors contributed to project discussion and manuscript preparation.

Competing Financial Interests

The authors declare no competing financial interests.

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